CONTRIBUTORS TO THE HAND, WRIST, AND FOREARM DISORDERS GUIDELINE

Editor-in-Chief:
Kurt T. Hegmann, MD, MPH, FACOEM, FACP

Evidence-based Practice Hand, Wrist, and Forearm Panel Chair:
J. Mark Melhorn, MD, FAAOS, FACOEM, FADEP, FACS, FASSH, FAAHS

Evidence-based Practice Hand, Wrist, and Forearm Panel Members:
James Ausfahl, MD
M. Felix Freshwater, MD
Charles P. Prezzia, MD, MPH, MMM, FACOEM
David M. Rempel, MD, MPH, FACOEM, FACP
Shawn C. Roll, PhD, OTR/L, RMSKS, FAOTA
Arlen J. Rollins, DO, MSc, FACOEM, FACPM
Robert A. Werner, MD, MS, FAAPMR
Jason Zaremski, MD, CAQSM

These panel members represent expertise in occupational medicine, orthopedic surgery, hand surgery, occupational therapy, physical medicine and rehabilitation, sports medicine, internal medicine, family practice, forensic medicine, and electrodiagnostic medicine. As required for quality guidelines (Institute of Medicine’s [IOM] Standards for Developing Trustworthy Clinical Practice Guidelines and Appraisal of Guidelines for Research and Evaluation [AGREE]), a detailed application process captured conflicts of interest. The above Panel has none to declare relevant to this guideline.

Methodology Committee Consultant:
Kurt T. Hegmann, MD, MPH, FACOEM, FACP

Managing Editors:
Production: Marianne Dreger, MA
Research: Julie A. Ording, MPH
Research Conducted By:
Kurt T. Hegmann, MD, MPH, FACOEM, FACP
Matthew A. Hughes, MD, MPH
Matthew S. Thiese, PhD, MSPH
Ulrike Ott, PhD, MSPH
Deborah Gwenevere Passey, PhDc, MS
Atim Effiong, MPH
Kristine Hegmann, MSPH, CIC
Alzina Koric, MPP
Brenden Ronna, BS
Austen J. Knudsen
Pranjal A. Muthe
Skyler Walker
Anh Tran
Jenna K. Lindsey
Dillon J. Fix
Leslie MC Echeverria, BS
Jeremiah L. Dortch, BS

Specialty Society and Society Representative Listing:
ACOEM acknowledges the following organizations and their representatives who served as reviewers of the Hand, Wrist, and Forearm Disorders Guideline. Their contributions are greatly appreciated. By listing the following individuals or organizations, it does not infer that these individuals or organizations support or endorse the hand, wrist, and forearm treatment guidelines developed by ACOEM.

American Academy of Physical Medicine & Rehabilitation

American Association of Occupational Health Nurses
David A. Allcott, MSN, APRN, ANP-BC, COHN-S, FAAOHN

Association for Applied Psychophysiology and Biofeedback
Gabriel E. Sella, MD, MPH, MSC, PhD, FAADP, FAAFP, FACPM

American College of Emergency Physicians
Charles Gerardo, MD, MHS, FACEP

American Society of Anesthesiologists
Richard W. Rosenquist, MD

The American Occupational Therapy Association, Inc.
Debbie Amini, EdD, OTR/L, CHT, FAOTA
Hand, wrist, and forearm symptoms in the workforce are common problems presented to health care providers and are among the five most common causes of reported work-related health symptoms and workers’ compensation claims. According to 2010 US Census data, there was an incidence rate of 67.6 upper extremity fractures per 10,000 persons.(1) In 2013, there were 345,560 work-related upper extremity disorders for an incident rate of 32.5 per 10,000 full-time workers.(2) This was the leading cause of work-related injury,(2) and it is estimated that 20% of the population in any given month will complain of at least one type of upper limb disorder.(3) In 1998, a study involving more than 10,800 participants concluded that 30.5% had a self-reported neck or limb disorder.(4) Results from another study concluded that in 2000, 5.3 out of every 1,000 workers would take an absence due to sickness because of a musculoskeletal upper limb disorder; by 2004 this number had risen to 6.3.(5) These disorders account for nearly one-third (31.4%) of the missed days of work.(2) They also account for about 7 to 8% of total lost workdays in workers’ compensation and 17 to 23% of cases and claims, ranking them in the top five for financial severity.

Overview
Recommendations on assessing and treating adults with hand, wrist, and forearm disorders are presented in this clinical practice guideline. Topics include the initial assessment and diagnosis of patients with acute, subacute, and chronic hand, wrist, and forearm disorders that are potentially work-related, identification of red flags that may suggest the presence of a serious underlying medical condition, initial management, diagnostic considerations and special studies to identify clinical pathology, work-relatedness, modified duty and activity, and return to work, as well as further management considerations including delayed recovery. The majority of peer-reviewed literature categorizes acute as <1 month duration, subacute as 1 to 3 months duration, and chronic as >3 months duration. These definitions have been adopted throughout this document. Throughout the Guideline, we refer to costs in the rationale for the recommendation – costs are defined as low (<$100), moderate ($100-$500), and high (>500).

Algorithms for patient management are included and schematize how to generally manage acute, subacute, or chronic hand, wrist, and forearm disorders. It is important to realize that there are few studies that evaluate patients with work-related hand, wrist, and forearm disorders; therefore, studies that include different populations were used to develop the recommendations. In addition, most studies that focus on pharmaceuticals, appliances, and specific devices are industry sponsored. In certain areas, this may have made little difference as the comparisons were between the medication and placebo and the results may be stark. However, in other studies, the comparison groups may have been suboptimally treated (e.g., a low-dose of ibuprofen) and produced a bias in favor of the medication or device. In addition, industry-sponsored studies have been shown to frequently have better results and lower complication rates than studies conducted by independent investigators. Readers should refer to the Low Back Disorders, Chronic Pain and Opioids guidelines for more information regarding medications, including adverse effects.

General Approach and Basic Principles
The principal recommendations for assessing and treating patients with acute, subacute, or chronic hand, wrist, or forearm symptoms are as follows:

- The initial assessment focuses on detecting indicators of potentially serious disease, termed red flags, which require urgent assessment and treatment as indicated.
- The foci for treatment of patients with hand, wrist, or forearm symptoms include optimal medical care, monitoring for complications, facilitating the healing process, assisting stay at work or early return to work in a modified or full-duty capacity, and include surgical intervention(s) when indicated.
- Relieving discomfort can frequently and most safely be accomplished by modifying activities and using either topical or systemic nonprescription analgesics.
- Encourage patients recovering from hand, wrist, or forearm problems to stay at work or consider early return to modified work as their condition permits.
- Address occupational factors where the disorder is believed to be caused by work.
address nonphysical factors (e.g., psychosocial, workplace, or socioeconomic problems) in an effort to resolve delayed recovery (see Cornerstones of Disability Prevention and Management and Chronic Pain guidelines).

This Guideline addresses the following hand, wrist, and forearm disorders which may present to the health care provider.

Carpal Tunnel Syndrome
CTS is the most common and widely known of the entrapment neuropathies in which the body’s peripheral nerves are compressed or traumatized,(6-11) affecting an estimated 4 to 10 million Americans.(9) Carpal tunnel syndrome (CTS) occurs when symptoms occur that are attributable to abnormal median nerve compression within the carpal tunnel – a narrow, rigid passageway of ligament and bones at the base of the hand, which houses the median nerve and flexor tendons. The median nerve supplies sensations to the palmar aspect of the thumb, index, middle and radial half of the ring finger, as well as the dorsal segment of each of those four digits from the DIP distally, but not the fifth digit, as well as innervation to some small muscles (lateral two lumbricals, opponens pollicis, abductor pollicis brevis and flexor pollicis brevis.) in the hand that allow the fingers and thumb to move. Often, the condition arises without apparent cause.(6, 12, 13) Patients who have open injuries, unstable fractures, wrist fractures, or acute gout attack that results in acute CTS require immediate referral to a surgeon since improvement may only be obtained through surgery. Sometimes, synovial thickening around tendons or other swelling narrow the carpal tunnel and cause the median nerve to become variously compressed or enlarged through poorly understood processes. The result may be tingling, numbness, pain, or weakness in the digits. Tingling and numbness are essential symptoms. Pain is not an essential symptom and it may indicate other conditions, but if present, may also radiate proximally.

Triangular Fibrocartilage Complex (TFCC) Tears
Triangular fibrocartilage complex (TFCC) tears are frequent wrist injuries involving the cartilaginous meniscus between the radius and ulna with symptoms often described as occurring on the ulnar side of the wrist joint. TFCC is apparently susceptible to the same anatomic, pathophysiologic, and degenerative issues as the knee menisci. Vascular supply is similarly analogous to a meniscus with radial penetration into the meniscal periphery and central avascularity(14, 15) and evidence that degeneration increases with age.(16) Since abnormalities are commonly found on MRI and/or arthrography,(17) indications for surgical interventions are somewhat unclear.

Crush Injuries and Compartment Syndrome
Crush injuries as well as compartment syndrome are usually surgical emergencies.(18, 19) Mild cases of crush injuries may be treated similar to non-specific hand, wrist, forearm pain with particular emphasis on RICE (rest, ice, compression, elevation). Not all crush wounds, especially those more extensive and prone towards swelling are sutured as additional problems may ensue from suturing including possible tissue necrosis and the intervention may help to inhibit expansion to relieve pressure.

Kienböck Disease
Kienböck disease involves changes in the lunate that eventually lead to collapse of the lunate bone, which results in progressive pain and disability. It is a controversial condition from the standpoint of work-relatedness, as it is a disease and there are no quality studies on cause.

Wrist Sprains
Wrist sprains (which are partially or totally disrupted ligaments) are a common result of occupational slips, trips, and falls. Evaluation for occult fracture should be considered especially as fracture(s) may be present in a minority of cases.

Mallet Finger
Mallet finger is a common occupational and sports injury,(20) although it may occur with minimal apparent trauma.(21) The injury involves rupture of the extensor mechanism of a digit at the distal upper extremity
joint with or without fracture of the distal phalangeal segment. The mechanism of injury most typically involves forcefully striking the tip of the extended digit on an object including balls, or from falls.(22)

**Flexor Tendon Entrapment (Tenosynovitis and Trigger Digit)**

Stenosing tenosynovitis is a generally non-inflammatory condition caused by hypertrophy of the A-1 pulley with ensuing narrowing of the canal through which the digital flexors travel, with resulting symptoms of pain and snapping of the digit particularly with extension of a flexed digit.(23-28) While some cases are thought to be occupational(29) and many cases have been reportedly idiopathic, there are other less frequent reported causes and associated conditions, including diabetes mellitus, rheumatoid arthritis, CTS, Dupuytren’s disease, osteoarthritis, amyloidosis, hypothyroidism, heredity, and partial tendon laceration.(30-35) There also is evidence these may be connective tissue disorders.(36, 37) Diabetes mellitus reportedly portends a worse prognosis for glucocorticosteroid injection.(38, 39) The disorder includes a spectrum from localized pain in the flexor compartment to triggering to locking of a digit.(40) The most common abnormality is thickening at or of the A1 pulley.(40) Less common pathophysiologic abnormalities include metacarpal-phalangeal joint abnormalities, disorders at the level of the carpal tunnel, and other pulley anomalies.(40)

**Extensor Compartment Tenosynovitis (Including de Quervain’s Stenosing Tenosynovitis and Intersection Syndrome)**

De Quervain’s stenosing tenosynovitis involves hypertrophy of the extensor retinaculum of the first extensor compartment(41) involving the abductor pollicis longus and extensor pollicis brevis tendons with signs of tenosynovial and retinacular fibrosis usually present.(41, 42) The condition may be occupational when jobs require repeated forceful gripping or sustained wrist extension. However, most cases are not likely occupational. Extensor compartment tenosynovitis, including de Quervain’s tenosynovitis, is considered a comparable disorder to trigger digit with somewhat similar pathophysiology, clinical presentation, and treatment issues. De Quervain’s is the most common of the extensor compartment tenosynovoses. Intersection syndrome with a reported prevalence of 0.37% of all patients with arm or hand pain is substantially less common(43) and is somewhat controversial regarding the actual site of entrapment of the dorsal compartment(44-46) with the principle site appearing to involve the musculotendinous junction of the first extensor compartment and the tendons of the 2nd extensor compartment.(47)

Similar clinical and pathophysiological conditions are believed to affect the flexor carpi ulnaris and flexor carpi radialis tendons at the wrist. There is a strong predisposition in women and among those in their 5th through 7th decades of life.(41, 48, 49) De Quervain’s is also considered a generally non-inflammatory condition caused by hypertrophy of extensor retinaculum and parietal layer of the tenosynovium with resulting symptoms of pain on use of the affected digit. Systemic diseases are potential causes, including rheumatoid arthritis, other rheumatic disorders, diabetes mellitus, amyloidosis, heredity and anatomic variants.

**Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome)**

Ulnar nerve entrapment involves delayed conduction of the ulnar nerve combined with symptoms. It has no quality evidence of work-relatedness, but theories of work-relatedness are proposed. Guyon’s canal is the space in which the ulnar nerve accompanied by the ulnar artery traverses the wrist. It is anatomically defined as the proximal medial wall formed by the pisiform, the distal lateral wall formed by the hook of the hamate, the floor formed by the flexor retinaculum and transverse carpal ligament, and the roof formed by the pisohamate carpal ligament.(50) Within the canal, the nerve bifurcates into the superficial (sensory) branch, and the deep (motor) branch. The superficial branch leaves the canal and provides a branch to the Palmaris brevis, and then continues subcutaneously to provide sensation to the fifth digit and the ulnar half of the ring finger. The deep branch loops around the hook of the hamate, and innervates the abductor digiti quinti, flexor digiti quinti, lumbricales and interossei as it crosses the palm in a curvilinear direction.(51) This canal is dissimilar to the carpal canal in that the tendons and their tenosynovium do not accompany the nerve, thus most of the usual postulated causal mechanisms for carpal tunnel syndrome...
are not possible. However, use of the hypothenar area of the hand as a hammer is a postulated occupational mechanism. (52-56)

The location of the lesion affecting the ulnar nerve as it crosses through Guyon’s canal and the wrist is predictive of clinical symptoms, and has resulted in several classification schemes. Much of the current literature references the classification scheme proposed by Wu, which details five locations for lesions identified in collective published case reports. Lesions proximal to the bifurcation of the ulnar nerve (Type I) will exhibit mixed motor and sensory involvement. Type II lesions involve only the superficial branch; therefore, clinical presentations are purely sensory. Type III lesions occur at the outlet of the canal and involve only the deep branch, thus they are purely motor. Type IV lesions occur involving the deep branch distal to the branch innervating the hypothenar, thus are purely motor with sparing of the hypothenar muscles. Finally, Type V lesions occur proximal to the branches innervating the first digital interosseous and abductor pollicis muscles, so that only the distal motor symptoms are involved. (51)

Guyon’s canal syndrome is relatively uncommon, occurring about 20 times less frequently than ulnar lesions at the elbow. (57) Pathological lesions resulting in ulnar entrapment at the wrist reportedly are associated with concurrent compression of the median nerve in the carpal tunnel in approximately one-third of patients; (50) although there is no quality evidence that median nerve neuropathy is similarly associated with ulnar nerve involvement.

Radial Nerve Entrapment
There are no quality studies linking radial nerve entrapment with work factors, although direct, significant trauma would be a presumptive cause. Radial nerve palsies affecting the hand and wrist usually occur at points along the course of the arm and forearm, well proximal to the wrist. (58-60) Upper arm lesions are generally associated with concurrent compression of the median nerve in the carpal tunnel or subsequent callous formation. Radial Tunnel Syndrome, or posterior interosseous nerve entrapment, occurs in the proximal forearm (see Elbow Disorders Guideline). Wartenberg’s Syndrome, or radial sensory nerve entrapment in the distal forearm, is uncommon. (60)

Compression of the radial sensory nerve has been attributed to wearing a tight wrist or forearm band, anomalous brachioradialis tendon, (61, 62) repeated wrist flexion and ulnar deviation, external compression and trauma, (60, 63, 64) or from mass or bony lesions. (65) Case studies have also hypothesized an association with de Quervain’s tenosynovitis, which occurs in roughly 50% of cases diagnosed with Wartenberg Syndrome. (66)

Non-Specific Hand/Wrist/Forearm Pain
Non-specific pain is thought to be common in initial presentations in primary occupational health clinical settings, although work-relatedness is naturally unclear for condition that is not well defined. (67) The initial step is a careful history and physical examination, particularly to attempt to ascertain a specific musculoskeletal disorder.

Scaphoid Fracture
Scaphoid fractures, also known as wrist navicular fractures, are among the most common fractures of the carpal bones, (68) occurring most commonly in young males. Most are not occupational, but some clearly are work-related. The scaphoid is located at the base of the thenar eminence (thumb side), just distal to the volar wrist crease, and acts to transfer the compression loads between the hand and forearm. It also maintains normal wrist motion, carpal stability and function of the wrist flexor and extensor tendons. (69) The primary mechanism of scaphoid injury is a fall on the outstretched hand, or from axial loading with a closed fist such as grasping a steering wheel in an auto accident. (70) Scaphoid fractures are prone to non-union and avascular necrosis, particularly those involving the proximal third of the navicular, and especially if displaced. Healing problems in the proximal third have been attributed to limited blood supply that is disrupted by the fracture plane. (71) A history of fracture, as well as non-union both increase risk for development of osteoarthrosis.
Distal Phalanx Fractures and Subungual Hematoma
Fingertip or distal phalangeal fractures are frequently cited as the most common fractures of the hand, with the tuft being the most common.(22) Fractures may or may not be occupational. There are no recent quality incidence data available for tuft fractures, but estimates are between 15 to 30% of all hand fractures are tuft fractures. Tuft fractures are most often usually due to a crush injury of the fingertip, resulting in comminuted or transverse fractures and are a common occupational injury. Often, they are accompanied with nail bed laceration and subungual hematoma.(75, 76) Tuft fractures are generally stable and heal uneventfully because of the soft tissue support of the fibrous septae and nail plate.(77, 78) Crush fractures or avulsion fractures involving the proximal base of the distal phalanx however may also involve flexor or extensor tendons and may require surgical intervention.(78)

Mallet fracture or mallet finger is a common fracture-dislocation injury of the distal phalanx involving loss of continuity of the extensor tendon over the distal interphalangeal joint. This common hand injury results in a flexion deformity of the distal finger joint and may lead to an imbalance between flexion and extension forces more proximally in the digit.

Middle and Proximal Phalangeal and Metacarpal Fractures
Fractures of the proximal and middle phalanges represent approximately 46% of fractures of the hand and wrist.(22, 79) The more severe fractures are among the most challenging injuries that hand surgeons and therapists treat.(80) Fortunately, most are uncomplicated and are non-surgical cases.(81-83) Fractures may or may not be occupational. Physicians who encounter hand fractures must be able to properly diagnose and manage these hand fractures, as improper management may result in permanent impairment and disability from bone shortening, permanent angulation, joint and finger stiffness, and loss of hand function. Proximal phalangeal fractures particularly have a significant potential for hand impairment particularly if suboptimally managed because of the importance of this bone in longitudinal transfer of axial forces between the carpal and distal phalangeal joints, and the PIP joint for digit mobility. Decisions for surgical intervention should be offered upon careful consideration balancing risk of superior radiographic reduction with higher risk of debilitating stiffness from the post-operative rehabilitative state, with confidence that non-operative therapy can be improved upon.(84)

Metacarpal fractures comprise roughly 1/3 of hand fractures, with fifth metacarpal neck fractures (sometimes called “Boxer’s fracture”) accounting for 1/3 to 1/2 of these injuries,(79, 85) and fractures of the thumb constituting another 25%.(86) They occur most commonly from a direct blow to the bone causing transverse shaft fracture or through an axial loading blow such as striking an object with a closed fist. Isolated fractures of the third and fourth metacarpals are uncommon and usually involve one or more the neighboring metacarpals.

Fracture type and displacement can be partially predicted by the underlying anatomic structures of the affected digit. Fractures of the proximal phalanx, which has no tendinous attachments, typically result in volar angulation. In contrast, the middle phalanx has insertions of the flexor digitorum superficialis along the volar surface, such that fractures at the base and shaft usually have a dorsal angulation because of the action of the flexor tendons, whereas fractures of the distal neck will usually have a volar angulation as the flexors act to pull the distal fragment.(87) Fifth metacarpal fractures usually displace at a volar angle because of the action of the interosseous muscles.(88) Other metacarpal fractures tend to angulate dorsally owing to the unbalanced pull of the interosseous muscles and extrinsic finger flexors on the distal fragment.(89)

Distal Forearm Fractures
Fractures of the distal forearm make up a significant proportion of injuries and fractures treated in the emergency room,(90) although no quality data regarding incidence or injury type in the workplace were found. Fractures may or may not be occupational, but most distal forearm fractures are not work-related. There are several types of distal forearm fractures in adults, the most common being Colles’ fracture, named after the surgeon and anatomist Dr. Abraham Colles who described it in 1814.(91, 92) Because it is the most common, the eponym Colles’ fracture is often mistakenly used as a generic reference term for
all forearm or wrist fractures in adult populations. However, Colles’ fracture specifically refers to a transverse fracture of the distal radial metaphysis, with or without extension into and disruption of the radiocarpal or radioulnar articular surfaces. The distinguishing feature for Colles’ fracture is that fracture fragments are displaced or angulated dorsally on a lateral view x-ray. Other adult distal radial fractures include displaced fracture fragments that have an anterior angulation and displaced fracture fragments that are displaced palmarly and may have an anterior angulation. A fracture of the distal radius with carpal displacement can be dorsal or palmar displaced, the latter being more common. That type of fracture is caused by a fall on an extended and pronated wrist increasing carpal compression force on the dorsal rim. Some fractures are limited to the radial styloid and some are frequently associated with fracture of the ulnar styloid,(91, 93, 94) as well as a high incidence of triangular fibrocartilage complex (TFCC) disruption.(95) In one report of 118 distal fracture cases, TFCC tears occurred in 53% of extra-articular distal radius fractures and 35% of intra-articular fractures.(96) Failure to recognize a torn TFCC may result in inadequate immobilization or surgical repair, resulting in distal radioulnar joint instability. Despite the severity of these injuries, with proper diagnosis and management most patients will have a satisfactory outcome.(97)

Distal radial fractures are the result of traumatic forces, most commonly related to falling on the outstretched hand. The typical mechanism for Colles’ fracture is breaking the fall with the hand outstretched and wrist in dorsiflexion, although a minority occur due to an impact on the dorsal aspect of the hand while the wrist is flexed (jam injury into the dorsum of hand) or a direct blow to the radial styloid.(91, 93, 94) In modern times, this injury more often results from a fall with the hand in ulnar deviation and midway between pronation and supination, or as a result of other force that is transferred axially from the scaphoid into the radial facet. Distal radial fractures are up to 6 times more common in women, with incidence of 7.3 vs. 1.7 per 1,000 human years.(98) In addition, people who walk regularly, which increases exposure to falling, have an associated higher incidence of fractures. Osteoporosis and low bone-mineral density is also an associated risk factor for fracture,(99) and likely explains much of higher incidence of fracture in women. A large population longitudinal study of osteoporosis has shown elderly women with high risk factors such as diabetes mellitus, cognitive impairment, and history of falls are at high risk for fall and distal forearm fracture, and should be considered for selective preventive strategies to reduce the incidence of these fractures.(100)

The triangular fibrocartilage complex (TFCC) plays a somewhat analogous role in the wrist to the meniscus and collateral ligaments in the knee. It is formed by a network of ligaments and articular cartilage originating on the medial border of the distal radius with insertion into the base of the ulnar styloid, and includes a meniscus at the distal radioulnar joint (DRUJ). The TFCC plays an important role in load bearing across the wrist as well as in DRUJ stabilization,(101) and in allowing for pronation and supination of the hand.(102)

Fracture Classification and Diagnostic Criteria
Several classification systems for distal radial fractures have been developed in an effort to evaluate treatment outcomes. However, studies of interobserver reliability and intraobserver reproducibility for the better-known classification systems (such as Frykman, Melone, Mayo, AO, and Fernandez) have demonstrated unsatisfactory reliability and reproducibility.(103, 104) Therefore, the key to recommending a treatment course is to determine the following criteria: is a fracture open or closed, stable or unstable, or likely to become unstable. Treatment can then be provided based on these criteria until better information is available to compare outcomes using a classification system that is reliable and reproducible.

In describing or in considering a specific treatment course of distal radial fractures, it may be more useful to determine the stability of fracture patterns according to radiological measurements rather than a specific classification system.(105) Stable fractures are most often defined as dorsal angulation of less than 10°, radial shortening of 2mm maximum, and no radial shift.(106) Fractures outside of these limits generally require reduction, with larger angulation, articulation step-offs, dorsal comminution, and lateral shift of more than 2mm considered contributory factors for fracture instability and indication for more aggressive therapies.(107, 108) Despite the importance placed on these criteria, not surprisingly there are conflicting
opinions regarding the reliability of these measurements, which provides reduced confidence that these
are absolute criteria, and leads to some uncertainty regarding measurements used in comparison studies
as well as in general practice.(109-112)

Ganglion Cyst
Ganglion cysts occur in nearly any joint of the hand and wrist and have an estimated prevalence rate of
14%.(113) although prevalence rates based on MRIs are approximately 50%, with asymptomatic ganglia
more likely to be volar (palmar) than dorsal.(114) Symptomatic onset is a common work-related claim, but
quality studies linking ganglia with work continue to be lacking. Wrist ganglia account for 50 to 70% of all
wrist masses identified.(115) Other causes include giant cell tumors also known as localized nodular
tenosynovitis and fibrous xanthoma, epidermal inclusion cysts and fibromas. Wrist ganglia are generally
classified as either dorsal or palmar, with dorsal ganglia comprising up to 80% and volar ganglia making up
approximately 20% of clinically detected ganglia.(116) Approximately 10% of all hand and wrist ganglia are
found on a flexor tendon sheath of the fingers.(117)

A ganglion is a cystic structure, although is not technically a cyst as it has no synovial lining.(118) Electron
microscopy shows the walls to be composed of randomly oriented collagen fibers. The gelatinous cystic
fluid is likened to synovial fluid, although the composition of hyaluronic acid, glucosamine, globulins, and
albumin is not the same.(118)

The pathogenesis of ganglia is unknown and the epidemiology sparse. Contributing factors are also
unknown. There are several theories of origin, although each has significant weaknesses and none have
been proven. These include the cyst being formed: 1) as a simple herniation of the joint capsule; 2) as a
result of an inflammatory process from overuse; 3) as a tear in the joint capsule with subsequent release
of synovial fluid and subsequent reaction to the mucinous fluid; 4) as a result of mucoid degeneration of
adjacent extra-articular connective tissue; and 5) from joint stress causing mucin secretion by
mesenchymal cells in surrounding tissue.(118-121) Each of these theories fails to wholly explain all of the
known facts, particularly because there seems to be no inflammatory process.

Most wrist ganglia are asymptomatic. Many patient visits are primarily for aesthetic reasons. A cross
sectional study of asymptomatic volunteers who underwent wrist MRI revealed a ganglion prevalence rate
of 51%. (114) A symptomatic ganglia were more likely to be volar (palmar) than dorsal.(114)

Hand Arm Vibration Syndrome
The term “hand arm vibration syndrome (HAVS)” has been used since the 1980s to describe the
constellation of adverse physiological responses causally associated with high-amplitude vibratory forces,
such as those experienced through the use of various hand tools including pneumatic drills, riveters and
chain saws(122-124) or from vibratory rich activities such as driving off-road vehicles.(125) Other terms
commonly used to describe these responses include Raynaud’s phenomenon of occupational origin, white
fingers, dead fingers, traumatic vasospastic disease (TVD), and “vibration-induced white finger.”(126)

The adverse effects of HAVS are characterized by circulatory disturbances associated with digital arteriole
sclerosis and manifest as vasospasm with local finger blanching; sensory and motor disturbances manifest
as numbness, loss of finger coordination and dexterity, clumsiness and inability to perform intricate tasks;
and musculoskeletal disturbances manifest as swelling of the fingers, bone cysts and vacuoles.(127, 128)
There are also several reports of association of CTS with HAVS and exposure to vibration.(127, 129-132)

Epidemiologic evidence indicates there is a latency period of from 1 to 16 years of exposure before onset
of HAVS, with a trend for decreasing prevalence as changes in work-practice and anti-vibratory tools and
dampening actions have been implemented.(133) The direct pathophysiological basis for the observed
vascular responses of HAVS is not known, but several theories are proposed including vibration causing
direct trauma to smooth muscle and smooth muscle vacuoles,(134) vascular spasm related to activation of
alpha-2 adrenoreceptor in the vessel walls,(135) or the release of a potent vasoconstrictor known as
salivary endothelin.(136) The pathophysiology of neurologic deficits is also unknown, but presumably is
related to vibration induced microvascular changes and demylelination.(133)
The pathophysiologic changes related to vibration are initially reversible, but with increasing duration and intensity of exposure, the disorder may continue to progress or become permanent. According to the International Organization for Standardization, the risk for developing HAVS is proportional to the total vibration energy measured in magnitude, duration, and frequency. The range of vibration frequencies thought to be harmful is 4Hz to 5000Hz dependent on the intensity, and whether or not it is oscillatory or impact force, with impact vibratory force thought to be more hazardous. There are several exposure limit guidelines for vibration, including for the United States. There are other guidelines adopted in the UK and Japan. There is limited epidemiological data to better define the exposure-response relationships for each of the various components of HAVS, however, recent animal models provide some insights into exposure-response patterns.

Laceration Management
Traumatic injuries resulting in skin lacerations of the upper extremity are a common reason for patient visits to an urgent care, occupational medicine clinic or emergency department. Lacerations result from blunt or crush injuries that produce shear forces, or more commonly from sharp objects which are abundant in the workplace. The majority of lacerations can be treated on an outpatient basis. The primary purpose of wound and laceration management is to avoid infection, detect if a nerve injury has occurred, manage tendon lacerations, and achieve a cosmetically acceptable result with the highest degree of function and patient satisfaction. The most optimal results are accomplished by preventing infection through thorough wound cleansing, approximating wound edges with appropriate closure techniques, and providing a proper dressing with a clean moist environment to accelerate wound healing.

Human and Animal Bites and Associated Lacerations
There are no recently reported rates of human and animal bites in the United States. However, extrapolation of emergency department visits and other epidemiological studies from the 1990s indicate there are an estimated 5.0 million dog bites annually, with roughly 750,000 to 800,000 of those bites of significant severity to require medical treatment. Data on cat bites are more limited, but they are the second most common animal bite, with an estimated 66,000 emergency room visits followed closely by human bites.

Although most bites occur from animals known to the victim, occupations that may be at higher risk for animal bites include veterinarians, animal handlers, police officers, utility services personnel who access private property, mail carriers, and other similar professions. Human bites are common in caregivers, educators, law enforcement officers, and in instances of accident or workplace violence that may involve the fist or hand being cut by contact with teeth.

Other than deep destruction of tissue requiring reconstruction, risk of infection is the primary concern for animal bites. There also are other zoonotic diseases such as rabies, cat scratch fever, and human blood borne pathogens exposures that should also be considered. The reported incidence of infection from non-complicated bite wounds from dogs is between 3 and 10%, from cats is 20 to 50%, and from humans is up to 50%. Rates may be higher for wounds of the hand, depth of penetration into the skin, and length of time before wound is irrigated and cleaned. For purposes of this guideline, discussion and recommendations are made based on bites to the extremities or trunk as well. Facial injuries are not considered in this guideline and there may be somewhat different indications as the significance of complications is generally more severe.

Hand/Finger Osteoarthrosis
Hand and finger osteoarthrosis is extraordinarily common affecting over 50% of the aged population. These are believed to be largely non-occupational issues but some may be covered under some workers' compensation jurisdictions, usually under fairly limited circumstances. This is particularly true for mono-articular arthrosis as a consequence of an occupational injury.
Initial Assessment

Thorough medical and work histories as well as focused physical examinations (see General Approach to Initial Assessment and Documentation Guideline) are sufficient for the initial assessment of the majority of patients with a potentially work-related hand, wrist, or forearm symptom(s). These evaluations should consider assessments of red flags, including the possibility of referred pain to the hand, wrist, or forearm from a disorder in another part of the body (e.g., cervical nerve root or heart). The absence of red flags largely rules out the need for special studies, surgical intervention, or inpatient care for the first 4 weeks, as during this time, spontaneous recovery is common (provided any inciting workplace or other factors are addressed).

Hand, wrist, and forearm symptoms can be classified into one of four working categories:

- **Potentially serious hand, wrist, or forearm condition**: fracture, acute dislocation, infection, neurovascular compromise, or tumor.
- **Mechanical disorders**: derangements of the hand, wrist, or forearm related to acute trauma, such as ligament sprain or muscle-tendon unit strain.
- **Degenerative disorders**: resulting from aging or symptoms associated with use, or a combination thereof, such as osteoarthrosis, other arthritides, tendinosis, or tenosynovitis.
- **Nonspecific disorders**: occurring in the hand or wrist without clear, specific pathophysiological correlates (most typically includes non-specific pain and sometimes erroneously called “forearm tendinitis”).

Carpal Tunnel Syndrome

CTS patients typically have a constellation of symptoms with some variation in clinical presentations(8, 167, 168) and a lack of a criterion standard.(10) Symptoms most typically start gradually in the thumb, index, and middle fingers with tingling, numbness, or burning.(11, 167) Symptoms may also include subjective hand swelling.(169) Symptoms often first appear during sleep, possibly due to sleeping with wrists flexed, edema, venous pooling or a combination of factors. The patient may awaken with the desire to “shake out” the hand or wrist. As symptoms worsen, patients may experience tingling during the day particularly after a sustained hand grasp (such as when using a steering wheel or hand tool). Wrist flexors are innervated by the median nerve proximal to the wrist. However, decreased thumb and grip strength sometimes occurs and may make it difficult to form a firm fist, sustain grasp particularly of small objects, or perform other manual tasks. In chronic, advanced, and/or untreated cases, the muscles of the thenar eminence may atrophy. Some severely affected patients are unable to differentiate between hot and cold. Symptoms are most commonly documented through detailed recording of symptoms and digits affected or with a hand symptom diagram.(168, 170-174) With the Katz hand diagram, the patient is provided with a form that shows outlines of the arms, and the palmar and dorsal surfaces of the hands. The patient identifies areas of discomfort on the diagrams and characterizes the symptoms (e.g., pain, numbness, tingling, or other). The results are scored by the clinician as “probable,” “possible,” or “unlikely,” depending upon specified criteria.

Patients with CTS should have paresthesias (tingling and/or numbness)(10, 167, 168) but pain in the wrist hand or fingers may or may not be present. In patients with only wrist or hand pain without paresthesias, a diagnosis other than carpal tunnel syndrome may be present. Symptoms of tingling, numbness and pain in the median nerve distribution of the hand are common in the general population (prevalence approximately 14 to 37%). However, based on clinical examination and electrophysiologic testing, CTS prevalence rates range from 2.7% to as high as 13.0%(175-187) while the incidence rate for working populations is near 2.3 per 100 person-years.(188) Differences in diagnostic criteria and population characteristics between these studies may play a role in the differences in reported CTS prevalence.(189)

Triangular Fibrocartilage Complex (TFCC) Tears

TFCC tears appear to occur either with acute discrete traumatic events and/or as degenerative cartilaginous changes. A primary focus of the patient history is ascertaining whether the TFCC is significantly torn, and if so, whether it is sufficiently symptomatic to require intervention(s). Following the
patient’s symptoms for healing without immediate surgical intervention is generally the most common approach. Some do not heal, continue to be symptomatic and do well with surgical repair or removal.

**Crush Injuries and Compartment Syndrome**
Patients with more severe injuries present with severe pain and may have vascular compromise. The initial assessment should focus on the degree of injury severity and if the injury requires surgical evaluation and treatment. Milder injuries may be managed non-operatively; however, the threshold for surgical consultation should be low. Those with milder injuries should be monitored for neurovascular compromise.

**Kienböck Disease**
Patient typically presents with progressive pain and disability and have characteristic wrist x-rays demonstrating changes in the lunate.

**Wrist Sprains**
Wrist sprains typically occur with acute traumatic events. The diagnosis is sometimes applied as a diagnosis of exclusion among patients with pain in the setting of trauma with negative fractures. However, the specific entity is properly defined as a partial ligamentous disruption rather than undefined pain generators. Sprains may also occur as an accompaniment to fracture.

**Mallet Finger**
Mallet finger is readily diagnosed based on the presentation of inability to extend the distal interphalangeal joint, generally in the context of trauma or distal interphalangeal joint arthrosis.(21)

**Flexor Tendon Entrapment (Tenosynovitis and Trigger Digit)**
Flexor tendon entrapment generally presents as a relatively simple clinical presentation. Some occur after acute injury, but most occur without specific inciting event.(40, 190-193)

**Extensor Compartment Tenosynovitis (Including de Quervain’s Stenosing Tenosynovitis and Intersection Syndrome)**
Extensor tendon entrapment generally presents as a relatively simple clinical presentation. Some occur after acute injury, but most occur without specific inciting event.

**Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome)**
Ulnar nerve entrapment at the wrist generally presents as numbness and/or tingling in the fourth and fifth digits. Certain patients may also experience a weakened grip or difficulty with finger coordination.

**Radial Nerve Entrapment**
Radial nerve entrapment usually presents as radial nerve palsies affecting the hand and wrist, most commonly occurring at points along the course of the arm and forearm, well proximal to the wrist.(58-60) The medical history should include a search for sensory symptoms. Symptoms may also include pain over the course of the nerve.

**Non-Specific Hand/Wrist/Forearm Pain**
Non-specific hand/wrist/forearm pain typically occurs in the absence of discrete trauma. Instead, it frequently occurs in settings of high physical job demands or ill-defined exposures. This is a “diagnostic” category to be utilized when symptoms are present, but in the absence of an identified, specific disorder. Most cases will resolve without significant difficulty. If there is no improvement after several weeks of treatment, focused diagnostic testing should be considered. Non-specific pain lasting more than 2 months is fairly rare. The search for a specific diagnosis should include proximal pathology including spine-related (e.g., radiculopathy, spinal tumor, infection) as well as psychological disorders particularly when widespread symptoms are elicited or a pattern or recurrent unexplained illnesses is present (see Chronic Pain Guideline).
Scaphoid Fracture
The main initial tasks are to confirm a fracture, identify those patients with fractures best treated with surgery, and treat those with a high clinical suspicion of fracture with appropriate splinting. A history of sufficient injury potential is important. Patients frequently complain of persistent swelling and tenderness near the thumb base in the area of the scaphoid. Gripping and wrist motion may be painful.

Distal Phalanx Fractures and Subungual Hematoma
Patients have swelling, reduced range of motion, and tenderness of the fingertip. Patients with accompanying subungual hematoma may have severe throbbing pain and obvious discoloration of the affected nail.

Middle and Proximal Phalangeal and Metacarpal Fractures
The initial assessment involves a search for confirmation of fracture. Limited or guarded range of motion with pain, local tenderness, swelling, deformity and possibly ecchymosis over the affected area are common.

Distal Forearm Fractures
Wrist injuries associated with significant pain, swelling, ecchymosis, crepitance, or deformity should be considered to be fractured until proven otherwise. Forearm fractures may also result in concomitant vascular, neurological, ligament and tendon injuries. Further, as distal forearm fractures are the result of trauma, careful inspection for other traumatic injuries should be included, such as elbow, shoulder, neck, head, and hip. In general, most distal forearm fractures should be managed by an orthopedic or hand surgeon and consultation is recommended.

Ganglion Cyst
Most ganglia present as a bump or mass. Occasionally patients with noticeable ganglia will complain of mild nuisance pain, and less often of severe pain. In the assessment of wrist pain in the absence of palpable ganglia, the unexplained wrist pain may be a result of occult ganglia and should be included in the differential diagnosis. The pain from an occult dorsal lesion has been linked to the compression of the posterior interosseous nerve.(194) Ganglia have also resulted in compression of the median and ulnar nerves as they pass through the carpal tunnel and condylar groove respectively (see section on Ulnar Nerve Entrapment and Elbow Disorders Guideline).

Hand Arm Vibration Syndrome
Initial assessment for HAVS is a detailed history and examination focusing particularly on high-amplitude vibratory exposure and sensorineural or vascular symptoms. The clinical symptoms may include episodic tingling, numbness, blanching white fingers, pain and paresthesia, burning sensation, clumsiness, poor coordination, sleep disturbance, hand weakness measured in grip strength, and diffuse muscle, bone and joint pain from the fingers to the elbow.(42) Differential diagnosis should consider other causes of Raynaud’s phenomenon, including the connective tissue diseases of scleroderma, systemic lupus erythematosus, rheumatoid arthritis, dermatomyositis, and polyarteritis nodosa.

Laceration Management
A thorough history of the injury, with particular attention to mechanism, potential degree of wound contamination, potential for foreign bodies, and presence of other trauma should be obtained. Crush wounds may be more susceptible to infection, and contamination. Additionally, inquiry of personal factors that may contribute to delayed healing or increased risk for infection, such as diabetes mellitus, chronic renal failure, or the use of immunosuppressive medications should be included.(195) Tetanus immunization status should be noted and are recommended to be updated per CDC guidelines (see Table 1).
Table 1. Guide to Tetanus Prophylaxis in Routine Wound Management

<table>
<thead>
<tr>
<th>History of adsorbed tetanus toxoid (doses)</th>
<th>Clean minor wounds Tdap or Td†</th>
<th>Clean minor wounds TIG§</th>
<th>All other wounds’ Tdap or Td†</th>
<th>All other wounds’ TIG§</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than 3 or unknown</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>3 or more doses†</td>
<td>No**</td>
<td>No</td>
<td>No††</td>
<td>No</td>
</tr>
</tbody>
</table>

* Such as (but not limited to) wounds contaminated with dirt, feces, soil, and saliva; puncture wounds; avulsions; and wounds resulting from missiles, crushing, burns, and frostbite.
† For children younger than 7 years of age, DTaP is recommended; if pertussis vaccine is contraindicated, DT is given. For persons 7-9 years of age, Td is recommended. For persons >10 years, Tdap is preferred to Td if the patient has never received Tdap and has no contraindication to pertussis vaccine. For persons 7 years of age or older, if Tdap is not available or not indicated because of age, Td is preferred to TT.
§ TIG is human tetanus immune globulin. Equine tetanus antitoxin should be used when TIG is not available.
¶ If only three doses of fluid toxoid have been received, a fourth dose of toxoid, preferably an adsorbed toxoid, should be given. Although licensed, fluid tetanus toxoid is rarely used.
** Yes, if it has been 10 years or longer since the last dose.
†† Yes, if it has been 5 years or longer since the last dose. More frequent boosters are not needed and can accentuate side effects.


Human Bites, Animal Bites and Associated Lacerations
A careful history for time and location of the bite should be obtained as it will help guide clinical decisions regarding prophylaxis. If possible, information about the type of animal and its health status as well as the circumstances related to why the bite occurred should be obtained. Tetanus and rabies immunization status should be established.

Hand/Finger Osteoarthrosis
Most cases of osteoarthrosis are believed to result from genetic factors, although discrete trauma is a potential cause. The initial assessment is usually relatively concise and generally involves securing a diagnosis and initiating treatment.

Medical History
Asking the patient open-ended questions allows gauging of the need for further discussion or make specific inquiries to obtain more detailed information. Hand dominance should be noted. Consider initiating the clinical visit with an open-ended question such as “What can I do for you?” to assure that the chief complaint is addressed. More specific questions for hand, wrist, and forearm conditions include:

Symptoms:
What symptoms are you having? For how long?
- Do you have pain, numbness, tingling, weakness, or limited movement?
- For traumatic injuries: Did the area swell? If swollen, how quickly did it swell (immediately or delayed)?
- Was the hand/finger deformed?
- Are your symptoms located primarily in the hand, wrist, or forearm? Do you have pain or other symptoms in the elbow, shoulder, or neck? Anywhere else?
- Are your symptoms constant or intermittent?
- What causes your symptoms to increase?
- What time of the day are your symptoms best? Worst? On getting out of bed? Morning? Mid-day? At work? Evening? While sleeping?
- If symptoms awaken you, how often a week? Each night?
- What makes the symptoms better or worse?
- Have your symptoms changed? How have they changed?
- Can you quantify your pain on a scale of 1 to 10 (10 being unbearable or worse possible pain). It is important to quantify and track the patient’s response to evaluate the effectiveness of treatment.
- What have you done to reduce your symptoms?
Onset (Occupational and Avocational):
How did your symptoms begin? Was there a single, sudden event (e.g., slip, trip, or fall) when your symptoms started or did the symptoms begin gradually?
- Are you able to do your usual job? How do these symptoms limit you?
- Do you work out and use weights/weightlifting while working out?
- Can you perform activities of daily living (e.g., dressing, bathing, grooming, etc.) or instrumental activities of daily living (e.g., shopping, food preparation, housekeeping, etc.)?
- What stops you from doing activities? Are the symptoms worse with workplace activities?
- Can you grasp? How much? Are you dropping things?
- What is your job? What are your specific job activities? Do you use your hand, wrist, or forearm to perform them? What are the most forceful hand activities? How? How often?
- Are there differences in exposures between hands (are symptoms not dissimilar or vice versa)?
- (For discrete trauma): Exactly how did you injure the hand/ﬁnger? (Record in detail)
- (For non-discrete trauma): What do you think caused these symptoms? (Record in detail) Proceed with other questions, but return to record details of maximum and typical force, repetition, posture, vibration as appropriate after securing a provisional diagnosis.
- Have the symptoms limited your activities? For how long?
- What are your hobbies? How often?
- Do you use vibrating tools or devices at work or at home (especially high amplitude, low frequency such as older model chain saws)? Do you ride a motorcycle or four wheeler? Do these activities seem to affect your symptoms?

Current Treatments Used:
What have you used to treat the current symptoms?
- Has any treatment helped? Or, not helped?

Prior Injuries and Prior Treatments:
- Have you had this problem or similar symptoms previously with this hand? The other hand?
- What makes it better and what makes it worse? Do you have symptoms at night? On weekends? On vacations?
- Have you had previous testing or treatment? Which? What were the results? What seemed to work best in the past?
- What do you think caused your symptoms? Do you think your symptoms are related to work?

Other Relevant Disorders:
- Do you have other medical conditions? (For example, overweight/obesity, diabetes mellitus, hypothyroidism, other endocrinopathy, pregnancy, osteoarthritis, rheumatoid arthritis, other arthritides, renal disease etc.)

Carpal Tunnel Syndrome
There are numerous purported risk factors for CTS (see Table 2), although many have not been conﬁrmed in prospective studies as true independent risk factors. Evidence appears most consistent in the retrospective studies for age, obesity, female gender, diabetes mellitus, and combinations of forceful and repetitive grasping.(6, 12, 196-210, 211, 212) Recent prospective cohort studies of CTS have conﬁrmed the above ﬁve factors as apparently true risk factors, including repeated high force grasping, overweight or obesity, female gender, and psychosocial factors.(146, 213-219)
Table 2: Possible Risk Factors for Carpal Tunnel Syndrome

This list is based on prospective, cross-sectional, and case-control studies, case series, and case reports. Note, this table is not meant to be all inclusive.

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>Possible Risk Factors</th>
</tr>
</thead>
</table>
| Trauma      | Any past or recent fracture of the wrist  
Carpal-metacarpal dislocation  
Casting following a fracture  
Crush injury  
Repeated contusions to the wrist  
Volkmann’s ischemic contracture |
| Developmental or Genetic Causes (Heredity) | Female gender, pregnancy, menopause  
Age >40  
Persistent median artery  
Enlarged lumbrical or/flexor digitorum superficialis muscle(s)  
Smaller cross sectional carpal tunnel area – females particularly have smaller wrists  
Squarer wrists – wrist depth to width ratio of more than or equal to 0.70  
Primary familial carpal tunnel syndrome due to thickening of the transverse carpal ligament – thus runs in families  
Hereditary neuropathic pressure palsies |
| Swelling and Masses | Ulnar bursitis  
Ganglion cysts  
Lipoma or fatty tumor/other tumors  
Overweight or obesity – usually measured with Body Mass Index – weight (kg)/height (m²)  
Acromegaly with oversized bones and soft tissues in the wrist  
Hypertrophic polyneuropathy with median nerve enlargement  
Proximal lesion of the median nerve (double crush syndrome) |
| Rheumatological Disorders, including Inflammatory and Non-Inflammatory Arthropathies | Nonspecific tenosynovitis with synovial swelling and thickening  
Osteoarthritis  
Rheumatoid arthritis  
Scleroderma  
Chondrocalcinosis  
Dermatomyositis  
Amyloidosis with amyloid deposits  
Multiple Myeloma  
Paget’s disease  
Gout, as well as other crystal arthropathies |
| Other Inflammatory and Infectious Conditions | Histoplasmosis  
Sporotrichosis  
Coccidiomycosis  
Rubella  
Leprosy with enlargement of the median nerve  
Hepatic disease  
Fibromyalgia  
Polymyalgia rheumatica  
Raynaud’s disease  
Infections of the wrist joint or other compartments  
Lyme disease  
Tuberculosis |
| Metabolic, Nutritional, and Alterations in Fluid Balance | Diabetes Mellitus  
Alcoholism  
Vitamin Bs deficiency  
Pregnancy – presumably due to increased body fluid and swelling  
Menopause with hormonal imbalance  
Eclampsia of pregnancy  
Hypothyroidism – particularly with fluid retention, although other history of thyroid disorders appears to be a risk  
Renal disease and renal failure – especially with fistulae for hemodialysis  
Oral contraceptive and estrogen use  
Glucocorticosteroid use |
<table>
<thead>
<tr>
<th>Activities and Avocations</th>
<th>Vocational Activities</th>
</tr>
</thead>
<tbody>
<tr>
<td>Musical instrument use (e.g., violin, piano)</td>
<td>Combinations of high force and high repetition especially meat and shellfish processing and some manufacturing positions. Some grocery scanning positions may also be at risk, particularly if handling high volumes of heavy product)</td>
</tr>
<tr>
<td>Prolonged driving</td>
<td>High force grasping on a stereotypical basis</td>
</tr>
<tr>
<td>Prolonged writing</td>
<td>Highly repeated grasping</td>
</tr>
<tr>
<td>Bowling</td>
<td></td>
</tr>
<tr>
<td>Motorcycle riding (e.g., vibration and handle bar grasp)</td>
<td></td>
</tr>
<tr>
<td>Snowmobiling</td>
<td></td>
</tr>
<tr>
<td>Sewing, knitting and crocheting</td>
<td></td>
</tr>
<tr>
<td>Bicycling</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Triangular Fibrocartilage Complex (TFCC) Tears**

Patients commonly complain of non-radiating ulnar sided pain and clicking. It is important to correlate the symptoms with the physical examination and mechanism of injury since MRI studies suggest TFCC tears are both prevalent while also apparently frequently asymptomatic.\(^{(14, 220, 221)}\) Ulnar deviation with axial loading tends to increase pain. A “click” or “clunk” in the ulnar wrist joint may be reproduced with forearm rotation (supination/pronation). Commonly reported mechanisms of injury include a fall on an outstretched hand(222-224) as well as sports.\(^{(225, 226)}\) Thus, some work-related accidents are reported causes of these tears. Those with occupational cases will tend toward symptomatic onset after a discrete traumatic event such as a slip and fall.

**Crush Injuries and Compartment Syndrome**

Patients have pain, and may have paresthesias. Those with vascular compromise may have a cool extremity compared with the unaffected limb. Crush injuries have clear mechanisms of injury on history. However, there are many causes of compartment syndrome including trauma, excessive traction from fractures, tight casts, bleeding disorders, burns, snakebites, intraarterial injections, infusions, and high-pressure injection injuries.\(^{(19, 227-231)}\)

**Kienböck Disease**

Patient complains of increasing wrist pain, pain with movement, pain with use, and limited range of motion.

**Wrist Sprains**

Patients invariably have incurred an acute traumatic event, usually a slip, trip, or fall with forceful loading of the wrist joint usually in a fully deviated position (e.g., full extension). They have pain in the wrist joint, and generally have no swelling.

**Mallet Finger**

The mechanism of injury most typically involves forcefully striking the tip of the extended digit on an object (e.g., balls caught by the hands in sports), as well as from falls.\(^{(22)}\) Unless there is a fracture, most cases present without significant, post-traumatic pain. Some occur without any trauma and are thought to mostly occur with osteoarthrosis and Heberden’s nodes or other chronic joint pathology.

**Flexor Tendon Entrapment (Tenosynovitis and Trigger Digit)**

Epidemiological evidence is weak, thus lines of query are unclear and causal conclusions tenuous.\(^{(40, 232)}\) The mechanism of injury for many appears to be typically idiopathic\(^{(40, 190, 192, 233)}\) or as a complication of medical conditions (especially diabetes mellitus and rheumatoid arthritis).\(^{(33)}\) However, available epidemiological and biomechanical evidence suggests that the disorder may also occur as a complication of repeated forceful use of a digit,\(^{(35, 40, 190, 192, 234-244)}\) or unaccustomed use,\(^{(35, 40)}\) thus many cases may be work-related.\(^{(29, 40, 245)}\) A careful history of occupational tasks as well as non-occupational exposures is recommended. Symptoms are variable and may include pain, stiffness, clicking, snapping, and locking.\(^{(28, 35, 192, 234, 244, 246-249)}\) Pain is generally in the volar digit and/or metacarpophalangeal joint area.\(^{(244, 246-248)}\) Certain patients report worse symptoms in the morning or after lack of use.\(^{(40)}\)
Extensor Compartment Tenosynovitis (Including de Quervain’s Stenosing Tenosynovitis and Intersection Syndrome)

Patients present with wrist pain that is augmented by movement and generally non-radiating,(42) although occasionally pain may spread along the course of the affected tendon sheath.(41, 42) Patients rarely have paresthesias unless there is an accompanying swelling or other mechanism to affect the superficial radial nerve or other digital nerves.(41) Some repeated hand postures with thumb pinching may be associated with de Quervain’s disease.(250) There is belief that superficial radial nerve entrapment may accompany de Quervain’s and may then produce paresthesias in the first dorsal web space. Triggering is rare.(251)

Ulnar Nerve Entrapment at the Wrist (including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome)

Ulnar nerve entrapment at Guyon’s canal typically first presents with symptoms of paresthesias followed by late symptoms of weakness. It is reportedly usually not associated with pain, in contrast with carpal tunnel syndrome that appears to more frequently involve pain. Patients with traumatic causes of ulnar neuropathy tend to have motor symptoms, whereas those with idiopathic or non-trauma related causes usually manifest sensory symptoms.(50)

Radial Nerve Entrapment

Successful localization of radial nerve entrapment can frequently be accomplished through a careful history and physical exam. The medical history should search for sensory symptoms including paresthesias with precision of the location of the paresthesias to a typical radial nerve distribution on the dorsal hand, particularly in the first dorsal web space.(59) Symptoms may also include pain over the nerve. Distinguishing from other sources of sensory symptoms is usually possible, particularly including radiculopathies and other entrapment syndromes. An assessment of motor symptoms, including wrist extensor weakness as well as wrist drop, are also helpful, particularly in conjunction with absence of weakness in other distributions.

Non-Specific Hand/Wrist/Forearm Pain

Patients most commonly give a history of gradual onset of pain or other symptoms in the absence of discrete trauma. Symptoms are most often in the forearm, and frequently are not well localized.

Scaphoid Fracture

Historical features most commonly involve a high-energy injury such as a fall on an outstretched, extended hand with immediate, non-radiating pain in the radial carpus. Other common mechanisms include grasping a steering wheel in a frontal motor vehicle crash, or direct blow to the scaphoid such as when using the heel of the wrist as a hammer.

Distal Phalanx Fractures and Subungual Hematoma

Tuft fracture should be suspected when a patient presents with a crush injury or perpendicular shearing force injury to the fingertip, particularly if there is a subungual hematoma. Injuries resulting in avulsion of the nail plate can also be associated with tuft fractures.

Middle and Proximal Phalangeal and Metacarpal Fractures

Careful history regarding the mechanism of injury including and direct axial blow or angular or rotational trauma will reflect substantially on the nature of the fracture and its inherent stability.(78)

Distal Forearm Fractures

There are several types of distal forearm fractures (see Distal Forearm Fractures in the section on General Approach and Basic Principles). There can be a high incidence of TFCC tears with these fractures.

Ganglion Cyst

Ganglion cysts are usually asymptomatic. The cause is unknown, although the condition appears associated with aging.
Hand Arm Vibration Syndrome (HAVS)
HAVS can manifest as vasospasm with local finger blanching; sensory and motor disturbances such as numbness, loss of finger coordination and dexterity, clumsiness and inability to perform intricate tasks; and musculoskeletal disturbances such as swelling of the fingers, bone cysts, and vacuoles. (128, 252)

Laceration Management
Lacerations of sufficient size, depth and those occurring over joints usually require suturing. Tetanus immunization status should be addressed.

Human Bites, Animal Bites and Associated Lacerations
A detailed medical history pertaining to tetanus and in the case of animal bites, rabies immunization status, and underlying medical conditions such as diabetes mellitus or other immune-compromising conditions is important. Most wounds are puncture wounds, but some wounds may be considered for suturing.

Hand/Finger Osteoarthrosis
Patients usually have no recalled acute traumatic event. A minority have a history of significant trauma, such as a fracture or dislocation. Regardless of cause, symptoms usually consist of gradual onset of stiffness and non-radiating pain. Gradual joint enlargement is often present, although frequently unnoticed by the patient. Swelling, erythema, warmth and other signs of infection or inflammation are not present, and if present signal an inflammatory, crystalline arthropathy, septic arthritis or other cause. The history should include symptoms affecting any other joints in the body, presence of other potential causes (e.g., psoriasis, rheumatoid arthritis, gout) to help ascertain the correct diagnosis.

Physical Examination
Guided by the medical history, the physical examination includes:
- General observation of the patient; and
- Appropriate regional examination of upper limbs (hands, wrists, forearms, elbows, arms, shoulders, and neck).

The general observation involves specification of which distal upper extremity is affected and observation of how much the affected hand or arm is used versus how much activity is avoided – e.g., does the patient shake the examiner’s hand or avoid all use of the hand or arm? Does the patient hold the arm without using it? Are there differences in use depending on whether there is active rather than casual observation and examination? These aspects of the physical examination are under-rated, yet perhaps the most important aspects for ascertainment of degrees of impairment and severity of the condition. Most components of the examination are at least in part, subjective since the patient must exert voluntary effort or state a response to a stimulus such as the sensory examination or tenderness. In many cases of hand, wrist, or forearm problems, there are no strictly objective findings. Exceptions include palpable trigger finger, ganglia, thenar atrophy, and fracture-related deformities.

The physician should seek objective evidence of pathology that is consistent with the patient’s symptoms. In some cases, careful examination will reveal one or more truly objective findings, such as swelling, deformity, atrophy, reflex changes or spasm, fasciculations, trophic changes, or ischemia. Regardless of whether completely objective findings are present, all findings should be documented in the medical record.

A. Regional Examination of Hand, Wrist, and Forearm
The inter-related hand, wrist, forearm, arm, shoulder, and neck should be examined individually and functionally together for observation of use, function, swelling, masses, redness, deformity, asymmetry, or other abnormality. The examination should extend to the proximal upper limb and neck. This examination may be followed by evaluating active and passive range of motion within the patient’s limits of comfort with the area as relaxed as possible for passive range of motion. Local tenderness may be accentuated by specific motions or stresses on specific joints, and active muscle contraction may produce pain, indicating
a specific tendinosis. If this latter finding is on the dorsoradial side of the wrist, it suggests a diagnosis of de Quervain’s tenosynovitis. Specific areas of decreased pinprick sensation may indicate median or ulnar nerve compression. Flexing the wrist for 60 seconds with elicitation of dysesthesias in the median innervated digits is considered a positive Phalen’s test.

Several purported signs of carpal tunnel syndrome (CTS) have limited specific diagnostic value and the history is believed to be of critical importance in securing a presumptive diagnosis of CTS. The various signs for CTS show a broad range of positive predictive value that is especially dependent on the patient population assessed. Physicians should primarily rely on the clinical history as well as the physical examination. The most sensitive screening methods appear to combine night discomfort, abnormal Katz hand diagram, and abnormal sensibility by monofilament Semmes-Weinstein testing comparing affected with unaffected nerve distributions. Hypalgesia in the median nerve distribution and thumb abduction strength testing also have been found to be helpful in establishing the diagnosis of CTS. The flick “sign” is another diagnostic tool. It reportedly has high sensitivity and specificity; however, it is a historical finding rather than a true physical examination sign. The historical feature is positive when a patient reports shaking his or her hand in an effort to relieve paresthesias. (253) The diagnostic utility of physical examination tests for CTS is unclear as the underlying studies supporting meta-analyses have methods that are not well described. Phalen’s maneuver is thought to be superior to Hoffmann-Tinel’s ("Tinel’s") sign over the median nerve, although neither perform particularly well. (254)

Trigger finger (tendon) nodules may be palpable with both active and passive range of motion. However, some patients only have tenderness over the flexor surface of the metacarpal phalangeal joints, which may make this examination more difficult. A ganglion may be present on either inspection, or for smaller ganglia, only on palpation. The severity of symptoms on physical examination is usually the basis for aspiration or surgical excisions.

Fractures are most commonly discovered by deformity in the context of focal pain and an inciting trauma history. Some occur without deformity and are only found on x-rays, although most have focal tenderness on a careful palpatory examination.

B. Neurovascular Screening
The neurologic and vascular status of the hand, wrist, forearm, and upper limb should include peripheral pulses, motor function, reflexes, and sensory status. Examining the neck and cervical nerve root function is also recommended for most patients. For example, a C6 radiculopathy may cause tingling in the thumb and index finger and may affect the wrist extensors while T1 radiculopathy can present as dysfunction of the intrinsic muscles of the hand.

C. Assessing Red Flags
Potentially serious conditions for the hand, wrist, and forearm are listed in Table 3. Early consultation by a hand or upper limb specialist, rheumatologist, or other relevant specialist is recommended depending on the provider’s training and experience in dealing with the particular disorder.

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Medical History</th>
<th>Physical Examination</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fracture</td>
<td>History of significant trauma</td>
<td>Significant swelling</td>
</tr>
<tr>
<td></td>
<td>History of deformities with or without spontaneous or self-reduction</td>
<td>Deformity with displaced, rotated or spiral fractures</td>
</tr>
<tr>
<td></td>
<td>Focal, severe non-radiating pain combined with history of trauma</td>
<td>Point tenderness</td>
</tr>
<tr>
<td></td>
<td>Inability to use the joint</td>
<td>Swelling, hematoma</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ecchymosis</td>
</tr>
<tr>
<td>Dislocation</td>
<td>History of significant trauma</td>
<td>Deformity present</td>
</tr>
<tr>
<td></td>
<td>History of deformities with or without spontaneous or self-reduction</td>
<td>Tenderness and instability with history of deformity with reduction</td>
</tr>
<tr>
<td></td>
<td>Inability to use the joint</td>
<td>Hemarthrosis</td>
</tr>
<tr>
<td>Infection</td>
<td>History of systemic symptoms: fever, chills/rigor</td>
<td>Tenderness with motion</td>
</tr>
</tbody>
</table>
History of immunosuppression (e.g., transplant, chemotherapy, HIV)
Diabetes mellitus
Portal of infection (e.g., laceration, distant infection)

Systemic signs of sepsis
Local heat, swelling, erythema
Drainage of a sinus tract
Painful, red, swollen area(s)

Tumor
History of rapidly growing, painful, firm or hard mass of hand or wrist not consistent with ganglion
History of immunosuppression (e.g., transplant, chemotherapy, HIV)
History of cancer

Mass of hand, wrist, or forearm, not consistent with ganglion or other benign lesion

Joint Inflammation
History of inflammatory arthropathy or crystal arthritis
Clinical history consistent with inflammatory or crystal arthropathies

Swelling and deformity
Mostly symmetrical joint involvement for more common inflammatory arthropathies (e.g., rheumatoid arthritis)
Erythematous, swollen, warm usually solitary joint for acute crystal arthropathy
Painful swollen joints, usually without systemic symptoms

Rapidly Progressive Neurologic Compromise
Rapidly progressive numbness, paresthesias, or weakness in radial, ulnar, or median nerve distribution
Inciting traumatic event or history to produce acute neurological compromise
Progressive weakness
Stroke, cervical spine disorders or other central nervous system compromise

Sensory deficit in ulnar, median, or radial distribution
Loss of finger or grip strength when picking up objects
Atrophy

Vascular Compromise
History of vascular disease
History of diabetes mellitus
Compartment syndrome
Inflammatory arthropathies with vasculitis

Decreased pulses
Decreased capillary filling
Cold, cool, or pale hand

Severe Carpal Tunnel Syndrome
Continuous median distribution tingling and numbness after acute trauma, especially fracture
Severe flexor compartment pain after repeated, unaccustomed, forceful use with continual median distribution tingling and numbness

Reduced median distribution sensation
Muscle atrophy (late) and severe weakness of thenar muscles

Carpal Tunnel Syndrome
The physical examination is particularly helpful for assuring other condition(s) are not present. Some believe the physical examination is highly useful(255) while others suggest the physical examination findings are of limited use in securing a diagnosis as compared with a careful history, and add little to a careful history combined with electrodiagnostic evidence.(10, 254) A recent analysis of signs of carpal tunnel syndrome reported considerable methodological issues, including spectrum biases that likely result in overstatement of the clinical utility of common tests for CTS.(256) Clinical testing for CTS may include several items outlined below. The following describes pertinent history and clinical testing:

- Thenar atrophy – Thenar eminence should appear small compared with the hypothenar eminence and the contralateral hand. This is an advanced sign.
- Hand sensibility – Multiple tests are tested to attempt to determine clinical sensibility. These include Semmes-Weinstein monofilament test, Ten Test, 2-point discrimination, paper clips and various devices. However, sensibility (ability to sense or detect cutaneous stimuli) decreases with age resulting in challenges in interpreting results. Comparison with unaffected digits or the opposite hand is often helpful.(257-260)
- Hypoalgesia in the median nerve territory – Diminished ability to perceive painful stimuli in the median nerve distribution (e.g., palmar aspect of the index finger compared with the ipsilateral fifth digit).
- Monofilament test – A test involving nylon monofilaments that collapse at specific amounts of force when pushed perpendicularly against the palm or fingers. A positive test results when a filament of greater than normal size is required in order for its application to be perceived by the patient.
- Vibration Testing – Diminished ability to perceive vibratory sensations using a standard vibrating tuning fork comparing the distal interphalangeal joint of the index finger to ipsilateral fifth finger.
- Weak thumb abduction strength – Weakness of resisted abduction (i.e., palm horizontal, thumb lifted as vertically as possible, then patient resists examiner pushing the thumb down towards the index finger).
- Hoffmann-Tinel’s Sign (“Tinel’s”) – Up to 6 taps of a reflex hammer or tip of examiner’s finger to the soft tissue overlying the carpal tunnel. A positive test occurs when the taps cause paresthesias or shooting pain in the median nerve distribution.(261)
- Phalen Sign – As originally described, flexion of the wrist by having the examiner passively flex the wrists of the patient for up to 60 seconds.(262) Clinically, this is more commonly performed by having the patient press the dorsal aspect of both hands together with approximately 90° of flexion for 60 seconds. It is unclear if these two means of performing this sign result in different sensitivities and specificities. A positive test produces paresthesias in the distribution of the affected median nerve.
- Carpal Compression Test – The examiner holds the supinated wrist in both hands, flexes the wrist 45° and applies direct, even pressure over the transverse carpal ligament with both thumbs for up to 30 seconds. A positive test is indicated by tingling or paresthesia into the thumb, index finger, and middle and lateral half of ring finger within 30 seconds.(263)
- Tourniquet Test – Paresthesias developing in the distribution of the median nerve when a blood pressure cuff is inflated above systolic pressure for 60 seconds.
- Hand volume – Hand volume change measured by water displacement in a graduated cylinder.

**Triangular Fibrocartilage Complex (TFCC) Tears**
The exam may reveal dorsal-ulnar wrist joint tenderness that is not focally tender over an extensor compartment. Swelling is generally not present, although it may be present with an acute, large tear. The examiner should generally attempt to reproduce catching or snapping in the ulnar wrist joint, either by having the patient place the wrist into a position that elicits the symptoms and/or moving the wrist and forearm through a combined supination movement with simultaneous movement of the wrist from flexion to extension.

**Crush Injuries and Compartment Syndrome**
The physical examination ranges from mild abnormalities with mild injuries (e.g., contusions) to severe with fractures, limited range(s) of motion and neurovascular compromise.

**Kienböck Disease**
The physical examination may be normal early, but generally the patient has mild to moderate dorsal wrist tenderness while also having asymmetric, limited range of motion. Tenderness and limited range of motion tend to progress.

**Wrist Sprains**
The exam may include wrist capsule tenderness, or it may be normal. Deformity suggests fracture. Scaphoid tubercle tenderness suggests scaphoid fracture.

**Mallet Finger**
The patient is unable to extend the distal phalangeal segment. Swelling often signifies a fracture fragment, while most are extensor tendon ruptures(264) and have no significant swelling.

**Flexor Tendon Entrapment (Tenosynovitis and Trigger Digit)**
Patients without triggering will typically have tenderness localized over the A1 pulley.(265) A palpable tendon nodule is frequently present. Triggering often occurs upon arising in the morning or after lack of use. Active movement is often required to demonstrate triggering as passive motion is often normal. Those rare cases with a locked digit are unable to extend (or flex) the digit.(31, 33-35, 40, 190-192, 198, 237, 242, 249, 266-282)
Extensor Compartment Tenosynovitis (Including de Quervain’s Stenosing Tenosynovitis and Intersection Syndrome)

The hand is usually normal in appearance, although there is visible tendon sheath swelling in a minority of cases. Swelling is more common with inflammatory conditions (e.g., rheumatoid arthritis) or infections. Some believe swelling and crepitus are also only present among those with peritendinitis if there is no inflammatory or infectious disease. Focal tenderness over the compartment is present. Finkelstein’s maneuver is the classic provocative maneuver and is nearly always present,(41) however, the predictive values are unclear. Pain in the affected compartment is generally present with use or any provocative maneuver (e.g., resisted use of the muscle-tendon unit).(42) Triggering may be demonstrated on rare occasions.

Ulnar Nerve Entrapment at the Wrist (including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome)

Dependent on the location of the lesion, motor, sensory, or mixed motor-sensory findings are detectable. Muscle atrophy may be present in the interosseous and hypothenar areas. Point tenderness may be present. Sensory loss is typically most prominent at the palmar tip of the 5th finger, in contrast with ulnar neuropathies at the elbow which present with sensory loss on the palmar and dorsal surfaces of the 5th digit. Motor weakness may be demonstrated by resisting spreading of the fingers to assess intrinsic muscle strength. A positive Tinel’s is purportedly helpful, but there are no quality studies evaluating the utility of this clinical test and its utility elsewhere has been questioned. Associated carpal tunnel compression should also be examined. A vascular exam and auscultation for bruits should be performed,(283) particularly for those cases thought to involve vascular symptoms and hypothenar hammer-like symptoms.

Radial Nerve Entrapment

The physical examination attempts to localize the site of nerve entrapment and should include sensory (especially sensation) and motor components (movement, range of motion, strength, reflexes) to localize the entrapment. Comparisons to the unaffected limb should be made. Differentiation from de Quervain’s tenosynovitis is a primary differential diagnostic consideration, yet Finkelstein’s is not particularly helpful as it may be positive with both conditions.

Non-Specific Hand/Wrist/Forearm Pain

The examination is generally without any unequivocally objective evidence. Instead, tenderness is most often the only physical examination finding. Qualitative muscle strength testing may be weak compared with the unaffected side. Precise documentation of the location of the pain should be made with consideration for photographing the location for future reference. In cases where the pain does not migrate, the probability of specifically defined pathology is believed to increase.

Scaphoid Fracture

Physical examination findings include antalgic behavior with avoidance of use of the hand, and tenderness over the scaphoid tubercle.(284-286) Scaphoid tubercle tenderness may be more sensitive and specific than snuffbox tenderness. The scaphoid tubercle is located at the volar wrist at the junction of the distal wrist crease under the flexor carpi radialis. The tubercle becomes prominent and readily palpable with radial deviation of the wrist. Patients may also have tenderness over the snuffbox, absence of tenderness in the distal radius, wrist joint effusion,(287-289) and scaphoid pain on axial loading of the thumb (“scaphoid compression test”).(285, 290) However, many of these findings may also be present without scaphoid fracture. An isolated finding of snuffbox tenderness appears to be sensitive, but has poor positive predictive value for scaphoid fracture.(71, 286, 291)

Distal Phalanx Fractures and Subungual Hematoma

Physical examination should include inspection and identification of localized swelling and open wounds. Neurovascular status should be described. The DIP joint should be palpated in each plane to assess point tenderness over ligament insertions. Passive range of motion and joint stability should be assessed through dorsal, volar, and lateral stressing. An estimate of subungual hematoma size relative to the nail bed surface should be noted. A case series demonstrated fractures in 63% of patients with hematomas...
greater than 50% of surface area, 43% of patients with hematomas 25 to 50% of nail surface, and 10% in those with less than 25% of nail bed surface involvement. (75) The DIP joint should be evaluated for flexion and extension range of motion.

**Middle and Proximal Phalangeal and Metacarpal Fractures**
Prior to fracture manipulation, physical examination includes evaluation of digital nerves using two point discrimination or pin prick, tendon and ligament integrity with active and passive range of motion at each joint, vascular status with capillary refill, and surrounding soft tissue structures of affected areas. (77) Finger shortening or knuckle depression may be present. Bone alignment should be checked for rotational deformity by finger flexion of hand, with the nails pointing toward the scaphoid tubercle. The natural alignment will be disrupted if a rotational fracture is present, such that one finger will overlap another.

**Distal Forearm Fractures**
Comprehensive physical examination for traumatic injuries at the wrist as well as elbow, shoulder, neck, head, and hip should be included. Examination of the injured wrist and hand should include neurological and vascular exam, as well as testing for tendon and ligament integrity. The ulnar styloid should be palpated for tenderness as well as the radial head. TFCC should be suspected for displaced or complex fractures, and DRUJ instability may be noted dependent on extent of pain and nature of fracture.

**Ganglion Cyst**
Wrist ganglia are usually well demarcated, firmly tethered, and have a consistency similar to a rubber ball, and are translucent. Lack of translucency should raise suspicion of other tumor type. The mass and surrounding skin should be inspected and palpated for erythema and infection. Examination should also include close inspection for mass effect, including neurovascular involvement, impairment of wrist or finger joint range of motion, impairment of tendon function, and triggering. Small occult dorsal wrist ganglia may result in tenderness over the scapholunate ligament and pain with hyperextension of the wrist. (292)

**Hand-Arm Vibration Syndrome**
A complete examination should include close attention to motor, sensory and vascular functions of the affected extremities. Evaluation should be extended to the shoulder and neck for upper extremity symptoms including tests for vascular insufficiency. Particular note should be made for blanching, coordination of movement, grip strength, tenderness and swelling of the digits and forearm tissue, and trophic changes of the skin. The value of cold provocation or neurophysiological tests in the diagnosis is controversial. (293, 294)

**Laceration Management**
Close inspection of the wound should be performed under proper lighting. Control of bleeding may be required, generally by applying appropriate pressure and elevation to the wound. The wound should be evaluated for damage to underlying structures including joint involvement, vessels, tendons, bone and nerves. Sensory examination should be accomplished prior to anesthetic administration. Examination of involved muscles should be conducted if nerve injury is suspected. Close inspection should be made for foreign bodies.

**Human Bites, Animal Bites and Associated Lacerations**
The wound should be carefully cleaned and inspected for depth of injury, potential associated crush injury or fracture, tendon or tendon sheath involvement, foreign body (e.g., teeth, fur, soil), and joint space involvement.

**Hand/Finger Osteoarthrosis**
Mild cases may show few, if any abnormalities. However, as the disease progresses, more findings develop. Boney enlargement of the affected joint(s) is present on inspection and range of motion is usually reduced. The most commonly affected joint is the thumb carpometacarpal (CMC) joint, which may become enlarged and deformed. Boney enlargement of the distal interphalangeal joints is termed “Heberden’s nodes” while of the proximal interphalangeal joints is called “Bouchard’s nodes.” Crepitus on range of
motion is often present. Joints are generally not warm, have no significant joint effusion and are usually non-tender.

**Diagnostic Criteria**
The criteria presented in the Diagnostic Criteria for Hand, Wrist, or Forearm Disorders table (Table 4) list the probable diagnosis or injury, potential mechanism(s) of illness or injury, symptoms, signs, and appropriate tests and results to consider in assessment and treatment.

**Table 4. Diagnostic Criteria for Hand, Wrist, or Forearm Disorders**

<table>
<thead>
<tr>
<th>Probable Diagnosis or Injury</th>
<th>Unique Mechanism (includes only physical factors; in some cases there are other factors)</th>
<th>Unique Symptoms</th>
<th>Unique Signs</th>
<th>Tests and Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ligament Sprain</td>
<td>Acute excess loading, generally from falling onto an extremity. Increased pain with motion.</td>
<td>Focal pain in ligament</td>
<td>Tenderness over ligament(s) Pain or weakness on strength testing of the affected ligament(s)</td>
<td>X-rays (normal)</td>
</tr>
<tr>
<td>Tuft Fracture</td>
<td>Crush injury to distal phalangeal segment</td>
<td>Pain and deformity of tip of digit. May have subungual hematoma or other deformity(ies)</td>
<td>Crush injury to tip of digit</td>
<td>X-rays with tuft fracture.</td>
</tr>
<tr>
<td>Mallet Finger</td>
<td>Forceful flexion of DIP joint while digit is extended. Ball striking tip of digit or digit extended during fall. Some rupture spontaneously, usually over a Heberden’s node from osteoarthrosis.</td>
<td>Unable to extend digit at DIP joint. Usually pain-free if no accompanying fracture.</td>
<td>Incapable of extension at DIP joint. May be swollen, particularly with fracture</td>
<td>X-ray occasionally may show fracture, but usually normal. May not have fracture if extensor mechanism ruptured without fracturing bone</td>
</tr>
<tr>
<td>Myotendinous Strain</td>
<td>Unaccustomed forceful use. May be from acute loading or fall. Worse pain with motion</td>
<td>Focal pain at a discrete myotendinous junction</td>
<td>Tenderness over myotendinous junction. Pain or weakness on strength or resistance testing of the affected myotendinous junction. Crepitus on motion particularly if accompanied by tenosynovitis or peritendinitis.</td>
<td>None</td>
</tr>
<tr>
<td>Tendinosis/ Tendinitis/ tenosynovitis</td>
<td>High force and repetition, stereotypical awkward motions, combinations of physical factors Direct pressure (unusual) Blunt trauma (rare) (Diagnosis of “tendinitis” also frequently used as a diagnostic label for “pain” without pathophysiological correlation.)</td>
<td>Pain localized to flexor or extensor compartment. Triggering may be present if digital flexor compartment involved</td>
<td>Tenderness over discrete flexor or extensor compartment tendon (usually not more than 1 or 2 compartments) Synovial thickening Triggering or locking Crepitus Pain or weakness on strength testing of the affected tendon</td>
<td>None</td>
</tr>
<tr>
<td>De Quervain’s Tenosynovitis</td>
<td>High force and repetition with forceful wrist and thumb motion Direct pressure (unusual) Blunt trauma (rare)</td>
<td>Pain over radial styloid in area of first dorsal compartment</td>
<td>Tenderness over radial styloid Mass over radial styloid (rare) Crepitus over extensor compartment Thick tendon sheath</td>
<td>None</td>
</tr>
<tr>
<td>Trigger Finger, Trigger Thumb</td>
<td>High force and repetition Blunt trauma (rare)</td>
<td>Triggering Pain at volar metacarpal phalangeal joint Locked finger</td>
<td>Triggering Tender volar metacarpal crease Tendon nodule Synovial thickening of specific parts of flexor retinaculum</td>
<td>None</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>---------------------------------</td>
<td>-------------------------------------------------</td>
<td>-------------------------------------------------</td>
<td>------</td>
</tr>
<tr>
<td>Carpal Tunnel Syndrome</td>
<td>High force and repetition, combinations of physical factors Vibration (Associated factors include cold temperatures and glove use. Posture is unclear factor, thought to be a relatively weak factor.)</td>
<td>Numbness/tingling in thumb, index, middle, radial half of ring finger, especially at night or with activity Volar hand pain radiating into forearm may be present. Decreased grip strength Difficulty picking up small objects Hand symptoms diagram</td>
<td>Atrophy or decreased strength of abductor pollicis brevis, opponens (advanced cases) Decreased sensation in median nerve distribution (including monofilaments)</td>
<td>Electrodiagnostic studies</td>
</tr>
<tr>
<td>Ulnar Neuropathy at the Wrist and Hypothenar Hammer Syndrome</td>
<td>Repeated striking of the heel of the hand/hypothenar region on a tool or object</td>
<td>Pain in hypothenar region, blanching of ulnar artery distribution (especially 5th digit), Paresthesias in small and ring fingers</td>
<td>Tender hypothenar region, blanching of ulnar artery distribution (especially 5th digit), decreased sensation in small and ring fingers</td>
<td>Ulnar artery Doppler/ultrasound, electrodiagnostic studies</td>
</tr>
<tr>
<td>Hand-Arm Vibration Syndrome</td>
<td>Repeated, prolonged use of low-frequency, high-amplitude vibrating tool, especially in cold environments</td>
<td>Pain in the fingers, episodic finger blanching</td>
<td>Blanching of fingers, worse with cold provocation. Ulceration of finger tips when severe.</td>
<td>None</td>
</tr>
<tr>
<td>Nonspecific Pain</td>
<td>Unknown as condition is idiopathic; possibly resulting from combination of risk factors. May be psychological condition.</td>
<td>Pain, but non-specific</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Ganglion</td>
<td>Unknown</td>
<td>Painful or painless mass on wrist, hand, or any other joint</td>
<td>Tender (or non-tender) mass most commonly over dorsal or volar wrist or hand</td>
<td>None</td>
</tr>
</tbody>
</table>

**Special Studies and Diagnostic Considerations**

For most patients presenting with non-traumatic hand, wrist, and forearm disorders, special studies are not needed during the first 4 weeks. Most patients improve quickly, provided red flag conditions are ruled out. Exceptions include the following:

- In cases of wrist injury, with tenderness over the scaphoid (especially over the scaphoid tubercle), but minimal other findings, a scaphoid fracture may still be present. Initial radiographic images should be obtained, but may appear negative in the presence of nondisplaced scaphoid fracture. If clinical symptoms continue, a re-evaluation with new radiographs is advised in approximately 2 weeks.
- An acute injury to the metacarpophalangeal joint of the thumb, accompanied by tenderness on the ulnar side of the joint and laxity when that side of the joint is stressed (compared to the other side), may indicate a gamekeeper’s thumb or rupture of the ulnar collateral ligament of the MCP joint. Radiographic images may show a fracture or stress views, if obtainable, may show laxity. The diagnosis may necessitate surgical repair of the ligament and surgical referral is warranted.

Also, of note, a number of patients with hand, wrist, and forearm symptoms will have associated disease such as diabetes mellitus, hypothyroidism, renal disease, and one or more of the arthritides which are often heretofore undiagnosed. When medical history and/or physical examination findings indicate or other risk factors are present, testing for these or other comorbid condition(s) is recommended.

**Work-Relatedness**

There are numerous occupational and non-occupational risk factors for hand, wrist, and forearm (upper limb) musculoskeletal disorders (MSDs). (6, 12, 196-198, 201, 202, 205, 206, 208-210, 212, 232, 295, 296) Most available quality evidence has been reported on CTS, with sparse information on other disorders. While some risk factors (e.g., age, obesity, (175, 213) diabetes mellitus, and metabolic syndrome(297)) generally appear in common with most MSDs, other risk factors do not appear in common across the disorders (e.g., low density lipoprotein,(298) thyroid disorders, pregnancy). The lack of common risk factors across the spectrum of disorders raises questions about the accuracy of generalizing any risk factor, whether occupational or non-occupational across all disorders. Some of these inconsistencies among studies may also be due to lack of statistical power to identify relationships between these factors and upper extremity MSDs.(29, 145, 175, 213, 215, 217, 299, 300)

Work-relatedness of hand, wrist and forearm MSDs is dependent on the precipitating exposure(s). For acute, traumatic injuries (e.g., dislocations, true ligamentous sprains, mallet finger, fractures), the work-relatedness is determined by whether the inciting event occurred out of, or in the course of employment. Such determinations of work-relatedness are rarely difficult or controversial.

Non-traumatic MSDs (e.g., CTS, tendinoses, tendinitis, trigger digit) are often difficult to attribute to work to a medical degree of certainty. There are many retrospective studies of these CTS and tendinoses. However, recently there are several prospective cohort studies evaluating risk. One cohort in industrial and clerical workers found the greatest predictors of upper extremity tendinosis were older age, body mass index over 30, shoulder or neck discomfort at baseline, history of CTS and a higher shoulder posture rating.(300) Another prospective study of production workers reported associations of hand/wrist tendinitis to repeated forceful pinching at work.(301) A prospective study of automobile workers found increased risks for incident cases of CTS included a history of wrist/hand/finger tendinoses, diabetes mellitus, nonneutral wrist and elbow postures, lower social support, and greater differences between baseline median and ulnar nerve peak latencies across the wrist.(185) One cohort study of repetitive work in Denmark found incidence rates over 3 to 4 years were too low to compare the risk among those doing highly repetitive work (0.62%) versus more variable work (0.44%).(302) However, combinations of forceful and repetitive hand activities at work as combined in the American Conference of Governmental Industrial Hygienists hand-activity level have been found to increase risk of CTS in several prospective studies(145, 146, 213, 216, 218, 219, 303) which is consistent with findings from numerous cross sectional studies.(197, 199, 204, 207, 304-310) Attributable CTS risk from high-risk occupations in France is estimated to range from 36 to 93%. (311)

A thorough work history is crucial to a foundation for establishing work-relatedness. Determining whether a complaint of a hand, wrist, or forearm disorder is related to work requires a careful analysis and weighing of all associated or possible causal factors operative at the time.(232, 312) A predominance of work factors suggests that worksite evaluation may be appropriate.

A broad range of ergonomic surveys and instruments is available for measuring range of activity, strain, weights, reach, frequency of motion, flexion, and extension, as well as psychological factors such as organizational relationships and job satisfaction (e.g., Motion Time Measurement analyses, ACGIH TLV for Hand Activity Level, Strain Index).(143, 313, 314) However, physicians should be aware that currently,
no job evaluation method has been fully validated. Nevertheless, documentation of job physical factors in conjunction with adverse health effects is often necessary to facilitate and substantiate engineering and organizational changes (see individual sections for discussions of work-relatedness of specific hand, wrist and forearm disorders).

Carpal Tunnel Syndrome
There are numerous occupational and non-occupational risk factors for CTS, as well as other hand, wrist, and forearm musculoskeletal disorders.(197, 199, 204, 211, 212, 232, 296, 304, 306) Many studies on CTS have not used objective measures that included electrodiagnostic testing in case definitions, rather they relied solely on symptoms or combinations of symptoms and physical examination findings (e.g., Hoffman-Tinel's sign).(197)

A thorough work history is crucial to a foundation for establishing work-relatedness (see Work-Relatedness Guideline for a method to determine work-relatedness). Non-occupational risk factors that have been most consistently identified in numerous studies for CTS include age, gender, body mass index (BMI), diabetes mellitus, and wrist depth/width ratio.(6, 12, 196-198, 201, 202, 205, 206, 208-211, 295, 311, 315-318) Physicians should also be aware of the high prevalence of CTS in the general population, its strong relationship with age, and the relatively high prevalence of nerve conduction abnormalities in the population, some of which are asymptomatic.(319) Determining whether a complaint of a hand, wrist, or forearm symptoms are related to work requires a careful analysis and all associated or possible causal factors operative at the time must be weighed.(232, 312) A predominance of work factors suggests that worksite evaluation is likely appropriate and intervention may also be appropriate. A careful ergonomic assessment, work management, and other preventive measures are also suggested when a cluster of cases in a work group occurs.

Based on recent prospective studies, the sustained or repeated application of forceful pinching or gripping is thought to be the most potent work-place activity related risk factor for CTS and hand or wrist tendinosis, particularly when combined with high rates of repetition.(145, 146, 185, 213, 216, 218, 219, 301) The risk appears present when pinch forces are greater than 10 N (1kg).(204, 216, 301) Carpal tunnel syndrome risk appears most strongly increased in jobs involving high-force gripping such as meat processing, manufacturing, and farming.(311, 320-322)

Keyboard use is often a highly repetitive, but very low-force task with very different physical exposures than non-computer work and having many mostly retrospective epidemiological studies previously reported.(302, 318, 323-336) Prospective cohort studies have failed to find associations between CTS and keyboard use,(318, 323, 337, 338) however, one of these studies reported increased risk with increased mouse use in both its baseline/cross-sectional analyses and cohort analyses.(323) Case-control studies have reported conflicting results, with one reporting reduced risk with increased hours spent typing(326) and one reporting increased risk with typing more than 4 hours per day.(333) In several large prospective studies, increasing hours of computer work was associated with tendinosis, de Quervain’s disease, and non-specific hand, wrist, forearm and elbow, neck and shoulder pain.(337, 339, 340) Split keyboards have been associated with reduction in pain and disorders.(341, 342)

Triangular Fibrocartilage Complex (TFCC) Tears
Work-relatedness of an acute TFCC tear sustained in the course of a slip, trip, fall, or heavy and awkward lift at work is generally considered an occupational injury and is not usually controversial, although apportionment is a consideration in applicable jurisdictions due to the prevalence of pre-existing degenerative abnormalities, as well as presence and degree of ulnar positive variance (longer ulna than radius, which is thought to be a risk factor for TFCC tears). However, other TFCC tears occurring without an acute, inciting event are of unclear etiology, as the injuries are more analogous to a disease or disorder and there is no quality epidemiological evidence to link them with work. Chronic TFCC tears in the presence of a positive ulnar variance are generally not thought to be occupational.
Acute Trauma, including Fractures (e.g., scaphoid, phalangeal, distal forearm), Crush Injuries, Compartment Syndrome, Subungual Hematomas, Lacerations, and Animal Bites

Caution is based on the specific major incident that produced the injury. Wrist sprains do not occur without an acute, precipitating significant mechanism of injury.

Kienböck Disease
This disorder is a disease without sound epidemiological support for work-relatedness. It may be reasonable to hypothesize work-relatedness in those cases where the onset is promptly after a discrete, significant traumatic event. However, in most cases, a physical cause is speculative.

Mallet Finger
Work-relatedness is generally non-controversial and is based on having an acute accident at work. However, in cases without precipitating injury, work-relatedness is speculative.

Flexor Tendon Entrapment (Tenosynovitis and Trigger Digit)
As the epidemiological evidence is weak, the etiological fraction for occupational tasks is unknown.(37) Thus, work-relatedness is somewhat unclear.(232) The available biomechanical evidence suggests pinch force may be a risk factor.(35, 40, 190, 192, 234-245)

Extensor Compartment Tenosynovitis (Including de Quervain’s Stenosing Tenosynovitis and Intersection Syndrome)
Work-relatedness is thought to be present in a significant proportion of cases,(277, 343, 344) although more recent studies have suggested less work-relatedness.(232) Risk factors have not been confirmed in cohort studies, but are thought to particularly involve combinations of force, repetition and posture.(41, 277, 337, 343-345) Direct trauma over the affected extensor compartment is reported in a minority of cases.(41) Risks for intersection syndrome are not well defined. Purported risks appear to be high-force sports related particularly if unaccustomed including rowing, canoeing, racket sports, and weight lifting.(346, 347) Work tasks reported to be risks appear similar with intensive agricultural workers (threshing, planting, hammering, hand washing, spraying, cementing)(43) and recent job change to supermarket cashiering being examples of reported risks.(348) Discontinuation of the high force, unaccustomed activity has been frequently reported to resolve intersection syndrome.(43, 45, 349, 350) Increasing hours of computer work has been associated with extensor compartment tenosynovitis, de Quervain’s disease, and non-specific wrist and forearm pain.(337, 339) Those risks may be due to contact stress at the wrist or sustained wrist postures. Split keyboards, which reduce awkward postures, have been associated with reduction in pain and disorders.(341, 342)

Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome)
Ulnar neuropathy at the wrist is reportedly most often caused by a space occupying lesion such as ganglion, scar, abnormal ulnar artery or aneurysm, and trauma(51, 57, 63) (i.e., resulting from hamate fracture). Experimental studies suggest that the nerve moves within the canal with wrist motion, thus traction on the nerve may be possible.(64) In a case series of 47 patients, suspected “cumulative trauma” was attributed to nearly 75% of cases. However, no definition or quantification of physical factors was given. Activities included both mechanisms with potential significant localized compression (e.g., cycling, wheelchair use), those without apparent compression (e.g., piano, truck driving), and those that may or may not have involved compression (e.g. boiler operator, machine press operator, and steel cutter).(50) Other described causes include aberrant muscles at the wrist affecting both median and ulnar nerves,(351) and distal neuropathy caused by systemic diseases, particularly diabetes mellitus and systemic sclerosis (scleroderma).(352-354) As there are no quality epidemiological studies among non-traumatic patients, work-relatedness is speculative in those populations.

Radial Nerve Entrapment
Radial neuropathy at the wrist is reportedly caused by local mechanical compression of the nerve at the wrist from external trauma, a tight wrist or forearm band, or anomalous brachioradialis tendon.(61, 62) It
has been attributed to repeated wrist flexion and ulnar deviation, however, there is no quality epidemiological evidence and thus when occurring in the absence of trauma, work-relatedness is speculative. There may be a better basis for work-relatedness for radial neuropathy with entrapment just above the wrist in the context of concomitant de Quervain’s tenosynovitis that is considered work-related.

**Non-Specific Hand/Wrist/Forearm Pain**
Work-relatedness is unclear as there are no quality studies of this condition. However, it is generally recommended that the condition be treated and it will generally resolve. Thus, in the absence of costly testing and/or treatment protocols or prolonged duration, the condition is generally non-controversial.

**Ganglion Cyst**
No quality epidemiological studies have shown work relatedness. However, ganglia may be accepted as work related in some legal jurisdictions. In a cross-sectional survey of more than 30,000 workers in the 1988 National Health Interview Survey, the prevalence of clinical ganglion cyst was estimated at 14%. (113) Of all cases, it was estimated based on patient report of physician diagnosis that nearly 6% were attributed to work. However, there were no analyses based on occupation or activity. There were no quality epidemiologic studies addressing work place or occupational physical factors.

**Hand Arm Vibration Syndrome**
Work-relatedness is based on confirmation of the diagnosis and a mechanism of occupational injury where there is an appropriate exposure which is generally low frequency high amplitude vibration.

**Hand/Finger Osteoarthrosis**
There is one cross sectional study from the textile industry that suggests some cases of hand osteoarthrosis may have a component of occupational tasks; (355) however, those jobs are likely no longer present in the U.S. In most patients, multiple joints are symmetrically affected. Yet, occupational exposures are frequently not symmetrical and do not explain this association, thus these cases are usually believed to be non-occupational. However, there are cases of monoarticular osteoarthrosis occurring in a joint affected by a remote, traumatic event such as a fracture involving the joint or adjacent to the joint, dislocation or significant sprain. Work-relatedness of those cases is generally non-controversial as it is believed to be a consequence of the acute traumatic event.

**Job Analysis**
Some hand, wrist, and forearm symptoms are occupational in origin, with the occupational aspect differing by industry, job task, or by the disorder in question. Decisions about which jobs to analyze, and their prioritization, is of increasing importance as the proportion of affected individuals has been identified as in excess of 50% of the workforce per annum in settings of combinations of high force and high stereotypical occupational activity. In general, prioritization of job analyses in workplace settings is based on the numbers of affected individuals, reported and perceived rates of MSDs, costs and severity of the disorders, and planned job redesigns. From an occupational health care perspective, ergonomic analysis of a job may also be indicated for failure to improve in the absence of other plausible explanations. The employer’s role in accommodating activity limitations and preventing further problems through ergonomic changes may be a key factor in hastening the employee’s return to full activity, particularly among workers with a history of high job physical factors. In some cases, it may be desirable to conduct an ergonomic analysis of the activities that may be contributing to the symptoms.

**Carpal Tunnel Syndrome**
In certain cases, conduct an ergonomic analysis of the activities that may be contributing to the symptoms should be conducted. A broad range of ergonomic surveys and instruments is available for estimating duration of hand intensive activities, grasp repetition rates, pinch force, part or tool weights, reach distance, frequency of motion, wrist and hand postures, as well as psychological factors such as organizational relationships and job satisfaction. (313) As the most robust data support two of those ergonomic instruments, it would appear most wise to utilize one or both of them (i.e., American Conference of Governmental Industrial Hygienists Threshold Limit Value for Hand Activity, (218, 356) and Strain Index(314)). Such
detailed measures may be necessary or useful for modifying work activity, selecting tools, redesigning the workstation, or recommending organizational and management initiatives. Such situations may call for referral to certified professional ergonomists or a human factors engineer, either through the patient or the employer. Some occupational therapists, physical therapists, occupational medicine physicians and other professionals also may have appropriate credentials and experiences to accomplish these evaluations.

**Triangular Fibrocartilage Complex (TFCC) Tears**
Job analysis is generally not indicated for most cases, particularly if the focus of the analysis is unclear. However, where there is potential to eliminate a hazard that precipitated the event (e.g., slippery surface), job analysis is recommended.

**Acute Trauma, including Fractures (e.g., scaphoid, phalangeal, distal forearm), Crush Injuries, Compartment Syndrome, Subungal Hematomas, Lacerations, and Animal Bites**
Job analyses may be of benefit to attempt to prevent future occurrences of these types of injuries (e.g., machine guarding, icy walkways, tool kickback). Some of these, particularly compartment syndrome and fractures should generally be analyzed for root cause and potential remediation, as these injuries are generally viewed as critical incident cases.

**Kienböck Disease**
As physical risk factors are undefined, job analyses are unhelpful.

**Mallet Finger**
Job analysis is generally not indicated for most cases, although where there is potential to eliminate a risk for slips, trips of falls it should be resolved.

**Flexor Tendon Entrapment (Tenosynovitis and Trigger Digit)**
Job analyses may be useful for evaluating for forceful finger use and localized contact stress (e.g., compression by sharp objects). (35, 40, 190, 192, 234-245) There is now prospective evidence that the Strain Index is predictive for trigger digit. (29)

**Extensor Compartment Tenosynovitis (Including de Quervain’s Stenosing Tenosynovitis and Intersection Syndrome)**
Job analyses may be useful to identify repeated, forceful digit use, sustained extreme digit postures, or localized compression by sharp objects.

**Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome)**
Job analyses may be useful to identify tasks involving considerable hypothenar area localized mechanical compression, as well as use of the hand as a hammer.

**Radial Nerve**
Job analysis may be useful to identify tasks involving external compression of the radial nerve at the wrist due to occupational tasks.

**Non-Specific Hand, Wrist, or Forearm Pain**
Job analysis is difficult for many of these conditions, particularly as the discrete entity to be evaluated and job analysis methods are unclear. However, job analyses may also be revealing particularly when there is a high exposure situation (i.e., high force or combinations of high force and other ergonomic risk factors). This may be especially indicated where other cases of musculoskeletal disorders are present in the workforce and may help with the treatment plan.

**Ganglion Cyst**
There is no quality evidence that the cause of these cysts is work related, thus job analyses are not generally indicated.

**Hand Arm Vibration Syndrome**
Job analyses are generally indicated for cases with this diagnosis, particularly for primary, secondary and tertiary prevention. The analyses are fairly technical and usually require special equipment to measure vibration exposures.

Hand/Finger Osteoarthrosis
Job analysis is generally not indicated for most cases, although where there is potential to eliminate a hazard that precipitated an acute event (e.g., icy sidewalk, tripping hazards), it should be resolved. There have been no quality job analysis tools developed to analyze jobs for risk of hand osteoarthrosis.

Ergonomic Interventions for Distal Upper Extremity Musculoskeletal Disorders with an Occupational Basis
In order to facilitate recovery and prevent recurrence of distal upper extremity musculoskeletal disorders, one may recommend work and activity modifications or ergonomic redesign of the workplace.(357) The employer’s role in accommodating activity limitations and preventing further problems through ergonomic changes is crucial in hastening the employee’s return to full activity. In some cases it may be desirable to conduct an ergonomic analysis of the activities that may be contributing to symptoms. A broad range of ergonomic surveys and instruments is available for estimating duration of hand intensive activities, grasp repetition rates, pinch force, part or tool weights, reach distance, frequency of motion, and wrist and hand postures, as well as psychological factors such as organizational relationships and job satisfaction. Such detailed measures may be necessary or useful for modifying activity, redesigning the workstation, or recommending organizational and management relief. Such situations may call for referral to a certified ergonomist or a human factors engineer. Alternate keyboard layouts have been used to reduce disorders.(358-361)

1. Recommendation: Ergonomic Interventions for CTS and Common Distal Upper Extremity Tendinoses
   In settings with combinations of risk factors (e.g., high force combined with high repetition), ergonomic interventions are recommended to reduce risk factors for CTS and common distal upper extremity tendinoses.
   
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   
   Level of Confidence – Low

2. Recommendation: Typing Posture for Prevention and Treatment of CTS and Common Distal Upper Extremity Tendinoses
   Mandating typing in a 90° traditional posture is not recommended for prevention or treatment of CTS and distal upper extremity tendinoses.

   Strength of Evidence – Not Recommended, Evidence (C) – Prevention
   
   Not Recommended, Insufficient Evidence (I) – Treatment

   Level of Confidence – Low

   The use of alternate or split keyboards is recommended among select patients with common distal upper extremity tendinoses.

   Strength of Evidence – Recommended, Insufficient Evidence (I)
   
   Level of Confidence – Low

4. Recommendation: Forearm Support for Typing to Prevent Neck/Shoulder Symptoms
   Forearm support for frequent computer keyboard users is recommended for potential prevention of neck and/or shoulder symptoms (see Figure 1).

   Strength of Evidence – Recommended, Evidence (C)
   
   Level of Confidence – Low
5. **Recommendation: Trackballs for Treatment of Select Patients with CTS**  
A trackball (instead of a mouse) is recommended for treatment of select patients with symptoms of CTS (see Figure 2).  
*Strength of Evidence – Recommended, Insufficient Evidence (I)*  
*Level of Confidence – Low*

**Figure 1. Forearm Support for Typing**  
Reprinted with permission from David Rempel, MD.

6. **Recommendation: Computer Typing Breaks for Patients with CTS, Other Common Extensor and Flexor Hand/Wrist Tendinoses, or for Primary Prevention**  
Computer typing breaks are recommended for select patients with symptoms of CTS or other common extensor and flexor hand/wrist tendinoses as well as for primary prevention.  
*Strength of Evidence – Recommended, Insufficient Evidence (I)*  
*Level of Confidence – Low*

7. **Recommendation: Ergonomics Training in Moderate- or High-risk Manufacturing Settings**  
Ergonomics training is recommended in moderate- or high-risk manufacturing settings.  
*Strength of Evidence – Recommended, Insufficient Evidence (I)*  
*Level of Confidence – Low*

8. **Recommendation: Ergonomics Training for Prevention of MSDs in Office Settings**  
There is no recommendation for or against the use of ergonomics training for the prevention of MSDs in office settings.  
*Strength of Evidence – No Recommendation, Insufficient Evidence (I)*  
*Level of Confidence – Low*

**Rationale for Recommendations**

Ergonomics interventions have been attempted in numerous occupational settings.(362-365) Randomized controlled trials (RCTs) of changes to manufacturing and production positions have not been reported. However, a few RCTs have been reported of keyboard workstations (“office ergonomics”).(342, 362, 363, 366) There is one RCT regarding comparing a dental pick and finding lower pain in the group with the lighter tool that has a wider handle.(365) There also have been quality studies reported regarding participatory ergonomics programs. However, those are mainly reports of patients with spine disorders in programs whose purpose is return to work (see Low Back Disorders Guideline).(367) Despite the lack of quality evidence in most settings, reductions in job physical factors, particularly high force, are thought to be beneficial(365, 368) (see Work-Relatedness). There also are experimental studies of different equipment,(369) although reductions in injuries have not been shown in quality studies.

There are no quality studies of ergonomic interventions for common distal upper extremity MSDs in physically demanding occupations. Interventions which reduce forceful, repeated pinching or alleviating localized compression by sharp objects may be theoretically helpful.(35, 40, 190, 192, 234-245) Quality evidence is not available for effectiveness of ergonomic interventions on MSD injury rates in typical manufacturing settings. However, given available evidence of risk factors, interventions are recommended...
where there are combinations of risk factors; particularly combined high force and high repetition (see Work-Relatedness). Management/supervisor and labor/employee support are often necessary for optimal success of these programs.

Quality evidence has reported no beneficial effects of the 90° typing posture (seated erect, feet on floor, knees, hips, and elbow joints all at 90° angles), instead it has been found to have the same injury rates as a “laid back” posture.(366) “Ergonomic keyboards” involve a split design that produces a neutral wrist posture in comparison with a standard keyboard that requires approximately 15° of ulnar deviation. However, those keyboards have not been reported in quality studies to result in injury reductions, with the exception of a physically split keyboard with sharply angled keyboard faces (Figure 3). (362) Evidence of superiority of these interventions is currently weak. Providers should be aware that not all split designs are equivalent and there is evidence that a widely split keyboard with sharply angled keyboard faces is not well tolerated. (342) Additional quality studies are needed. Forearm supports for typing have been reported to result in fewer neck/shoulder symptoms. (363, 370) Quality evidence suggests reductions in symptoms may be realized from use of a trackball; however, providers should also be aware that there was a small, non-statistically significant increase in pain complaints among a minority of users. (363) This suggests that careful worksite or clinical observation, combined with instructions to discontinue use if symptoms materially increase, may be desired during this intervention. It also suggests that having multiple options available for workstations is desirable. Quality evidence suggests reductions in neck/shoulder symptoms may be realized through utilization of a forearm support. (363, 370)

Breaks from computer typing have been addressed in a low-quality study which reported reductions in symptoms, but no additional benefit from utilizing exercise during breaks. (371) Various types of breaks have been utilized including stretching breaks and exercise programs. (372, 373, 374, 375-380) Quality evidence supporting the efficacy of breaks is weak, especially for symptomatic patients. (371-373) One low-quality randomized study among an apparently asymptomatic population of temporary data-entry workers suggested fewer symptoms among those taking breaks; however, compliance was low (ranging from 25 to 39%). Breaks are not invasive, have no substantial adverse effects, are low cost, and do not appear to impair productivity. (371, 373, 378, 380-385) Widespread use of these programs has not been reported in quality studies; however, with no apparent significant cost impacts and studies suggesting potential benefits, breaks are recommended for both primary prevention and treatment of symptomatic patients.

While quality evidence is lacking for the use of ergonomics training, it is thought to be beneficial in high-risk settings. One study suggested that training is inferior to a combination of other interventions in an office setting (363) and another found benefits for the neck, but not distal upper extremity. (386) Thus, other benefits of training may be possible. However, an RCT comparing wrist splinting with ergonomic education found splinting superior. (387) Thus, if there is a benefit, it may be modest, and it is suggested that such training should consist of quality information.

Figure 3. Split Keyboard

Reprinted with permission from David Rempel, MD.

Evidence for the Use of Ergonomic Interventions
There is 1 high-(365) and 5 moderate-quality (342, 362, 363, 366, 370) RCTs incorporated into this analysis. There are 4 low-quality RCTs (372, 388-390) in Appendix 2.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rempel 2012</td>
<td>Cluster RCT</td>
<td>8.0</td>
<td>N = 110 (100 females/10 males) dentists and dental hygienists. Mean±SD age: narrow handle 42.9±10.8 years; wide handle 46.6±9.8 years.</td>
<td>Heavy instrument, Narrow Handle (34g, 8mm diameter handle) (n = 56) vs. Light Instrument, Wide Handle (14g (curette tips and 11mm–diameter handle) (n = 54). Follow-up for 4 months.</td>
<td>Mean (SEM) adjusted score change shoulder pain: Heavy instrument 0.19 (0.16) vs. light instrument 0.52 (0.17); p = 0.02. Mean (SEM) adjusted score change wrist/hand pain: Heavy instrument 0.14 (0.17) vs. light 0.40 (0.18); p = 0.15.</td>
<td>&quot;To prevent or reduce arm pain, practitioners should consider using lightweight instruments with large diameters when performing scaling and root planning procedures.&quot;</td>
<td>Data suggest use of wider handled and lighter instrument associated with improved pain scores for distal upper extremity and shoulder.</td>
</tr>
<tr>
<td>Rempel 1999</td>
<td>RCT</td>
<td>7.5</td>
<td>N = 20 (13 females/7 males) with hand or wrist symptoms who used keyboard ≥10 hours per week. Mean age 42.6 years.</td>
<td>Keyboard A- Protouch keyboard, Key Tronic Corporation (n = 12) vs. Keyboard B-MacPro Plus keyboard with 2-ounce rubber domes, Key Tronic Corporation (n = 12). Both keyboards were of conventional layout (101 keys). Follow-up for 3 months.</td>
<td>Pain ratings significantly lower (p = 0.05) for keyboard A (6 weeks: 2.7 vs. 2.9; 12 weeks: 1.9 vs. 4.3).</td>
<td>&quot;We conclude that use of keyboard A for 12 weeks led to a reduction in hand pain and an improved physical examination finding when compared with keyboard B.&quot;</td>
<td></td>
</tr>
<tr>
<td>Rempel 2006</td>
<td>RCT</td>
<td>5.5</td>
<td>N = 182 (173 females and 8 males) customer service works who perform 20 hours or more of computer work per week. No neck, shoulder or upper extremity workers compensation claims. Mean Age was 40.02 years.</td>
<td>Ergonomic Training only: included conventional recommendations such as chair height and position (n = 46) vs Ergonomic training and trackball (n = 45) vs Ergonomic training and arm board-arm board is wraparound, padded arm support that attaches to top, front edge of work surface (n = 46) vs Ergonomic training and trackball and arm board (n = 45). Follow-up for 1 year.</td>
<td>Sixty-three (63) participants diagnosed with 1 or more incident MSDs. 12 month incidence rates for any upper body MSD by intervention group (47.7% vs. 35.7% vs. 29.5% vs. 31.8%). Adjusted hazard rate ratios for armboard for neck/shoulder disorders (HR = 0.49, 95% CI 0.24 to 0.97), reduced neck/shoulder pain (p = 0.01) and right upper extremity pain (p = 0.002).</td>
<td>&quot;Providing a large forearm support combined with ergonomic training is an effective intervention to prevent upper body musculoskeletal disorders and reduce upper body pain associated with computer work among call centre employees.&quot;</td>
<td>Small sample size. Keyboard associated with fewer symptoms required modestly greater force (0.71N vs. 0.58N) and greater displacement (1.69mm vs 0.58mm) to activate. Suggests lower typing force may not be helpful.</td>
</tr>
</tbody>
</table>

Copyright© 2016 Reed Group, Ltd.
<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Age</th>
<th>Design</th>
<th>Findings</th>
<th>Recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conlon 2008 RCT</td>
<td>5.0</td>
<td>42.87</td>
<td>Conventional Mouse Group (n = 52) vs. Alternative Mouse Group- neutral forearm posture (n = 52) vs. Board and conventional mouse-Forearm support board (n = 51) vs. Board and alternative mouse- Forearm support board (n = 52). Follow-up for 1 year.</td>
<td>No significant differences for use of an alternative mouse or use of forearm ergonomic support board vs. use of conventional mouse for both crude and adjusted hazard ratios (p&gt;0.05). Unadjusted model showed significant decrease in discomfort score in right upper extremity using forearm support board; -0.41 (-0.83 to -0.001) (p ≤0.05).</td>
<td>“In engineers who use a computer for more than 20 h per week, a forearm support board may reduce right upper extremity discomfort attributed to computer use.”</td>
</tr>
<tr>
<td>Gerr 2005 RCT</td>
<td>4.5</td>
<td>≥18</td>
<td>Group A: Alternate Intervention based on protective factors for both neck/shoulder and hand/arm (n = 122) vs. Group B: Conventional Intervention based on recommendations from OSHA, NIOSH and private industry (n = 125) vs. Group C-Control group, no intervention (n = 115). Follow-up for 6 months.</td>
<td>Among other differences, alternative involved greater elbow extension and keyboard position further recessed from edge of desk. No significant differences in distal upper extremity or neck/shoulder symptoms (p &gt;0.05).</td>
<td>“This study provides evidence that two specific workplace postural interventions are unlikely to reduce the risk of upper extremity musculoskeletal symptoms among computer users.”</td>
</tr>
<tr>
<td>Tittiranonda 1999 RCT</td>
<td>4.5</td>
<td>43.65</td>
<td>Placebo Group-Standard Keyboard (slope 8.0°) (n = 20) vs. Keyboard 1- Apple adjustable keyboard (slope 3.8 to 7.0°) (n = 20) vs. Keyboard 2- Comfort Keyboard System (slope -44.0 to 38.5°) (n=20) vs. Keyboard 3- Microsoft natural keyboard (slope 5.5 or -2.6°) (n = 20). Follow-up for 6 months.</td>
<td>High dropouts among keyboard that was completely split in two with sharply angled, but somewhat adjustable slopes. Changes in overall pain severity: placebo (-0.29±1.5) vs. split1 (0.52±2.0) vs. split/sharply angled (0.84±1.9) vs. split2 (1.21± 3.1), p = 0.11. More differences present in tendonitis subgroup (p = 0.088) than CTS (p = 0.57).</td>
<td>“These results provide evidence that keyboard users may experience a reduction in hand pain after several months of use of some alternative geometry keyboards.”</td>
</tr>
</tbody>
</table>

**Notes:**
- CTS and tendinitis were combined.
- Dropouts high in keyboard group with widely separated hands and more steeply angled surfaces.
- No meaningful differences in outcomes between conventional mouse and experimental mouse designs.
Return-To-Work Programs

Return-to-work programs have not been well studied among patients with hand, wrist, or forearm injuries (see Chronic Pain Guideline for discussion of principles). Several studies suggest that job physical demands, lack of job accommodation, and psychosocial conditions are the most important factors in predicting work disability.(391-393)

1. **Recommendation: Return-to-Work Programs for Subacute or Chronic Hand, Wrist, or Forearm MSDs**

   Return-to-work programs are recommended for treatment of subacute or chronic hand, wrist, or forearm MSDs, particularly patients with significant lost time.

   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – **Moderate**

2. **Recommendation: Return-to-Work Programs for Acute Hand, Wrist, or Forearm MSDs**

   Return-to-work programs are not recommended for treatment of acute hand, wrist, or forearm MSDs.

   Strength of Evidence – **Not Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – **Moderate**

**Rationale for Recommendations**

There are no quality studies that review the types of return-to-work programs typically found in the U.S. There is one quality study from Spain;(394) however, most patients had spine disorders and the program otherwise may have limited applicability due to longstanding, early active management of these issues in the U.S. These programs are thought to reduce morbidity and improve function. They are not invasive, have minimal potential for adverse effects, and are not costly. Return-to-work programs are recommended for management of select patients with hand, wrist, and forearm musculoskeletal disorders with lost time, and may be helpful for proactive emphases on functional recovery.

**Evidence for the Use of Return-To-Work Programs**

There is 1 moderate-quality RCT incorporated into this analysis.(394) There is one other study(395) in Appendix 2 (see Chronic Pain Guideline for additional studies).

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Compariso n Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abasolo 2007</td>
<td>RCT</td>
<td>4.0</td>
<td>N = 13,077 (gender not specified) workers on sick leave with diagnosis of MSD. Mean age for intervention and control groups: 40.8 and 40.6.</td>
<td>Multifaceted intervention program vs non-interventional control</td>
<td>Mean durations of temporary work disabilities for CTS patients (n = 74) 100.4 in controls vs. 27.8 days in intervention group (p &lt; 0.001).</td>
<td>“The implementation of this type of specialist-run, protocol-based early intervention program would be very beneficial in the treatment of patients with work disability related to MSDs, except for those with knee pain (excluding osteoarthritis).”</td>
<td>Scored for CTS patients within trial. Overall participation rate 62.8%.</td>
</tr>
</tbody>
</table>

**Work Activities**

Key factors to consider in disability duration are age and job activities. By communicating with patients and employers, physicians can make it clear that:

- Forceful repetitive grasping may increase forearm, hand, and wrist symptoms.
- Modified work and workplace activity guides may allow for recovery or time to (re)build activity tolerance through exercise.

Significant reductions in unnecessary lost work time can occur when the patient, physician, and employer work together to develop and apply modified work activities.(396-398, 399,400)
Carpal Tunnel Syndrome
Some physicians place work restrictions on patients with CTS; others do not. There is no quality evidence to suggest that restrictions are required.

Recommendation: Work Restrictions for CTS
For patients with CTS, it is recommended that their work be restricted to those tasks that do not involve high-force, stereotypical hand gripping or pinching or the use of high acceleration vibrating hand-held tools.

Indications – Select patients with combined forceful and repeated, stereotypical use of the hands or use of high amplitude vibrating tools. Of note, these types of jobs involve a minority of patients with CTS.

Indications for Discontinuation – Resolution, lack of improvement, or desire of the patient to remove limitations.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
There are no quality studies evaluating workplace restrictions; thus, whether patients improve more quickly with activity limitations has not been proven. However, based on available evidence associating combined forceful and repeated, stereotypical use of the hands or use of high amplitude vibrating tools with CTS, work restrictions are recommended for select patients with CTS. These types of jobs involve a minority of patients with CTS. Restrictions are not invasive, likely have few adverse effects, and may be moderate to high cost depending on length.

Evidence for Work Restrictions
There are 5 moderate-quality RCTs incorporated into this analysis. (342, 362, 363, 366, 370) There are 2 low-quality RCTs in Appendix 2.(389, 390)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: work restriction, ergonomics, carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, burning, tingling, itching, numbness, hand, palm, finger, pain controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 37 articles in PubMed, 609 in Scopus, 13 in CINAHL, and 45 in Cochrane Library. We considered for inclusion 3 from PubMed, 3 from Scopus, 1 from CINAHL, 0 from Cochrane Library and 6 from other sources. Of the 13 articles considered for inclusion, 7 randomized trials and 6 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rempel 1999</td>
<td>RCT</td>
<td>Sponsored by Northwest Trade Adjustment Assistance Center and by Key Tronic Corporation. No mention of COI.</td>
<td>7.5</td>
<td>N = 20 (13 females/7 males) with hand or wrist symptoms who used a keyboard ≥10 hours per week. Mean age 42.6 years.</td>
<td>Keyboard A- Protouch keyboard, Key Tronic Corporation (n = 12) vs. Keyboard B-MacPro Plus keyboard with 2-ounce rubber domes, Key Tronic Corp. (n = 12). Both keyboards conventional layout (101 keys). Follow-up for 3 months.</td>
<td>Pain ratings significantly lower (p = 0.05) for keyboard A (6 weeks: 2.7 vs. 2.9; 12 weeks: 1.9 vs. 4.3).</td>
<td>“We conclude that use of keyboard A for 12 weeks led to a reduction in hand pain and an improved physical examination finding when compared with keyboard B.”</td>
<td>Small sample size. Keyboard associated with fewer symptoms required modestly greater force (0.71N vs. 0.58N) and greater displacement (1.69mm vs 0.58mm) to activate. This suggests lower typing force may not be helpful.</td>
</tr>
<tr>
<td>Rempel 2006</td>
<td>RCT</td>
<td>Sponsored in part by grant from Centers for Disease Control/National Institutes for Occupational Safety and Health. COI: Dr Rempel has done consulting work for Logitech Corp., company which markets trackball tested in study.</td>
<td>5.5</td>
<td>N = 182 (173 females/8 males) customer service works who perform 20 hours or more of computer work per week. No neck, shoulder or upper extremity workers compensation claims. Mean age 40.02 years.</td>
<td>Ergonomic Training only: Included conventional recommendations such as chair height and position (n = 46) vs. Ergonomic training and trackball (n = 45) vs. Ergonomic training and arm board- arm board is a wraparound, padded arm support that attaches to the top, front edge of work surface (n = 46) vs. ergonomic training and trackball and arm board. Follow-up for 1 year.</td>
<td>Sixty-three (63) participants diagnosed with 1 or more incident MSDs. 12 month incidence rates for any upper body MSD by intervention group (47.7% vs. 35.7% vs. 29.5% vs. 31.8%). Adjusted hazard rate ratios for armboard for neck/shoulder disorders (HR = 0.49, 95% CI 0.24 to 0.97), reduced neck/shoulder pain (p = 0.01) and right upper extremity pain (p = 0.002).</td>
<td>“Providing a large forearm support combined with ergonomic training is an effective intervention to prevent upper body musculoskeletal disorders and reduce upper body pain associated with computer work among call centre employees.”</td>
<td>Dropout rate 31.3%. Return on investment estimated at 10.6 months.</td>
</tr>
<tr>
<td>Conlon 2008</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>5.0</td>
<td>N= 206 (57 females/149 males) engineers who worked at computer for at least 20 hours per week. Mean age 42.87 years.</td>
<td>Conventional Mouse Group- (n = 52) vs. Alternative Mouse Group- neutral forearm posture (n = 52) vs. Board and conventional mouse- Forearm support board (n = 51) vs. Board and alternative mouse- Forearm support board (n = 52). Follow-up for 1 year.</td>
<td>No significant differences for use of alternative mouse or forearm ergonomic support board vs. use of conventional mouse for crude and adjusted hazard ratios (p&gt;0.05). Unadjusted model showed significant decrease in discomfort score in right upper extremity using forearm support board; -0.41 (-0.83 to -0.001) (p ≤0.05).</td>
<td>“In engineers who use a computer for more than 20 h per week, a forearm support board may reduce right upper extremity discomfort attributed to computer use.”</td>
<td>No meaningful differences in outcomes between conventional mouse and experimental mouse designs.</td>
</tr>
<tr>
<td>Study</td>
<td>Title</td>
<td>Design</td>
<td>N</td>
<td>Description</td>
<td>Results</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>-------</td>
<td>--------</td>
<td>---</td>
<td>-------------</td>
<td>---------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gerr 2005</td>
<td>RCT</td>
<td>Sponsored by US National Institute</td>
<td>4.5</td>
<td>N = 362 (279 female/83 male) workers who operated a computer at least 15 hours or more per week. Age ≥18 years. Group A: Alternate Intervention - based on protective factors for both neck/shoulder and hand/arm (n = 122) vs. Group B: Conventional Intervention based on recommendations from OSHA, NIOSH and private industry (n = 125) vs. Group C: Control group, no intervention (n = 115). Follow-up for 6 months. Among other differences, alternative involved greater elbow extension and keyboard position further recessed from edge of desk. No significant differences in distal upper extremity or neck/shoulder symptoms (p&gt;0.05).</td>
<td>“This study provides evidence that two specific workplace postural interventions are unlikely to reduce the risk of upper extremity musculoskeletal symptoms among computer users.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tittiranonda 1999</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>4.5</td>
<td>N = 80 (46 female/34 male) with CTS syndrome and/or tendonitis. Mean age 43.65 years. Placebo Group - Standard Keyboard (slope 8.0°) (n = 20) vs. Keyboard 1: Apple adjustable keyboard (slope 3.8-7.0°) (n = 20) vs. Keyboard 2: Comfort Keyboard System (slope -44.0-38.5°) (n = 20) vs. Keyboard 3: Microsoft natural keyboard (slope 5.5 or -2.6°) (n = 20). Follow-up for 6 months. High dropouts among keyboard that was completely split in two with sharply angled, but somewhat adjustable slopes. Changes in overall pain severity: placebo (-0.29±1.5) vs. split1 (0.52±2.0) vs. split/sharply angled (0.84±1.9) vs. split2 (1.21±3.1), p = 0.11. More differences present in tendonitis subgroup (p = 0.088) than CTS (p = 0.57).</td>
<td>“These results provide evidence that keyboard users may experience a reduction in hand pain after several months of use of some alternative geometry keyboards.”</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*CTS and tendinitis were combined. Dropouts were high in the keyboard group with widely separated hands and more steeply angled surfaces.*
Triangular Fibrocartilage Complex (TFCC) Tears
TFCC tears may not require work limitations. However, the more forceful the work and more significant the symptoms, the more likely work limitations will be needed. Work limitations typically include reducing forceful use, wrist rotation, or other activities that provoke symptoms.

Crush Injuries and Compartment Syndrome
These injuries generally require work limitations depending on task demands. More severe cases require time away from work for recovery from surgery, pain management, and generally require a gradual resumption of usual activities dependent on injury severity and rate of healing.

Kienböck Disease
There is no evidence that work restrictions are helpful, yet as the condition often progresses, patients typically incur increasing degrees of disability with a progressive need for work limitations. Advanced cases generally require temporary removal from work and surgery, with return to work post-operatively. Post-operative limitations are generally based on a combination of the clinical results (i.e., severity of pain and symptoms) and work demands. Patients with light to medium work may require no limitations, while those with medium to heavy work, particularly with post-operative pain may require significant limitations.

Wrist Sprains
This injury may or may not require work limitations depending on task demands. However, moderate to severe wrist sprains likely necessitate splinting and limitations.

Mallet Finger
This injury requires splinting; however, whether there is any need for work limitations involving the digit other than a requirement to wear the splint continuously is unclear. Provided there is no difficulty with wearing the splint, no work limitations are generally needed.

Flexor Tendon Entrapment (Tenosynovitis and Trigger Digit)
Whether work limitations are indicated or helpful is unknown, but may be reasonable for select cases, particularly where contributions from physical factors are more probable such as localized compression from sharp objects or tools.

Extensor Compartment Tenosynovitis (Including de Quervain’s Stenosing Tenosynovitis and Intersection Syndrome)
Job modifications are thought to be needed in most of these work-related cases to facilitate recovery.

Recommendation: Modification of Work Activities for Extensor Compartment Tenosynovitis
Removal from job tasks thought to have caused extensor compartment tenosynovitis is recommended.

Indications – Patients with combined forceful and repeated use of the hands or combined use with substantially non-neutral wrist postures.
Indications for Discontinuation – Resolution, lack of improvement, or desire of the patient to remove limitations.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There are no quality studies evaluating the modification of work activities for extensor compartment tenosynovitis. However, where occupational factors are significant, a trial of removal from that type of work may be indicated.
Ulnar Nerve Entrapment at the Wrist (including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome)

Job modifications are thought to be needed in some cases to facilitate recovery.

**Recommendation: Modification of Work Activities for Ulnar Neuropathy**

**Removal from job tasks thought to have caused ulnar neuropathy at the wrist is recommended.**

**Indications** – Patients with forceful use of the hand, especially use of the hand as a hammer with striking of the hypothenar area and development of hypothenar hammer syndrome.

**Indications for Discontinuation** – Resolution, lack of improvement, or desire of the patient to remove limitations.

  - **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
  - **Level of Confidence** – Low

**Rationale for Recommendation**

There are no quality studies evaluating the modification of work activities for ulnar neuropathies at the wrist. However, where occupational factors are significant, especially for patients with hypothenar hammer syndrome, a trial of removal from that type of work may be indicated.

Radial Nerve Entrapment

Job modifications are thought to be needed in a few cases to facilitate recovery.

**Recommendation: Modification of Work Activities**

**Removal from job tasks thought to have caused radial neuropathy at the wrist is recommended.**

**Indications** – Patients with radial neuropathies thought to be caused by an ongoing job physical exposure (e.g., striking the radial nerve).

**Indications for Discontinuation** – Resolution, lack of improvement, or desire of the patient to remove limitations.

  - **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
  - **Level of Confidence** – Low

**Rationale for Recommendation**

There are no quality studies evaluating the modification of work activities for radial neuropathies at the wrist. However, where occupational factors are significant, a trial of removal from that type of work may be indicated.

**Non-Specific Hand/Wrist/Forearm Pain**

Non-specific pain may or may not require work limitations depending on task demands. For patients with high exposures, work limitations are more likely to be helpful. However, in the absence of high force or high force combined with other ergonomic factors, work limitations are at times counterproductive because they enforce debility and do not produce meaningful improvements. In those settings, work limitations may be trialed; however, in the absence of improvement, resumption of regular work activities may be helpful for long-term functional gain.

**Scaphoid Fracture**

Activities should be modified to allow for the splinting and immobilization of the carpal bones. In a moderate-quality study comparing surgical fixation to non-operative treatment,(401) the mean range of time for Scottish patients with non-displaced scaphoid fracture to return to normal daily activities living with non-operative treatment was 1 week for dressing, 1 week for washing, 2.8 weeks for shopping, and 2.7 weeks for housework. The mean time for returning to full employment was 11.4 weeks, and to full sports 15.5 weeks. The mean return time of the operative group was not statistically different except to full employment and full sports, which were 3.8 weeks and 6.4 weeks respectively.(401) While operative fixation of non-displaced scaphoid fracture may reduce short term disability, there is a reported 11-fold increased risk of scaphotrapezial arthritis compared with those non-operatively treated.(402)
Distal Phalanx Fractures and Subungual Hematoma

All work activities that can be accomplished while wearing a finger splint are appropriate. Athletes may return to sports after the initial swelling and pain have resolved, approximately 7 to 10 days. Activities requiring full distal joint mobility and forceful use may be delayed as long as 4 to 6 weeks. Residual tenderness may be present for up to 6 months. (78)

Middle and Proximal Phalangeal and Metacarpal Fractures

Activities restrictions should provide for immobilization of affected finger or hand, but otherwise activities should be allowed.

Distal Forearm Fractures

Functional restrictions of the affected extremity are limited by immobilization technique. Activities should be modified to allow for splinting and immobilization of the forearm. Return to work will likely be influenced by the patient and provider’s subjective assessment of disability and perception of job difficulty. It may be helpful to refer the patient to an occupational therapist to address the appropriate activity modification, compensatory strategies, adaptive equipment, and environmental modification throughout the period of the patient’s recovery and rehabilitation.

Ganglion Cyst

There is no indication for limiting work activity except for ganglia that are causing significant pain, as there is no reported strong association between activity and exacerbation or causation of ganglia. Those with considerable pain may require limitations to avoid activities provoking increased symptoms, most typically involving forceful use.

Hand-Arm Vibration Syndrome (HAVS)

1. Recommendation: Vibration Exposure Work Restrictions for HAVS

   For patients with HAVS, it is recommended that their work be restricted to those tasks that do not involve high-amplitude, low-frequency vibration exposures from hand-held tools.

   Indications – HAVS from high-amplitude, low-frequency vibration exposures through vibrating hand-held tools.

   Indications for Discontinuation – Resolution or desire of the patient to remove limitations. If the exposure(s) are confirmed and the clinical findings are significant, re-exposure is not believed to be indicated.

   Strength of Evidence – Recommended, Insufficient Evidence (I)

   Level of Confidence – Moderate

2. Recommendation: Cold Exposure Work Restrictions for HAVS

   For select patients with HAVS, it is recommended that their work be restricted to those tasks that do not involve cold exposures.

   Indications – HAVS that is not controlled through avoidance of vibration exposures, or patients having recurring problems with vasospasm or other complications that are unresolved with other treatments.

   Indications for Discontinuation – Resolution or desire of the patient to remove limitations.

   Strength of Evidence – Recommended, Insufficient Evidence (I)

   Level of Confidence – Moderate

Rationale for Recommendations

Limitation of exposure to total vibration dose should be achieved particularly by limiting the duration and frequency to high-amplitude, low-frequency vibration. Reducing transmission of vibration through isolation and damping techniques may also be attempted, although in a patient with established HAVS, avoidance is generally preferable. Avoidance of cold temperatures that provoke symptoms or wearing gloves if sufficient to control symptoms is warranted. (127) Anti-vibration gloves are sometimes utilized.
Recognition and reduction of other ergonomic factors including repeated and sustained exertion, forceful exertions, contact stress, and stressful postures may be helpful.

**Laceration Management**
There are no quality studies on return to work and restrictions for upper extremity laceration repair. Movement of injured body parts is thought to promote earlier recovery and minimize disability. Most patients should be able to return to work with appropriate task specific restrictions while the wound is healing. Accommodation for prescribed medications, elevation, splinting and modalities such as use of heat or ice may be necessary. While there is no quality evidence for any of these modalities, keeping the wound dry for the first few days, splinting, elevation, and heat or ice are simple techniques that are believed to be helpful. Splinting is generally limited to extensor surface lacerations that cross a joint and involve sufficient tension to pull wound edges apart.\(^{(403)}\)

**Human and Animal Bites**
Work activities are expected to be minimally impacted except for limitations related to treatment of laceration or infection.

**Hand/Finger Osteoarthrosis**
Hand osteoarthrosis generally requires no work limitations. When the disease progresses to moderate or severe disease, work limitations may be required due to the impairment and or pain.

**Initial Care**
Initial treatment should generally be guided by implementing the strongest evidence-based recommendations that are considered first-line interventions. Exceptions include those treatments that are accepted as best practices, but have not been subjected to RCTs or crossover trials (e.g., antibiotics for diabetics with “dirty” lacerations). Careful consideration of the indications and limitations described in the full text for each recommendation is critical to understanding the best application for each intervention. If treatment response is inadequate (i.e., if symptoms and activity limitations continue), second- and third-line recommendations may be considered. Physicians should consider the possibilities of diagnosed and previously undiagnosed medical diseases such as diabetes mellitus, hypothyroidism, and various arthritides. Adverse effects, cost, and provider and patient preferences should be considerations in guiding the choice of recommendations. Part of the initial treatment plan for all disorders should include patient education. For most diagnoses, this is critical to successful treatment.

**Recommendation: Education for Hand, Wrist, or Forearm Disorders**

**Education is recommended for select patients with hand, wrist, or forearm disorders.**

**Frequency** – One or 2 appointments for educational purposes. Additional appointments may be needed if education is combined with occupational or physical therapy treatments. Follow-up educational visit(s) for more severe disorders as part of a progression towards normal functional use is sometimes helpful.

- **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
- **Level of Confidence** – **Low**

**Rationale for Recommendation**
There are no quality studies specifically evaluating efficacy of patient education for utility or necessity in treatment of hand, wrist, or forearm disorders. Yet, for many disorders (e.g., criticality of maintaining splinting of mallet finger, cast management, monitoring for signs of infection) education appears essential. Some physicians accomplish this in the course of extended patient visits, while others routinely refer patients to an occupational or physical therapist for education. Regardless of the approach, a few appointments for educational purposes are recommended for select patients. The number of appointments is dependent on the diagnosis, severity of the condition, and co-existing conditions. Although education is usually incorporated as part of the overall treatment plan, an additional 1 or 2 appointments for purely educational purposes may be helpful midway through a treatment course for the more severely affected patient. In addition, education is low cost and thus is recommended.
**Follow-up Visits**

Patients with potentially work-related hand, wrist, and forearm symptoms should generally have a follow-up visit approximately every 3 (severe disorders) to 7 days (typical disorder severity) to monitor function, medication use and/or a physical or occupational therapist visit for counseling regarding contributing physical factor avoidance (e.g., reducing force, avoiding static positions), sleep posture, and other concerns. More frequent follow-up is usually required for patients who are not working. Care should be taken to answer questions and make these sessions interactive so that the patient is involved in his or her recovery including identifying potential barriers to recovery and return to normal function and work. More specific guidance for follow-up visits may be included in the discussion of each disorder topic.

**Amputations and Indications for Replantation**

The decision for amputation or replantation should be made by a physician who has training and experience in treating amputations and replantations. The key for the initial physician or health care provider is to reduce the warm ischemia time of the amputated part – the time without any preparation of the amputated part. This is best done by washing the amputated part in saline and wrapping it in saline soaked gauze, putting it into a plastic bag if possible, and then placing it onto cardboard that is laid over ice in a cooler or jug. The part of the body where the amputation has occurred should be covered with a compression dressing. Vascular control is important. Attempts to use clamps to control bleeding often damage the neurovascular structures and should not be used. Indicators that are used to suggest replantation success include thumb amputation, multiple digit amputations, amputation at a metacarpal amputation, almost any body part amputated in a child, wrist or forearm amputation, and individual digit amputated distal to flexor digitorum superficialis (FDS) insertion.

Contraindications may include ring avulsion injuries, severely crushed or mangled parts, amputations at multiple levels, amputations in patients with other serious injuries or diseases, arteriosclerotic vessels, mentally unstable patients, distal amputations (finger tip injuries), individual finger in adult proximal to the FDS insertion and prolonged warm ischemia. Prolonged warm ischemia is defined as more than 6 hours for proximal replantations (wrist), and 12 hours for digits, although some physicians will attempt replantation after 6 hours of warm ischemia, and 24 to 30 hours ischemia time (time from amputation until replant with the digit stored in cool container as described above) for digital replantations.

**Carpal Tunnel Syndrome (CTS)**

**Diagnostic Criteria**

Patients with a presumptive diagnosis of CTS should have both paresthesias in the median nerve distribution and symptoms that are either nocturnal or provoked. Paresthesias are tingling or numbness in a median nerve distribution, (vibrotactile testing has been utilized to attempt to objectify sensory findings, but appears to not perform particularly well)(404) generally involving at least two median nerve-served digits (they may also have pain – pain is not the primary symptom, there is also some evidence for more non-neurologic pain in workers’ compensation patients(405) – or burning in a median nerve distribution, but should have paresthesias); and 2) symptoms that are provoked either nocturnally or with sustained grasp (e.g., holding a tool, steering wheel or newspaper).

Patients with a confirmed diagnosis of CTS should have both symptoms as with a presumptive diagnosis above, and either: 1) a confirmatory electrophysiological study (EDS) interpreted as consistent with CTS, or 2) largely or completely resolved symptoms with injection of a glucocorticosteroid.

The differential diagnosis for carpal tunnel syndrome (CTS) particularly includes pronator syndrome; C6 and C7 cervical radiculopathies; and other neurological entrapments located between the spinal cord and median nerve in the carpal canal including thoracic outlet syndrome, diabetic neuropathy, neuropathy from alcohol, other systemic neuropathies, stroke, other cerebrovascular events, and central nervous system tumors. Most other causes may be eliminated or the probability reduced by conducting a careful history, physical exam, or focused testing.
Special Studies and Diagnostic and Treatment Considerations

ELECTRODIAGNOSTIC STUDIES

Appropriate electrodiagnostic studies (EDS), including nerve conduction studies (NCS), may help differentiate CTS from other conditions such as cervical radiculopathy, other median nerve neuropathies, or ulnar neuropathy at the elbow.\(10, 177, 180, 183, 184, 406-448\) In select or more difficult cases, especially if cervical radiculopathy is a concern, electromyography (EMG) studies should be incorporated.\(406\) NCS and EMG may be normal particularly in some mild cases of CTS. If EDS are negative, tests may be repeated later in the course of treatment if symptoms persist. It is also important to recognize that electrodiagnostic studies are abnormal in a large proportion of patients who are without symptoms and thus without CTS.\(171\) Thus, EDS testing in a patient with a low pre-test probability of CTS may result in inappropriate diagnosis of CTS. EDS has been purportedly not useful in diagnosing clear-cut CTS cases.\(449\)

The American Association of Electrodiagnostic Medicine, the American Academy of Neurology, and the American Academy of Physical Medicine and Rehabilitation jointly published a practice parameter for electrodiagnostic studies in CTS.\(406\) However, the quality of EDS varies widely in practice\(450\) and this practice guideline is sometimes not adhered to, requiring the treating physician to be familiar with these issues to better interpret the findings in a clinical context. Additionally, cut-off points for abnormal values have yet to be fully standardized and the correlations between symptom severity and EDS severity are not uniformly strong.\(11, 167\) In patients with suspected CTS where electrodiagnostic confirmation would alter treatment plans, the following EDS studies are recommended (in the majority of these studies, the hand temperature should be \(32^\circ\text{C or warmer}\)):\(406\)

1. Perform a median sensory NCS across the wrist with a conduction distance of 13 to 14cm. If the result is abnormal, compare the result of the median sensory NCS to the result of a sensory NCS of one other adjacent sensory nerve in the symptomatic limb.
2. If the initial median sensory NCS across the wrist has a conduction distance greater than 8cm and the result is normal, one of the following additional studies is recommended:
   a. Comparison of median-sensory- or mixed-nerve conduction across the wrist over a short (7 to 8cm) conduction distance to the ulnar sensory-nerve conduction across the wrist over the identical 7 to 8cm conduction distance, or
   b. Comparison of median sensory across the wrist with ipsilateral radial or ulnar sensory conduction across the wrist, or
   c. Comparison of median sensory or mixed nerve conduction through the carpal tunnel to sensory or mixed NCS of proximal or distal segments of the ipsilateral median nerve.
3. Motor conduction study of the median nerve recording from the thenar muscle and of one other ipsilateral nerve with distal latency.
4. Optional comparisons may include ipsilateral median-ulnar motor nerve distal latencies and median-ulnar motor conduction differences.
5. Needle EMG is optional as it is primarily used for evaluation of cervical radiculopathy.\(406\)

1. **Recommendation: EDS for Diagnosis and Pre-operative Assessment of CTS**
   Quality EDS (see above) are recommended to assist in securing a firm diagnosis for those patients without a clear diagnosis of CTS. EDS are also recommended to objectively secure a diagnosis of CTS prior to surgical release in workers compensation patients.\(451\) If EDS is elected, in most cases of CTS, only the sensory and motor conduction studies are necessary for diagnosis confirmation. If the examination is more complex, then addition of the EMG component of the EDS should be obtained.

   **Strength of Evidence** – **Recommended, Evidence (C)**

   **Level of Confidence** – **Moderate**

2. **Recommendation: EDS for Initial Evaluation of CTS Patients**
   EDS is not recommended for initial evaluation of most CTS patients with a confirming history and clinical signs as it does not change the management of the condition. EDS is also not
recommended prior to glucocorticosteroid injection as a good history and clinical suspicion is believed to be sufficient to warrant the intervention which would not likely be altered by EDS.

Strength of Evidence – Not Recommended, Insufficient Evidence (I)
Level of Confidence – High

3. Recommendation: Commercial Products for Performing EDS for CTS Patients

Automated devices to accomplish EDS is recommended for highly selected CTS patients.(452, 453)

Indications – EDS that is not readily obtainable by an American Board of Electrodiagnostic Medicine-certified electrodiagnostician (e.g., geographic distance). There also should be no concern about other potentially confounding conditions such as cervical radiculopathy.

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – Low

Rationale for Recommendations
EDS are the only unequivocally objective measures of median nerve function.(10, 406, 451, 454-456) However, there are both false-positive and false-negative test results that demand that the physician understand the pre-test probabilities and be capable of interpreting the results and placing them in an appropriate clinical context. For example, EDS should not be ordered in settings where the clinical history suggests a low likelihood of CTS because the probability of a false-positive test result may be well above 50%. EDS are primarily of assistance in: 1) identifying an anatomic location of nerve conduction slowing; 2) identifying objective evidence for alternate diagnostic considerations (e.g., cervical radiculopathy); and 3) quantifying nerve function to assure the physician that an operative state such as CTS is present. EDS are not invasive or minimally invasive (depending on whether the EMG component is required), have minimal adverse effects, and are high cost. They are recommended for evaluation of select cases, especially if the diagnosis is unclear or surgery is planned.

There are other commercial diagnostic products;(452, 453, 457, 458) and some studies have suggested there may have sufficient accuracy,(445, 452) however, there are relatively few studies available and thus the use of these studies may be currently limited to where there is both no concern about radiculopathy and other disorders and the EDS test is not readily available (e.g., due to distance geographical issues).

Evidence for the Use of Electrodiagnostic Studies
There are 20 moderate-quality studies incorporated into this analysis.(319, 445, 451-453, 455, 456, 459-471) There are 4 low-quality studies in Appendix 2.(472-475)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: electrodiagnostic studies, nerve conduction study (NCS), electromyography (EMG); carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 96 articles in PubMed, 371 in Scopus, 23 in CINAHL, and 23 in Cochrane Library. We considered for inclusion 20 from PubMed, 30 from Scopus, 5 from CINAHL, 6 from Cochrane Library and 30 from other sources. Of the 91 articles considered for inclusion, 67 trials and 7 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Population/ Case Definition</th>
<th>Investigative Test</th>
<th>Gold Standard / Comparative Test</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dale 2015 Diagnostic</td>
<td>7.0</td>
<td>N = 62 (19 females and 43 males) subjects that originally underwent NC-Stat automated NCS; mean age 33.66 (9.43).</td>
<td>NC-Stat an automated Nerve Conduction Studies (NCS) machine</td>
<td>Traditional NCS using a NeuroMax 1002 device in an electrodiagnostic lab.</td>
<td>Higher agreement between Median nerve parameter rather than Ulnar nerve parameter. Highest reciever operating curve (ROC) area 0.97 and 0.96 for median nerve parameter. 100% sensitivity and 74% specificity for Ulnar Distal Motor latency and Distal sensory latency. Highest specificity in median ulnar sensory difference, 100%.</td>
<td>“In conclusion, the NC-stat device has been previously shown to have excellent agreement with traditional methods of median nerve testing in clinical populations; this study shows that this excellent agreement extends to use in a general worker population with low prevalence of disease.”</td>
<td>Study reports automated nerve conduction study was comparable to the traditional EDS for detection of median nerve conduction abnormalities in a general worker population.</td>
</tr>
<tr>
<td>Buch-Jaeger 1994 Diagnostic</td>
<td>7.0</td>
<td>N = 112 with signs of carpal tunnel, 60 bilaterally. Patients confirmed through clinical analysis. The mean age of 52 years, ranging from 29-81 years.</td>
<td>Nerve Conduction studies (NCS), positive when distal motor latency in the abductor brevis muscle was greater than 4ms.</td>
<td>Clinical evaluation focusing on 11 different criteria including paraesthesiae in territory of median nerve, occasional pain, nocturnal recrudescence of symptoms, numbness leading to clumsiness of hand, Phalen’s test, Tinel’s test, dealt, Vibratory sensibility, Thershold sensibility, Giliat’s test, McMurhtry’s sign, Static 2-point discrimination.</td>
<td>NCS positive in 68 cases (61%) and negative in remaining 44 cases (39%). Of negative NCS patients, 10 spontaneously recovered, 4 unchanged, 5 had symptoms after heavy tasks, 3 thought to be malingering, and 20 diagnosed with other disease. Of NCS confirmed CTS group 33 had surgical findings; 40 (93%) had complete disappearance and intervention.</td>
<td>“Our findings suggest that typical clinical features and positive provocation tests are not sufficient to lead a surgeon to decompress the carpal tunnel, and we feel that electrodiagnostic examination is necessary in every case.”</td>
<td>Study supports nerve conduction studies to be a key component in diagnosis of CTS as other clinical tests have fair sensitivity and specificity.</td>
</tr>
<tr>
<td>Year</td>
<td>Study Title</td>
<td>Methods</td>
<td>Results</td>
<td>Conclusion</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>------</td>
<td>-------------</td>
<td>---------</td>
<td>---------</td>
<td>------------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2003</td>
<td>Atroshi</td>
<td>N = 125 (gender not specified) CTS group and symptomatic controls with possible/unlikely CTS (n = 155) and asymptomatic Control group (n = 124) no signs of CTS (n = 124) Mean age 51±14. All participants collected from 3,000 sample in Sweden. Mean age 52 ± 13. Bilateral Nerve Conduction Tests including median nerve distal motor latency (M) DML. Long Finger-wrist sensory latency, and sensory conduction velocity (SCNV) in forearm, wrist-Palm, and palm digit segments. Also an ulnar nerve small finger-wrist sensory latency. Patients clinically diagnosed using Phalen’s Test, Tinel’s Test, recurrent numbness or tingling, and filled out a hand diagram.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2008</td>
<td>Leffler</td>
<td>N = 75 symptomatic hands referred to electrophysiological lab; Mean age 49 ± 12 vs. n = 22 asymptomatic volunteers. An automated electro diagnostic device (AEND). A conventional diagnostic device conducted within a lab by a neurologists.</td>
<td>Linear regression showing AEND and conventional results correlation was 0.90 (p &lt;0.001). AEND sensitivity for very symptomatic hands 89% specificity 90%. Lower severe had sensitivity of 87%, also 90% specificity.</td>
<td>“This study demonstrated that the Distal Motor latency provided by an AEND is highly correlated with the Distal Motor Latency obtained by conventional testing.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2008</td>
<td>Graham</td>
<td>N = 143 clinically diagnosed with with CTS Standard electrodiagnosti c tests, Sensory nerve conduction by technician and evaluated by neurologist, use of stringent and Lax criteria used to confirm CTS. CTS-6 evaluation which is a clinical diagnosis aid. Using CTS-6 the pretest probability was 0.81 ± 0.22. After the Stringent Criteria posttest probability was 0.91 and Lax was 0.83. Average change in probability was -0.02 ± 0.10 with stringent and -0.06 ± 0.16 with lax.</td>
<td>“For the majority of patients who are considered to have carpal tunnel syndrome on the basis of their history and physical examination alone, electrodiagnostic tests do not change the probability of diagnosing this condition to an extent that is clinically relevant.”</td>
<td>“Using the clinical diagnosis of CTS as the criterion standard, nerve conduction tests had moderate sensitivity and specificity and a low positive predictive value in population-based CTS. Measurement of median-ulnar sensory latency difference had the highest diagnostic accuracy.”</td>
<td>Study suggests nerve conduction study to diagnose CTS had only modest sensitivity and specificity and measuring the median-ulnar sensory latency difference was a better predictor of true CTS diagnosis.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study Year</td>
<td>Diagnostic Type</td>
<td>Sample Size</td>
<td>Methodology</td>
<td>Results</td>
<td>Findings</td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------</td>
<td>-----------------</td>
<td>-------------</td>
<td>-------------</td>
<td>---------</td>
<td>----------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pastare 2009</td>
<td>Diagnostic</td>
<td>66</td>
<td>Nerve Conduction Studies vs. Ultrasound</td>
<td>Clinical Diagnosis of CTS</td>
<td>Nerve Conduction studies showed greater diagnostic sensitivity than ultrasound; 54 wrists 82% vs. 41 62% for highly likely clinical diagnosis of CTS.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nathan 1993</td>
<td>Diagnostic</td>
<td>2,334</td>
<td>Maximum latency difference (MLD) determined by centimetric technique.</td>
<td>Clinical diagnosis of CTS. MLD was compared with 8-cm latency (S8) and 14-cm latency (S14).</td>
<td>MLD most sensitive measurement (86%) and had greatest efficiency of correct classification (84%). The S14 was most specific measurement (94%).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lee 2009</td>
<td>Diagnostic</td>
<td>153</td>
<td>Electrodiagnost ic testing including Median Terminal latency differences, motor conduction study and sensory conduction study.</td>
<td>Clinical criteria and diagnosis was used as the parameter to test for sensitivity.</td>
<td>Sensitivity of top EDX testing: Wrist-Palm Sensory Conduction Velocity (SCV): 90.5%, Distal-Proximal ratio SCV 92.3%, Wrist-Digit 2 SCV 89.1%, Wrist-Digit 3 89.1%. Terminal Latency ratio of Wrist-Palm Motor conduction 81.8%.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

“Based on these findings, we recommend that confirmatory nerve conduction studies be performed in all cases where CTS is suspected.”

“Reports nerve conduction studies superior to sonography in detecting CTS. But, sonography may be used as first-line screening tool if clinical index of suspicion for CTS is high.”

“Controls younger than CTS group. Study reports maximum latency difference (MLD) most reliable measurement for predicting CTS. Study recommends nerve conduction studies be performed when high index of suspicion for CTS.”

“The terminal latency ratio of the wrist to the palm is a valuable technique for the diagnosis of carpal tunnel syndrome, and it requires only a simple additional stimulus compared to existing methods.”

Study suggests median terminal latency ratio in the third finger as the most sensitive technique for detection of CTS.”
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Procedures</th>
<th>Findings</th>
<th>Observations</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concannon 1997 Diagnostic</td>
<td>6.0</td>
<td>N = 349 (460 hands) patients who underwent carpal tunnel release.</td>
<td>Electrodiagnostic Studies</td>
<td>N/A</td>
<td>398/460 hands had positive electrodiagnostic studies. 60 clinical CTS diagnosis but normal electro-diagnostic studies. Phalen's only significant test with regression coefficient: -0.91; OR 0.40 CI: 0.17 – 0.95 (p = 0.04). Indicated model predicts higher probability of negative electromyogram than positive electromyogram. 76% (n = 348) of affected hands had mild to moderate electrodiagnostic findings, 11% had severe CTS (n = 50), and 13% had normal electrodiagnostic findings. Patients who were older tended to have severe electrodiagnostic findings (p = 0.0001). Significant association between gender and maximal electrodiagnostic findings (p = 0.02). Patients with severe CTS had highest incidence of muscle wasting (22%, p &lt;0.02).</td>
<td>“Electrodiagnostic studies in suspected carpal tunnel syndrome should be reserved for use in the patient with equivocal findings and should not be considered a necessary criterion when history and clinical examination provide this diagnosis.”</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>N</td>
<td>Suspected CTS Patients and Controls</td>
<td>Median Wrist-Palm Motor Conduction Velocity (W–P MCV)</td>
<td>Standard Sensory Conduction Techniques</td>
<td>Abnormal hand number, sensitivity (%), and specificity (%) of Motor DL/Sensory DL (D1)/Sensory DL (D2)/Sensory DL (D4)/W–P MCV/W–P SCV/W–P SCT/median–radial sensory latency difference/median–ulnar sensory latency difference were: 234, 65, and 99.3/289, 80.3 and 98.7/261, 72.5 and 99.3/276, 76.7 and 100/294, 81.7 and 100/265, 73.6, and 100/291, 80.8 and 100/312, 86.7 and 98.7/314, 87.2 and 96.7</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Chang 2006 Diagnostic</td>
<td>6.0</td>
<td>N = 280 suspected CTS patients (360 hands).</td>
<td>Median wrist–palm motor conduction velocity (W–P MCV)</td>
<td>Standard sensory conduction techniques</td>
<td>Abnormal hand number, sensitivity (%), and specificity (%) of Motor DL/Sensory DL (D1)/Sensory DL (D2)/Sensory DL (D4)/W–P MCV/W–P SCV/W–P SCT/median–radial sensory latency difference/median–ulnar sensory latency difference were: 234, 65, and 99.3/289, 80.3 and 98.7/261, 72.5 and 99.3/276, 76.7 and 100/294, 81.7 and 100/265, 73.6, and 100/291, 80.8 and 100/312, 86.7 and 98.7/314, 87.2 and 96.7</td>
<td>“W–P MCV is a valuable motor conduction technique for the diagnosis of CTS and it is confirmed again that W–P MCV is equal to or more sensitive than W–P SCV and W–P SCT.”</td>
</tr>
<tr>
<td>Wang 2013 Diagnostic</td>
<td>6.0</td>
<td>N = 162 CTS patients (248 hands) and 83 controls (166 hands).</td>
<td>Median-to-ulnar comparative Nerve conduction studies: Sensory median-ulnar difference (MS-US), Mixed median-ulnar palm latency difference (PM-PU), and Distal latency differences between second lumbrical and interossei (2L-INT).</td>
<td>N/A</td>
<td>168/248 (67.7%) hands had abnormal findings. 80 (32.3%) hands received 2L-INT, MS-US, and PM-PU additional tests. 88.3% symptomatic hands had at least an abnormal findings. The sensitivity of MS-US/2L-INT/PM-PU were: &gt;0.5 ms in 21.3% of hands/ &gt;0.4 ms in 27.5% of hands/ &gt;0.4 ms in 47.5% of hands. MP-UP had the greatest sensitivity in contrast to L2-INT and MS-US (p = 0.014 and p&lt;0.001). Conventional EDX with PM-PU had a sensitivity of 83%.</td>
<td>“For CTS patients with normal results from the standard methods, PM-PU is a good additional comparative test to further improve diagnostic rate.”</td>
</tr>
<tr>
<td>Study</td>
<td>Age</td>
<td>Description</td>
<td>Nerve Conduction Studies</td>
<td>Normal Control Group Values</td>
<td>Average Sensitivity of Different Segment Lengths: Long Segment 39.5%. Short Segment 56%. Two Segment 40.5%</td>
<td>The results obtained in this study demonstrate that patients with CTS form a heterogeneous group with a wide variation in a specific nerve conduction parameter between individual patients, reflecting the different degrees of nerve pathology. It is therefore not recommended to use a specific procedure for the evaluation of each patient suspected of CTS, but (i) to use only sensitive parameters with high specificity as an optimal routine for the investigation of the average CTS patient, (ii) to perform needle EMG in all suspected CTS patients.</td>
</tr>
<tr>
<td>---------------</td>
<td>-------</td>
<td>------------------------------------------------------------------------------</td>
<td>--------------------------</td>
<td>-----------------------------</td>
<td>------------------------------------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Lew 2005</td>
<td>5.5</td>
<td>control healthy hands; Mean Age 44.0 ± 12.9 (n = 44) vs. symptomatic hands suspected of CTS; Mean Age 51.5 ± 18.2 (n = 136).</td>
<td>Nerve Conduction Studies varying in segment length. Sensory Nerve conduction velocity of Long segment from wrist to Digit 1, 2, 3, and 4. Transcarpal mixed nerve conduction velocity of Short segment palm to wrist. Transcarpal sensory Nerve Conduction Velocity wrist-digit and palm to digit difference.</td>
<td>Nerve Conduction Study (NCS) results from control group.</td>
<td>Average Sensitivity of the different segment lengths: Long segment 39.5%. Short segment 56%. Two segment 40.5%</td>
<td>“Our Study showed that among the 8 median NCV tests, the short, segment, onset latency-based transcarpal NCV was most sensitive in diagnosing CTS. This study also suggests that direct measurement of a single nerve segment is superior to either long-segment studies or differential subtraction between 2 segments of the same nerve.”</td>
</tr>
<tr>
<td>Kuntzer 1994</td>
<td>5.5</td>
<td>N = 75 healthy subjects with no symptoms of CTS vs. 102 patients suspected on clinical grounds of having CTS</td>
<td>19 different sensorimotor and sympathetic parameters in electrophysiologic studies.</td>
<td>Normal control group values for different electrophysiologic tests. (EDX)</td>
<td></td>
<td>3:1 matched study suggests a single short nerve segment measurement was superior to both long segment studies or differential subtraction between 2 segments of the same nerve for CTS diagnosis.</td>
</tr>
</tbody>
</table>
Bodofsky 2005 Diagnostic 5.5  Patients randomly sampled from electrodiagnostic studies. Divided into 3 groups. 1) Normal Patients (Confirmed using physical exam, history, EMG and NCS) 2) Probable CTS (Symptoms, Physical Exam consistent with CTS. Normal EMG and NCS) 3) Definite CTS (Symptoms, Physical Exam consistent with CTS. EMG and NCS also consistent with CTS) (Median Sensory - Ulnar Motor) Latency difference (MSUMLD) as a more sensitive and specific diagnostic tool for CTS. Other Electrodiagnostic techniques including, Median Sensory Latency, Ulnar sensory latency, Ulnar Motor Latency, (Median-Ulnar) Sensory Latency Difference. MSUMLD had a median value of 0.4 msec in group 1, 1.0 msec in group 2, 2.0 in group 3 (p<0.0001). 95% CI for MSUMLD in normal group is 0.1-0.7 msec. 83% of group 2 patients were added to diagnostically confirmed CTS. 100% of group 3 were diagnosed with CTS using MSUMLD. Sensitivity and Specificity of MSUMLD is 95% and 100%, respectively. "[T]he results in this study strongly suggest that, in patients with symptoms and signs of CTS, the (Median Sensory-Ulnar Motor) Latency difference is an easy simple, highly sensitive and specific test." Data suggest median sensory ulnar latency is obtainable and yields a good sensitivity and specificity in the detection of mild CTS.
<p>| Khosrawi 2013 | Diagnostic | 5.0 | N = 100 healthy hand volunteers and 64 hands of patients with clinical symptoms of CTS | Electrodiagnostic tests (EDX) including Sensory Distal Latency (SDL), Distal Motor Latency (DML), Motor Nerve Conduction velocity (MNCV), Residual Latency (RL) | Clinical Diagnosis of Carpal Tunnel Syndrome. Also comparison of values of Electrodiagnostic readings in control vs diagnosed patients. | Sensitivity and Specificity (%) (95% CI) of EDX tests: SDL 87.3 (83.6-89.1) and 91.2 (89-95.6), DML 70.3 (65.6-71.9) and 100 (96.5-100), MNCV 97.2 (94.4-98.6) ad 90.4 (88.5-94.2), RL 85.9 (84.4-87.5) and 91.1 (87.8-92.2). Median-Ulnar DML difference 84.0 (82.6-85.1) and 89.9 (89-91.1). Median and Ulnar SDL 90.5 (88.1-93.4) and 93.7 (90.2-95.6). | “It seems that, in mild cases of CTS which traditional NCS shows abnormalities only in sensory studies, RL may better demonstrate the effect on median nerve motor fibers.” | Data suggest in mild CTS cases, RL may be a tool to demonstrate the effect on the median nerve motor fibers thus increasing the sensitivity of NCS. |
| Zagnoli 1999 Diagnostic | 5.0 | N = 20 patients (40 wrists) with CTS. Mild (n = 13), moderate (n = 12), severe (n = 8). Follow-up at 31 months. | Electrodiagnostic Studies (Vickers HME device) | MRI | 33/40 wrists showed abnormal electrodiagnostic findings. 11 had isolated sensory abnormalities, and 13 cases showed sensory and motor abnormalities. 2 symptomatic wrists showed normal electrodiagnostic findings (sensitivity 94%) and 2 asymptomatic wrists showed mild to moderate findings (specificity 94%). 32 cases (94%) had sensory abnormalities, 25 had decreased sensory nerve conduction velocity, 29 had decreased sensory nerve potential amplitude. MRI: 20 control wrists normal, 9 clinical symptoms of CTS, 10 had electrodiagnostic abnormalities. 73% sensitivity and 92% specificity of MRI for the diagnosis of CTS. Of 26 MRI studies, 70% had bowing of the transverse carpal ligament. There were 55% of median nerve enlargement and 57% of high median nerve signal. These were correlated with moderate or severe CTS (p &lt;0.001). | &quot;When electrodiagnostic abnormalities suggest more severe disease than expected otherwise discordant with clinical findings, demonstration by magnetic resonance imaging of high median nerve signal and/or median nerve enlargement may help to select those patients most likely to benefit from surgical treatment.&quot; | Small sample size. Data suggest MRI is useful in diagnosing more severe CTS diseases after electrodiagnostic abnormalities have been found. |</p>
<table>
<thead>
<tr>
<th>Study</th>
<th>Diagnostic</th>
<th>N</th>
<th>Population</th>
<th>Methodology</th>
<th>Findings</th>
<th>Implications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Violante 2004</td>
<td>5.0</td>
<td>114 meat workers (228 hands) at risk of CTS; mean age 38.0±10.0 years.</td>
<td>median nerve conduction studies (NCS)</td>
<td>N/A</td>
<td>Significant difference between symptomatic and asymptomatic hands in WSL, SCV-WP, WML, MCV-WP, and the SCV-WP/SCV-EW ratio (all p &lt;0.001). NCS parameters and symptoms had more agreements in non-dominant hand, which was shown in WSL (95% CI: 0.31–0.82) and SCV-WP (95% CI: 0.22–0.59), (p &lt;0.001 and p &lt;0.001).</td>
<td>&quot;Given the importance of the dominant hand in working populations, these data support use of SCV-WP (or WSL) as an informative NCS parameter for occupational studies on CTS.&quot;</td>
</tr>
<tr>
<td>Sheu 2006</td>
<td>5.0</td>
<td>N = 131 hands of CTS patients and 136 hands of controls. Mean age 49.5 years.</td>
<td>Nerve conduction studies</td>
<td>Carpal tunnel diagnosis.</td>
<td>The distoproximal latency ratio (DPLR) of the median nerve showed the highest sensitivity (77%) but had a misclassification rate of 6.9%). The sensitivity of DPLR was not significantly greater than D1M-D1R (p&gt;0.05).</td>
<td>&quot;Optimal transformation of NCS data is mandatory to diminish the effect of skewness and enhance the diagnostic accuracy. As compared to the comparative tests, the segmental study of the median nerve is more easily applied and yields higher sensitivity in detecting mild CTS.&quot;</td>
</tr>
<tr>
<td>Aydin 2004</td>
<td>4.5</td>
<td>N = 525 (818 hands) with suspected CTS confirmed through electrophysiologic evaluation. Mean age 49.1±11.7 years.</td>
<td>Compared sensitivity of first 3 digital branches of median nerve.</td>
<td>Electrophysiological testing was used as the standard diagnostic test in this study.</td>
<td>Most common abnormal physiological findings in Sensory Nerve Conduction Velocity over palm-wrist segment and Digit 1-Wrist segment with sensitivity of 98.5% and 95.4%, respectively.</td>
<td>&quot;The sensory nerve conduction velocity test of the digit 1–to-wrist segment has the most sensitivity among the three digital branches of the median sensory nerve, and it may be used more widely in the electrodiagnosis of carpal tunnel syndrome.&quot;</td>
</tr>
</tbody>
</table>

Data suggest segmental study of the median nerve has application ease and has a higher sensitivity when detecting mild CTS.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Population</th>
<th>Description</th>
<th>Findings</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elkowitz 2005 Diagnostic</td>
<td>2005</td>
<td>N = 72 who had traditional electrodiagnostic testing (EDX) as well as portable NC-Stat testing</td>
<td>A portable Electrodiagnostic testing device (9NC-Stat)</td>
<td>All patients who underwent both types of testing indicated that NC-Stat more comfortable. Both tests had a significantly (p&lt;0.001) linear relationship between Distal motor latencies.</td>
<td>“This portable electrodiagnostic device provides a reliable, convenient, and relatively inexpensive way to obtain objective data and that can be used in diagnosing, evaluating, and treating CTS.”</td>
</tr>
</tbody>
</table>
ULTRASOUND (DIAGNOSTIC)
Ultrasound and high resolution sonography have been investigated for the evaluation and diagnosis of CTS.\(^{(476-487)}\)

**Recommendation:** Ultrasound for Evaluation and Diagnosis of CTS

Ultrasound **is not recommended** for diagnosing CTS.

*Strength of Evidence – Moderately Not Recommended, Evidence (B)*

*Level of Confidence – Moderate*

**Rationale for Recommendation**
Multiple moderate-quality comparative studies report that ultrasound does not outperform and often modestly underperforms compared with EDS for the diagnosis of CTS.\(^{(465, 482, 484, 488, 489)}\) Thus, ultrasound is not recommended for diagnosing CTS. There are other diagnostic uses of ultrasound at the wrist (e.g., evaluating a cyst).

**Evidence for the Use of Ultrasound**
There are 4 moderate-quality studies incorporated into this analysis. \(^{(465, 488-490)}\) There are 3 low-quality studies in Appendix 2.\(^{(475, 491, 492)}\)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: ultrasound diagnostic studies; carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, burning, itching, numbness, tingling, hand, palm, finger, wrist, pain; diagnostic, sensitivity and specificity, positive predictive value, negative predictive value, Predictive Value of Tests, efficacy, efficiency. We found and reviewed 304 articles in PubMed, 370 in Scopus, 4 in CINAHL, and 13 in Cochrane Library. We considered for inclusion 35 from PubMed, 15 from Scopus, 3 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 53 articles considered for inclusion, 43 diagnostic studies and 10 systematic review met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Area of Upper Extremity</th>
<th>Diagnoses</th>
<th>Type of Ultrasound</th>
<th>CT Used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes assessed</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ziswiler 2005</td>
<td>Diagnostic</td>
<td>7.0</td>
<td>Wrist</td>
<td>CTS. Mean age 51±16 years.</td>
<td>5-12 MHz linear array transducer (ATL 3500, Philips Medical System)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>CTS present wrists: 81. CTS absent wrists: 26. ROC curve area under the curve: 0.89 (95% CI 0.82, 0.96); cutoff value 10 mm²; sensitivity 82%; specificity 87%. Likelihood ratios (LR): cutoff of 8 mm² satisfactory power to rule out CTS, fitted-negative LR 0.13 for cross-sectional areas &lt;8 mm²; cutoff of 12 mm² excellent power to rule in CTS, fitted-positive LR 19.9 for areas ≥12 mm².</td>
<td></td>
<td></td>
<td>“Depending on setting and purpose, different cutoff values for the largest cross-sectional area may be used to accurately rule in or rule out CTS.”</td>
<td>Data suggest high correlation between sonography and nerve conduction studies with almost equal sensitivity and specificity.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>N</td>
<td>Gender</td>
<td>Age ± SD</td>
<td>Site</td>
<td>Specimen Preparation</td>
<td>Sensitivity/Specificity (%)</td>
<td>Conclusion</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>---------------</td>
<td>------</td>
<td>---</td>
<td>--------</td>
<td>----------</td>
<td>--------------------</td>
<td>-----------------------</td>
<td>-----------------------------</td>
<td>------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pastare 2009</td>
<td>6.5</td>
<td>66</td>
<td>Men 53, Women 84</td>
<td>52 ± 14</td>
<td>Forearm, Wrist</td>
<td>CTS based on clinical signs and symptoms without previous splinting or surgical treatment for CTS. Mean age 52±14 years, controls 46±15 years.</td>
<td>Ultrasound was performed using a 12-MHz linear array transducer</td>
<td>- - - + - + - - + -</td>
<td>Nerve Conduction studies showed greater diagnostic sensitivity than ultrasound; 54 wrists 82% vs. 41 62% for highly likely clinical diagnosis of CTS.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visser 2008</td>
<td>6.0</td>
<td>168 (53 men, 84 women)</td>
<td>52 (± 14)</td>
<td>Forearm, Wrist</td>
<td>CTS based on clinical signs and symptoms without previous splinting or surgical treatment for CTS. Mean age 52±14 years, controls 46±15 years.</td>
<td>Ultrasound was performed using a 12-MHz linear array transducer</td>
<td>- - - + - + - - + -</td>
<td>Ultrasound was performed using a 12-MHz linear array transducer</td>
<td>Sensitivity/specificity (%): Wrist cross-sectional area &gt;0.1 cm²: 78 (70-84)/91 (86-95), Median- ulnar digit 4 difference &gt;0.4 msec: 82 (75-88)/97 (89-100), Median nerve &gt;3.8 msec: 74 (66-81)/97 (88-100).</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Conclusion:**

- In summary, our study shows that NCS have better sensitivity in supporting a diagnosis of CTS. However, because of its high positive predictive value, lack of discomfort, and ease of use, US can be used as a screening method for CTS.

- Data suggest nerve conduction studies are superior to sonography in detecting CTS.

- In patients with a clinical diagnosis of CTS, the accuracy of sonography is similar to that for EMG.

- Data suggest sonography is comparable to EMG in patients with a clinical diagnosis of CTS but study states EMG should still be first diagnostic test utilized in patients with atypical symptoms.
<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Study Design</th>
<th>Sample Size</th>
<th>Age</th>
<th>Symptom Duration</th>
<th>Ultrasound Equipment</th>
<th>Ultrasound Parameters</th>
<th>Diagnostic Criterion</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wang</td>
<td>2008</td>
<td>Diagnostic</td>
<td>N = 37 (20 controls)</td>
<td>Mean age CTS patients (44±9.4 years) and healthy subjects (43.7 ± 12.91 years)</td>
<td>Classic or probable symptoms of CTS for 1-60 months</td>
<td>Sequoia 512 with 8-15 MHz broad line transducer</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
MAGNETIC RESONANCE IMAGING AND DIFFUSION TENSOR IMAGING

MRI and especially diffusion tensor imaging (“diffusion MRI) are being investigated for the evaluation and diagnosis of CTS.(493-543)

**Recommendation: MRI and Diffusion Tensor Imaging for Evaluation and Diagnosis of CTS**

**MRI and Diffusion Tensor Imaging is moderately not recommended for diagnosing CTS.**

- **Strength of Evidence** – Moderately Not Recommended, Evidence (B)
- **Level of Confidence** – Moderate

**Rationale for Recommendation**

Multiple moderate-quality comparative studies report that MRI and Diffusion Tensor Imaging do not outperform and often modestly underperform compared with EDS for the diagnosis of CTS.(469, 544-546) Thus, MRI and Diffusion Tensor Imaging are not recommended for diagnosing CTS. There are other diagnostic uses of MRI at the wrist.

**Evidence for the Use of Magnetic Resonance Imaging and Diffusion Tensor Imaging**

There are 6 moderate-quality studies incorporated into this analysis.(469, 544-548) There are 5 low-quality studies in Appendix 2.(475, 549-552)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: magnetic resonance imaging, carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, diagnostic, sensitivity and specificity, positive predictive value, negative predictive value, predictive value of tests, efficacy, efficiency. We found and reviewed 287 articles in PubMed, 383 in Scopus, 8 in CINAHL, and 5 in Cochrane Library. We considered for inclusion 66 from PubMed, 6 from Scopus, zero from CINAHL, zero from Cochrane Library and 3 from other sources. Of the 75 articles considered for inclusion, 68 diagnostic studies and 1 systematic review met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Number</th>
<th>Area of Upper Extremity</th>
<th>Diagnoses</th>
<th>Type of MRI used</th>
<th>T1 weighted images</th>
<th>T2 weighted images</th>
<th>X-ray</th>
<th>Myelography</th>
<th>More than one rater</th>
<th>Surgery Performed</th>
<th>Clinical outcomes assessed</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jarvik 2002</td>
<td>7.0</td>
<td>N = 120</td>
<td>W</td>
<td>CTS</td>
<td>MRI using 1.5 Tesla Magnets</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>Intrareader reliability was substantial to near perfect (kappa = 0.76 - 0.88). Interreader lower but still substantial (kappa = 0.60 - 0.67). Sensitivity of MRI was greatest for the overall impression of the images (96%) followed by increased median nerve signal (91%) and with lower specificities (33 - 38%).</td>
<td>“The reliability of MRI is high but the diagnostic accuracy is only moderate compared to a research-definition reference standard.”</td>
<td>Study used a mixed cohort (both men and women) to enhance diagnostic accuracy (those who were true positive for CTS) using high resolution MRI. Data suggest MRI has a “moderate” diagnostic accuracy at best compared to the reference standard for CTS. Also, assumption that a high STIR signals within the palmar bursa as being a marker for CTS was likely incorrect as normal signals within palmar bursa were associated with CTS presence.</td>
</tr>
<tr>
<td>Study Year</td>
<td>Authors</td>
<td>N</td>
<td>Gender</td>
<td>Type</td>
<td>Imaging System</td>
<td>Wavelet/Region</td>
<td>Results/Comments</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------</td>
<td>---------</td>
<td>---</td>
<td>--------</td>
<td>------</td>
<td>----------------</td>
<td>----------------</td>
<td>-----------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2014 Bulut</td>
<td>No mention of</td>
<td>N = 120 (90 females and 30 males)</td>
<td>W</td>
<td>Carpal Tunnel Syndrome</td>
<td>1.5-T whole-body MRI system was used for all MRI examinations.</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>Diffusion tensor imaging (DTI) showed significant correlations with electrophysiological studies (EPS). DTI parameter (Fractional anisotropy-FA and apparent diffusion coefficients (ADC)) evaluated and significant difference between CTS and controls with CTS patients showing significantly lower FA and ADC scores (p ≤0.001).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2005 Uchiyama</td>
<td>Diagnostic</td>
<td>105 wrists of 105 women. 36 wrists of 36 female volunteers.</td>
<td>W</td>
<td>Idiopathic CTS</td>
<td>1.5 Tesla with a circular extremity coil.</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>Flattening of nerve more significant at distal TCL level than other levels. Cross sectional area larger in mild to moderate group vs. controls at DRUJ/pisiform/hook of hamate/distal TCL levels: 14.1 (4.8) vs. 9.0 (2.5)/14.6</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

"Severity of the disease could be judged by evaluating not only longitudinal changes of signal intensity and configuration of the median nerve, but also palmar bowing of the TCL. Increased palmar bowing of the TCL was found to be associated with an increase in the area of the carpal tunnel."

Data suggest significant differences between all subgroups for mean FA and ADC suggesting FA and ADC threshold values could be useful for diagnosing and grading CTS. The DTI parameters well significant versus EPS for assessment of severity.
| Zagnoli 1999 | 5.0 | 20 | W | Carpal tunnel syndrome | MRI vs. electrodiagnostic (Vickers HME device) | - | + | + | - | - | + | - | 31 months | 33/40 wrists showed abnormal electrodiagnostic findings. 11 cases showed isolated sensory abnormalities, and 13 cases showed sensory and motor abnormalities. | When electrodiagnostic abnormalities suggest more severe disease than expected otherwise discordant with clinical findings, demonstration by magnetic resonance imaging of high median nerve signal and/or median nerve may detect abnormalities after electrodiagnostic abnormalities have been found. | Small sample size. Data suggest MRI may detect abnormalities after electrodiagnostic abnormalities have been found. |
Symptomatic wrists showed normal electrodiagnostic findings (sensitivity 94%) and 2 asymptomatic wrists showed mild to moderate findings (specificity 94%). 32 (94%) had sensory abnormalities, 25 had decreased sensory nerve conduction velocity and 29 had decreased sensory nerve potential amplitude. In MRI, 20 control wrists normal, 9 had clinical CTS symptoms and 10 wrists had electrodiagnostic abnormalities. 73% sensitivity and 92% specificity of MRI for diagnosis of CTS. Of 26 MRI studies, 70% had bowing of transverse enlargement may help to select those patients most likely to benefit from surgical treatment.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Subjects</th>
<th>Methodology</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brienza 2014</td>
<td>4.5</td>
<td>30</td>
<td>Carpal tunnel syndrome 3-Tesla magnetic resonance imaging with diffusion tensor imaging (DTI)</td>
<td>Carpal ligament, 55% of median nerve enlargement and 57% of high median nerve signal. These were correlated with moderate or severe CTS (p &lt;0.001). Results do not reflect MRI, focused only on Electroneurography. Data suggest a high degree of correlation between DTI and ENG of the peripheral nervous system.</td>
</tr>
<tr>
<td>Wang 2012</td>
<td>4.0</td>
<td>40</td>
<td>Carpal tunnel syndrome Diffusion tensor imaging (DTI). 1.5-T whole body with a microcopy coil.</td>
<td>Overall results of FA and ADC at different levels (distal radius, pisiform bone, middle of tunnel, and hamate bone) were similar. Only CTS had significant effects on FA and ADC (p &lt;0.05). Linear correlation between distal latency of motor conduction and FA and ADC at the distal radius, pisiform bone, in the carpal tunnel and at the hamate bone. The correlations of FA and ADC were significant as compared with EP for CTS.</td>
</tr>
</tbody>
</table>

Small study population (n = 40). Data suggest FA and ADC were independent of finger posture and measuring location. Mean FA was decreased by CTS and ADC was increased by CTS. Study reports DTI imaging of FA and ADC were significant as compared with EP for CTS.
velocity of median nerve (MNDL) and length of abnormal intensity of median nerve (N_Len). If N_Len >15mm used as criteria for CTS, there was 1 false negative case and no false positive cases ($r^2= 0.529, p <0.001$).
**Monitoring Progress**

The clinical evaluation and progress of patients is most commonly monitored qualitatively from appointment to appointment. Particularly, physicians seek information regarding the degree to which symptoms are present and whether the patient believes there has been improvement. However, there are several instruments that may be utilized for monitoring the progress of patients with CTS. These include the DASH (554-575) and Boston Carpal Tunnel Questionnaire. The Short Form-36 (SF-36), the Flinn Performance Screening Tool (FPST), the Patient Evaluation Measure (PEM), the Amadio questionnaire, the Historical-objective-distribution based scale (Hi-Ob-Db), and the Alderson-McGali hand function questionnaire (AMHFQ) have been used to diagnose CTS. VAS symptoms and pain scores may also be used (561, 565, 576) even though many patients with CTS have no pain. Functional status scores (407, 554, 567, 571, 577, 582, 586, 587, 589, 592, 594, 598-600) and Global Symptom Scores (601) are also used, particularly in some research studies. Grip strength (560, 565, 576, 583, 584, 589, 596, 602-605) and pinch strength measures (560, 565, 576, 581, 583, 584, 588, 596, 602, 604) may be utilized. However, patients who have mild symptoms generally have normal grip strength. All of these questionnaires are subjective and strength measures are effort-dependent, although the strength measures attempt to provide a quantitative measure that may help to gauge improvement over time especially post-operatively (554, 558, 564, 578, 586, 594, 601, 606-609).

**Recommendation: Instruments for Monitoring the Progress of Patients with CTS**

**There is no recommendation for or against the use of instruments to monitoring the progress of patients with CTS.**

*Strength of Evidence – No Recommendation, Insufficient Evidence (I)*

*Level of Confidence – Low*

**Rationale for Recommendation**

There is no quality evidence that any of these instruments meaningfully contribute to improving clinical care. They may be more useful in the post-operative setting.

**Evidence for the Use of Monitoring Progress**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: monitoring progress: disabilities of the arm, shoulder and hand questionnaire, Boston carpal tunnel questionnaire, VAS symptoms score; VAS pain score, functional status scores, global symptom scores, grip strength, pinch strength, carpal tunnel syndrome, median neuropathy, carpal tunnel syndrome/diagnosis, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, predictive value of tests, efficacy, efficiency. We found and reviewed 527 articles in PubMed, 123 in Scopus, 32 in CINAHL, and 23 in Cochrane Library. We considered for inclusion 59 from PubMed, 2 from Scopus, zero from CINAHL, zero from Cochrane Library and zero from other sources. Of the 61 articles considered for inclusion, 59 diagnostic studies and 1 systematic review met the inclusion criteria.
Specific Treatment Interventions

Activity Modification and Exercise

EXERCISES

Various exercise regimens have been utilized to treat patients with CTS, most commonly tendon-gliding and nerve-gliding exercises.(610-617) These exercises are thought to help prevent adhesion formation.(615, 618-620)

1. Recommendation: Exercises for Treatment of Acute, Subacute, or Chronic CTS

   There is no recommendation for or against the use of exercises for treatment of chronic CTS as quality evidence is lacking.

   Strength of Evidence – No Recommendation, Insufficient Evidence (I)

   Level of Confidence – Low

2. Recommendation: Exercises for Rehabilitation of Post-operative CTS Patients with Significant Deficits

   Exercise is recommended for rehabilitation of post-operative CTS patients with significant deficits.

   Strength of Evidence – Recommended, Insufficient Evidence (I)

   Level of Confidence – Low

Rationale for Recommendations

There are multiple moderate quality studies, but none has clearly found benefit of exercises, including tendon-gliding, for treatment of CTS. One moderate-quality study suggested no statistically significant incremental benefit from adding tendon-gliding exercises to wrist splinting,(610) although modest trends towards benefit appear present. Another moderate-quality study found a combination of tendon-gliding exercise with ultrasound and splinting superior to two other combinations.(611) Thus, it is unclear if there is an independent benefit from tendon-gliding exercises. Additionally, as many believe that physical activity is a risk factor for CTS, the logic of performing exercises for treatment is somewhat dissonant. However, exercise programs are not invasive, have few if any adverse effects, and are low cost if performed independently after receiving initial instructions.

Evidence for the Use of Exercise for CTS

There are 5 moderate-quality RCTs incorporated into this analysis.(610, 611, 621-623) There are 4 low-quality RCTs in Appendix 2.(624-627)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: gliding exercise, tendon-gliding, tendon gliding, nerve-gliding, nerve gliding, neurodynamic mobilization, upper limb tension test, ULTT; carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 26 articles in PubMed, 19 in Scopus, 8 in CINAHL, and 31 in Cochrane Library. We considered for inclusion 13 from PubMed, 1 from Scopus, 1 from CINAHL, 1 from Cochrane Library and 1 from other sources. Of the 17 articles considered for inclusion, 10 randomized trials and 4 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brininger 2007</td>
<td>6.0</td>
<td>N = 61 (41 females and 10 males) with a positive Tinel sign or Phalen maneuver and complaints of nocturnal numbness and tingling. Mean age 50 years.</td>
<td>Neutral wrist and metacarpophalangeal (MCP) splint, custom splint positioning MCP joints from 0º to 10º of flexion, NW/MCP (n=17) vs. neutral wrist and MCP exercise group (tendon and nerve gliding exercises 3 to 5 times a day with 10 repetitions in each position, and to hold each position for 5 seconds), NW/MCP-X (n=16) vs. wrist cock-up splint prefabricated that immobilized the wrist in 20º of extension, WCU (n=12) vs. wrist cock-up splint and exercise, WCU-X (n=16). All groups wore the splint during sleep for 4 weeks and received and educational brochure on CTS. Assessments at baseline, 4 weeks, and 8 weeks.</td>
<td>All groups saw significant decrease in CTS symptoms (no p-value reported).</td>
<td>&quot;Our results provide further evidence of the effectiveness of splinting, designed to target an underlying anatomic problem, for reducing symptoms and improving functional status in patients with mild-to-moderate CTS.&quot;</td>
<td>Small group numbers. No table or graphic for results. Baseline comparability for group strength different between groups.</td>
</tr>
<tr>
<td>Baysal 2006</td>
<td>5.5</td>
<td>N = 36 females EDS confirmed CTS, all bilateral, all right handed. Mean age: Group I 47.8±5.5 years, Group II: 50.1±7.3 years, Group III: 51.4±5.2 years.</td>
<td>Group I: tendon- and nerve-gliding exercises 5 sessions daily, each exercise repeated 10 times/session for 3 weeks plus splinting full-time for 3 weeks (n = 12) vs. Group II: ultrasound 15 minutes per session to palmar carpal tunnel at frequency pf 1 MHz and intensity of 1.0 W/cm² once a day 5 days a week, 3 weeks plus splinting (n = 12) vs. Group III: ultrasound, splinting and tendon-nerve-gliding exercises (n = 12). Follow-up at end of treatment at after 8 weeks.</td>
<td>Pain score before treatment/after treatment I/after treatment II: Group I: 4.8±2.3/3.3±2.9/ 2.6±2.8; Group II: 5.7±2.7/ 2.2±1.9/ 2.5±2.8; Group III: 5.6±3.5/ 1.3±1.8/ 0.8±0.9. Functional status score: Group I: 20.6±7.8/14.8±7.5/ 14.9±6.6; Group II: 21.9±9.1/16.1±8.5/ 16.1±8.7; Group III: 20.5±7.1/11.7±3.6/ 12.6±3.4. NS between groups.</td>
<td>&quot;The result of this study emphasizes the efficacy of conservative treatment in CTS. In all patients groups, the treatment combinations were significantly effective immediately and 8 weeks after the treatment.&quot;</td>
<td>All groups were splinted precluding judgment of utility of splinting. Unclear if there is an independent effect of exercise.</td>
</tr>
<tr>
<td>Bialosky 2009</td>
<td>5.5</td>
<td>N = 40 females with &gt;12weeks</td>
<td>Neurodynamic technique (n = 20) vs. Sham technique (n = 20)</td>
<td>Values for between-group comparisons of clinical</td>
<td>&quot;Collectively, these findings suggest that NDT specific to...&quot;</td>
<td>Few differences between treatment arms were seen.</td>
</tr>
<tr>
<td>RCT</td>
<td>Schmid 2012</td>
<td>No sponsorship. No mention of COI.</td>
<td>N = 21 with mild to moderate CTS. Mean age: 53.9 years.</td>
<td>Nerve and tendon gliding exercise home program (n = 11) vs. Night splinting (n = 10). Follow-up at 1-week.</td>
<td>No significant differences present between groups. Within group Baseline vs. Follow-up – Exercise: Pain intensity VAS (0.7 vs. 0.8; p&gt;0.16). Numbness VAS (1.5 vs. 1.6; p&gt;0.16). Splinting: Pain intensity VAS (1.2 vs. 1.1; p&gt;0.16). Numbness VAS (2.3 vs. 1.9; p&gt;0.16).</td>
<td>“The findings of this study suggest that a reduction in intraneural edema is a therapeutic mechanism of both nerve and tendon gliding exercises and splinting… there seems to be no preference for splinting or nerve and tendon gliding exercises.”</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>4.5</td>
<td>N = 28 EDS confirmed CTS. Mean age 51.93±5.1 years.</td>
<td>Full-time splint (n=14) vs. full-time splint plus nerve tendon gliding exercises 5 sessions daily with each exercise repeated 1- times per session (n=14) for 4 weeks. Follow-up 8 weeks after treatment.</td>
<td>Grip strength (mean ± SD) – Pre-/post-treatment: Group I (splint): 38.44±14/49.88±15.3; Group II (exercise + splint): 38.61±13.8/54.94±17 p (between groups) = 0.14. Symptom severity score (mean ± SD): Group I: 36.11±9.0/21.88±8.8; Group II: 35.9±6.0/18.2±5.85 p (between groups) = 0.210</td>
<td>“Although the results in group 2 were better than group 1, the difference was not statistically significant. Further investigations are required to establish the role of nerve and tendon gliding exercises in the treatment of carpal tunnel syndrome.”</td>
<td>No clear evidence of benefit.</td>
<td>Small sample size (N=21). Data suggest no differences.</td>
</tr>
</tbody>
</table>
Yoga has been used to treat CTS, although its main uses have been in treating spine pain and other more widespread MSDs (see Chronic Pain and Low Back Disorders Guidelines).

**Recommendation:** Yoga for Treatment of Acute, Subacute, or Chronic CTS

There is no recommendation for or against the use of yoga for treatment of acute, subacute, or chronic CTS.

**Strength of Evidence – No Recommendation, Insufficient Evidence (I)**

**Level of Confidence – Low**

**Rationale for Recommendation**

There is one moderate-quality RCT that suggested improvements in grip strength; however, the comparative population had an inactive splint for treatment which may have created an artificial difference in grip strength. While yoga appears beneficial for treatment of spine patients, there is no evidence of efficacy for distal upper extremity MSDs. Yoga is not invasive, has low potential for adverse effects, and is low cost. Compliance and adherence are reportedly not good, as patient motivation must be high and there is much self-selection in studies assessing yoga’s efficacy.

**Evidence for the Use of Yoga for CTS**

There is 1 moderate-quality RCT incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: yoga and carpal tunnel syndrome, CTS, median nerve neuropathy, median neuropathy, median nerve disease, entrapment, neuropathy, nerve compression, burning, itching, numbness, tingling, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 16 articles in PubMed, 183 in Scopus, 7 in CINAHL, 17 in Cochrane Library and zero in other sources. We considered for inclusion 2 from PubMed, zero from Scopus, zero from CINAHL, zero from Cochrane Library and zero from other sources. Of the 2 articles considered for inclusion, 1 randomized trials and 1 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Garfinkel 1998 RCT</td>
<td>6.0</td>
<td>N = 51</td>
<td>Standard splint to supplement current treatment (n = 26) vs. yoga (n = 25). Follow-up at 8 weeks.</td>
<td>Grip strength yoga (161.6±70.4 to 187.4±68.8) vs. splint (183.3±69.5 to 190.5±68.2mm Hg). Pain reduced (p = 0.02). Median nerve sensory conduction yoga (4.40±1.5ms to 3.97±1.5) vs. splint (4.66±1.4 to 4.36±1.6ms) (NS).</td>
<td>“In this preliminary study, a yoga-based regimen was more effective than wrist splinting or no treatment in relieving some symptoms and signs of carpal tunnel syndrome.”</td>
<td>Grip strength improvement may be from activity in yoga as comparison was presumably an inactive splint which may have caused greater improvement not related to CTS. Lack of description of controls limits interpretations.</td>
</tr>
</tbody>
</table>

**Medications**

**NSAIDS/Acetaminophen**

Nonsteroidal anti-inflammatory drugs (NSAIDs) have been widely used to address beliefs in inflammatory mechanisms of CTS or to manage pain associated with CTS (630-635) (see Chronic Pain Guideline for detailed discussion on mechanisms of action, classes of medications, adverse effects, etc.). Acetaminophen and paracetamol are sometimes utilized to treat CTS, although their effects on cyclooxygenase activity are minimal and they are not anti-inflammatory.

1. **Recommendation:** NSAIDs or Acetaminophen for Subacute or Chronic CTS
NSAIDs or acetaminophen are not recommended as a primary treatment for subacute or chronic CTS. (636)

**Strength of Evidence** – Not Recommended, Evidence (C)
**Level of Confidence** – Low

2. **Recommendation: NSAIDs for Post-operative Management of CTS-related Pain**

NSAIDs are moderately recommended for post-operative management of CTS-related pain.

**Indication** – Patients having recently undergone carpal tunnel surgical release. Generally, treat 2 weeks up to 6 weeks post-op unless complications occur.

**Frequency/Dose** – See manufacturer’s recommendations.

**Indications for Discontinuation** – Resolution of pain, adverse effects, intolerance.

**Strength of Evidence** – Moderately Recommended, Evidence (B)
**Level of Confidence** – High

3. **Recommendation: Acetaminophen for Post-operative Management of CTS-related Pain**

Acetaminophen is recommended for post-operative management of CTS-related pain.

**Indication** – Patients having recently undergone carpal tunnel surgical release. Generally treat 2 weeks up to 6 weeks post-op unless complications occur.

**Frequency/Dose** – See manufacturer’s recommendations.

**Indications for Discontinuation** – Resolution of pain, adverse effects, intolerance.

**Strength of Evidence** – Recommended, Evidence (C)
**Level of Confidence** – Low

**Rationale for Recommendations**

While NSAIDs have been widely used to attempt to address a theoretical inflammatory basis for CTS and/or to treat pain associated with CTS, the one quality study comparing an NSAID to placebo found no benefit from the NSAID. (636) This same study also found no difference between NSAIDs and diuretics which also appear ineffective. There is also no quality evidence that there is a difference among NSAIDs (see Hip and Groin Disorders Guideline). Thus, there is quality evidence that NSAIDs do not have a role in the treatment of typical cases of CTS. (636)

Other studies comparing NSAIDs with manipulation plus ultrasound (637) and lidocaine patch (638) did not find benefits of NSAIDs compared with those treatments. A trial combining splinting (which appears effective) plus NSAID versus glucocorticosteroid injection did not find one arm to be superior. (631) While some patients may benefit from NSAIDs, evidence is lacking that there is any beneficial effect of NSAIDs for treatment of CTS and aggregate analyses of these studies also suggest NSAIDs are ineffective. (636-638) Acetaminophen is thought to also be ineffective.

NSAIDs are not invasive and have low adverse effects profiles, particularly when used for short courses in occupational populations. Generic or over-the-counter formulations are low cost. However, there is quality evidence that other interventions are effective. A short course of an over-the-counter NSAID may be reasonable for select patients; however, routine use of NSAIDs for treatment of CTS is not recommended. Select patients with acute CTS due to unaccustomed forceful use may be potential candidates for treatment with NSAIDs; however, that population has not been studied in quality trials.

There is one high-quality study in post-operative patients indicating that for post-operative pain management, naproxen is superior to acetaminophen, which in turn is superior to placebo. (639) NSAIDs and acetaminophen may also facilitate the rehabilitation process without the impairments associated with opioids. Thus, NSAIDs and acetaminophen are recommended for post-operative pain management.

**Evidence for the Use of NSAIDs and Acetaminophen for CTS**

There are 2 high- (639, 640) and 5 moderate-quality (631, 636-638, 641) RCTs incorporated into this analysis. There is 1 low-quality RCT in Appendix 2. (642)
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: anti-inflammatory agents, non-steroidal, non-steroidal, anti-inflammatory, NSAIDS, aspirin, diflunisal, salsalate, ibuprofen, dexibuprofen, naproxen, fenoprofen, ketoprofen, dexketoprofen, flurbiprofen, oxaprozin, loxoprofen, indomethacin, tolmetin, sulindac, etodolac, ketorolac, diclofenac, nabumetone, piroxicam, meloxicam, tenoxicam, d Roxicam, lornoxicam, isoxicam, celecoxib, etodolac, etoricoxib, lumiracoxib, meclofenamic acid, mefenamic acid, nimesulide, parecoxib, rofecoxib, toltenamic acid, valdecoxib; carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, burning, tingling, itching, numbness, hand, palm, finger, wrist, pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 41 articles in PubMed, 302 in Scopus, 10 in CINAHL, and 2 in Cochrane Library. We considered for inclusion 11 from PubMed, 0 from Scopus, 1 from CINAHL, 0 from Cochrane Library and 1 from other sources. Of the 13 articles considered for inclusion, 9 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chang 1998</td>
<td>RCT</td>
<td>Sponsored by NSC 86-2314-B-075B-012 to Ming-Hong Chang. No mention of COI.</td>
<td>7.0</td>
<td>N = 73 (53 female/20 male) with clinical signs and symptoms of CTS, EDS confirmed without abnormalities in radial and ulnar nerves. Mean age diuretic 45.7±4.8 years, NSAID-SR 47.4±5.7 years, steroid 45.4±5.2, placebo 44.2±5.4.</td>
<td>Trichlor-methiazide (diuretic), 2mg daily for 4 weeks (n = 16) vs. Tenoxicam-SR (NSAID-SR), 20mg daily for 4 weeks (n=18) vs. prednisolone (steroid) 20mg/day for 2 weeks, then 2-week dose of 10mg daily (n = 23) vs. placebo for 4 weeks (n = 16). Assessments at baseline, 2 weeks and 4 weeks.</td>
<td>Meant±SD global symptom score (GSS) baseline/2 weeks/4 weeks: placebo 22.9±5.9/21.6±6.4/20.8±6.6 vs. diuretics 26.0±3.8/22.3±5.5/21.6±6.3 vs. NSAID-SR 29.7±8.4/24.7±8.6/24.0±9.7 vs. steroid 27.9±6.9/15.0±6.8/10.0±7.5 (p &lt;0.0005 at week 2 steroid vs. other treatment groups; p &lt;0.0001 at week 4 steroid vs. placebo).</td>
<td>“For patients with mild to moderate CTS who opt for conservative treatment, corticosteroids are of greater benefit.”</td>
<td>Suggests oral steroids effective but diuretic and NSAID are not effective compared with placebo.</td>
</tr>
<tr>
<td>Yildiz 2011</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>8.0</td>
<td>N = 51 (43 females/8 males) with signs and symptoms of CTS for more than a month and mild-to-moderate CTS after electrodiagnostic test confirmation. Age range 39-66 years.</td>
<td>Group 1: sham ultrasound (US), ultrasound in off mode 15 minute sessions 1x a day, 5x a week for 2 weeks plus splinting with neutral custom-molded thermoplastic volar wrist splint at night and during day for 8 weeks (n = 17, 25 median nerves) vs. Group 2: US, pulse mode (1:4) with gel without medication at 1 MHz frequency and 1 W/cm² intensity plus splinting (n = 17, 26 median nerves) vs. Group 3: ketoprofen phonophoresis (PH), US pulse mode (1:4) with 2.5% ketoprofen gel at 1 MHz frequency</td>
<td>Meant±SD VAS (baseline/2 weeks/8 weeks): Group 1, 5.76±2.45/2.72±2.07/3.28±2.74 vs. Group 2, 4.96±2.50/2.41±2.43/2.77±2.74 vs. Group 3, 6.04±2.40/3.03±1.96/0.98±1.65 (p = 0.002, Group 3 &gt; Group 1; p = 0.004, Group 3 &gt; Group 2).</td>
<td>“Our results suggest that ketoprofen PH in addition to splinting is superior to the combination of US and splinting with respect to pain only in middle term patients with CTS.”</td>
<td>Ultrasound plus splinting not superior to splinting alone. Ketoprofen plus splinting was associated with a reduction in pain at 8 weeks.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>Sponsorship</td>
<td>Notes</td>
<td>N</td>
<td>Description</td>
<td>Intervention</td>
<td>Primary Outcome</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>--------</td>
<td>-------------</td>
<td>-------</td>
<td>----</td>
<td>-------------</td>
<td>--------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>Chang 1998</td>
<td>See above.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jarvik 2009</td>
<td>RCT</td>
<td>Sponsored by the Intramural Research Program of the NIH Clinical Center. No COI.</td>
<td></td>
<td></td>
<td>7.0</td>
<td>N = 116 (62 female/54 male) considering surgery for diagnosed carpal tunnel syndrome. Mean age 50.7 years.</td>
<td>Surgery group: open surgery or endoscopic surgery depending on surgeon's preference (n = 57) vs. non-surgical therapy group: 6 visits with hand therapist focused on ligament stretching, tendon gliding, and review of splint use (splint use at night) plus prescribed NSAIDS, ibuprofen 200mg 3x a day. If no improvement after 6 weeks, received 12 sessions (2-4 per week for up to 6 weeks) of focused ultrasound at 1 MHz, 1-0 W/cm² in pulsed mode 1:4, 15 minutes each (n = 59). Follow-up at 3, 6, 9, 12 months.</td>
<td>Primary outcome was Carpal Tunnel Syndrome Assessment Questionnaire (CTSAQ). Surgical group showed significantly lower CTSAQ function score vs. non-surgical group at 6 months: 1.91 vs. 2.44 (p = 0.0006) and at 12 months: 1.74 vs. 2.17 (p = 0.0081). Secondary outcome of CTSAQ symptoms was also significantly lower in surgical group vs. non-surgical group at 6 months; 2.02 vs. 2.42 (p = 0.018) and 12 months; 1.74 vs. 2.07 (p = 0.036).</td>
</tr>
<tr>
<td>Celiker 2002</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td>5.5</td>
<td>N = 23 with unilateral or bilateral CTS, EDS confirmed. Mean age Group A 49.6±15.3 years, Group B 46.9±10.0 years. Group A: acemetacine 120mg a day with splints at night, light-weight, neutral-positioned (n=11) vs. Group B: 40mg methylprednisolone acetate 1ml (n=12). Follow-up at 2 weeks and 8 weeks.</td>
<td>VAS pain scores (baseline/2nd week/8th week): NSAID plus splint 7.9±1.4/4.3±0.9/1.7±1.0 vs. injection 7.0±2.2/3.1±2.5/1.8±1.9 (p&lt;0.05). Symptom severity scale results similar.</td>
<td>&quot;Both splinting combined with the use of a nonsteroidal anti-inflammatory drug and steroid injection into the carpal tunnel resulted in significant improvement in carpal tunnel syndrome.&quot;</td>
</tr>
<tr>
<td>Davis 1998</td>
<td>RCT</td>
<td>Sponsored by a grant from the National Chiropractic Mutual Insurance Company. No mention of COI.</td>
<td></td>
<td></td>
<td>5.0</td>
<td>N = 91 with self-reported symptoms of CTS and EDS confirmed CTS. Mean age ibuprofen group 38±5 year, manipulation</td>
<td>Ibuprofen (800mg 3x a day for 1 week, then 2x a day for 1 week, then PRN 7 weeks) and nocturnal cock-up wrist supports (n = 46) vs. high velocity, low amplitude manual thrust procedures:</td>
<td>CTS outcome assessment physical distress (mean±SD) baseline to end of study: IBU 14.66±9.89 vs. injection 12.47±6.28 vs. ultrasound and manipulation 9.25±8.14 (p = 0.0132). CTS outcome assessment mental distress</td>
</tr>
</tbody>
</table>

At 12 months, surgical group was significant for improved symptoms and function.

Baseline did not exclude prior ibuprofen use or manipulation, but prior use of these treatments is likely differential between the 2 groups and is a potentially fatal study flaw. Ibuprofen use was PRN after 2 weeks and subject contact differed.
<table>
<thead>
<tr>
<th>Study</th>
<th>Chief Investigator</th>
<th>Sample Size</th>
<th>Study Design</th>
<th>Intervention</th>
<th>Outcome Measures</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nalamachu 2006 MedGenMed RCT</td>
<td>Sponsored by Endo Pharmaceuticals. COI, Nalamachu received research grants and consulting fee from Endo Pharmaceuticals. PharmD is employed by Endo Pharmaceuticals as Senior Director, Medical Affairs, and receives annual stock options from Endo. Gould is employed by Endo Pharmaceuticals as Associate Director, Medical Affairs, and receives annual stock options from Endo.</td>
<td>4.5</td>
<td>N = 100 age 18-75 with CTS, clinical and EDS confirmed. Mean age lidocaine patch 55.7±16.0 years, naproxen 51.5±11.8 years.</td>
<td>Brief Pain Inventory (BPI) scores reduced between baseline and Week 6 for both lidocaine patch 5% (p &lt;0.0001) and naproxen 500 mg twice daily (p = 0.0064), but no between group differences (p = 0.083). Clinical Global Impression of Improvement (CGI-I) scores also favored patch (51.1% vs. 24.3%, p = 0.016). Percentages satisfied or very satisfied 71.8% lidocaine patch vs. 63.2% naproxen (NS).</td>
<td>“This study demonstrates that the lidocaine patch 5% is effective in significantly relieving the pain associated with CTS and is well tolerated. The patch may offer patients an effective, non-systemic, noninvasive treatment for the management of their symptoms. Further controlled studies are warranted.”</td>
<td></td>
</tr>
<tr>
<td>Husby 2001 RCT</td>
<td></td>
<td>8.0</td>
<td>N = 77 who underwent surgery for CTS</td>
<td>Post-op naproxen 500mg BID (n = 26) vs. paracetamol 1,000mg</td>
<td>Postoperative CTS swelling as a percentage of preoperative volume 3.5±3.3</td>
<td>Naproxen might have a clinical relevant effect on swelling when used on</td>
</tr>
<tr>
<td>No mention of sponsorship or COI.</td>
<td>of Dupuytren's contracture (DC). Mean age 59 years.</td>
<td>QID (n = 26) vs. placebo tablets (n = 25) for 3 days immediate post-op carpal tunnel release surgical treatment; 2nd trial included 35 with Dupuytren's contracture. Opioid analgesic allowed for supplementary analgesic. No mention of follow-up time.</td>
<td>vs. 4.6±3.2 vs. 3.8±2.6. For Dupuytren's contracture releases: 5.6±3.8 vs. 6.9±3.7 vs. 8.2±5.1. Additional analgesics used were 0, 2, and 8 in naproxen, paracetamol, and placebo groups.</td>
<td>minor surgery in the hand, unlike paracetamol. Naproxen might be a useful analgesic during the immediate post-operative phase.</td>
<td>superior to placebo, which the studies were not powered to detect.</td>
<td></td>
</tr>
</tbody>
</table>
Systemic Glucocorticosteroids (AKA “Steroids”)

Oral
Glucocorticosteroids are used to treat CTS and other tendinoses through both oral and injection routes (injections for CTS and other tendinoses).(636, 643-648) Although these medications are considered to be anti-inflammatory corticosteroids, absent an inflammatory arthropathy or infection, CTS does not typically evidence inflammation. Thus, the exact mechanism of action is uncertain. Regardless, evidence indicates that carpal tunnel injections are superior to oral steroids for treatment of CTS.(648)

**Recommendation:** Oral Glucocorticosteroids for Treatment of Acute, Subacute, or Chronic CTS

Oral glucocorticosteroids are moderately recommended for treatment of acute, subacute, or chronic CTS among patients who decline carpal tunnel injection.

**Indication** – CTS unresponsive to splinting. Most patients should be injected rather than given oral steroids.(648) However, for patients declining injection, oral glucocorticosteroids may be warranted. Oral glucocorticosteroids are relatively contraindicated for patients with diabetes mellitus and may worsen glucose intolerance among those who are pregnant.

**Frequency/Dose** – It is unclear what dose and duration of treatment is optimal. Two trials used 10 days of treatment with prednisolone acetate 25mg a day.(646, 648) A third used prednisolone 20mg a day for 2 weeks, then 10mg a day for 2 weeks.(636, 647) Another used prednisone 20mg a day for 1 week, then 10mg a day for 1 week.(644) Another used prednisolone 20mg a day for 2 weeks on one treatment arm.(643) There is evidence that 2 weeks of treatment is as effective as 4 weeks.(643) It is recommended that one course (10 to 14 days) of oral glucocorticosteroid be prescribed rather than repeated courses. Prescriptions of low rather than high doses are recommended to minimize potential for adverse effects.

- **Strength of Evidence** – Moderately Recommended, Evidence (B)
- **Level of Confidence** – Moderate
- (Note: Injections are recommended as superior to oral forms.)

**Rationale for Recommendation**
There is strong evidence that injected glucocorticosteroids are more effective(648) with longer duration of benefits. Nevertheless, there is consistent evidence that oral glucocorticosteroids are superior to placebo,(636, 643, 644, 646) as well as compared with diuretics and NSAIDs.(636) Unlike glucocorticosteroid injections, long-term follow-up studies have not been reported, thus duration of benefit is unclear. However, oral glucocorticosteroids are not invasive, have relatively few adverse effects for a short course, and are low cost.

**Evidence for the Use of Oral Glucocorticosteroids**
See Intracarpal Tunnel Glucocorticosteroid Injections (“Steroid Injections”) Section.

**Diuretics**
Diuretics have been used to treat CTS, in part due to observations of swelling in some patients.(359, 634, 636, 649-652)

**Recommendation:** Diuretics for Treatment of Acute, Subacute, or Chronic CTS

Diuretics are moderately not recommended for treatment of acute, subacute, or chronic CTS in the absence of fluid retention states.

- **Strength of Evidence** – Moderately Not Recommended, Evidence (B)
- **Level of Confidence** – Moderate

**Rationale for Recommendation**
There are two quality studies evaluating diuretics for treatment of CTS patients and both failed to find evidence of efficacy compared with placebo.(636, 652) Thus, diuretics are not recommended for routine treatment of CTS patients. Whether they are effective for treatment of patients with CTS accompanied by
fluid retention states, such as third trimester pregnancy, has not been determined in quality studies, and thus their use in select cases may be a reasonable intervention.

Evidence for the Use of Diuretics for CTS
There are 2 moderate-quality RCTs incorporate into this analysis. (636, 652)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Diuretics, Trichlormethiazide, Hydrochlorothiazide, carpal tunnel syndrome, CTS, median nerve neuropathy, median neuropathy, median nerve disease, entrapment, neuropathy, nerve compression, burning, itching, numbness, tingling, wrist, hand, palm, finger, pain, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, and prospective studies. We found and reviewed 14 articles in PubMed, 1556 in Scopus, 27 in Cochrane Library and 2 in other sources. We considered for inclusion 2 from PubMed, 1 from Scopus, 0 from CINAHL, 1 from Cochrane Library and 2 from other sources. Of the 6 articles considered for inclusion, 2 randomized trials and 4 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chang 1998</td>
<td>RCT</td>
<td>Sponsored by the National Science Council Grants. No mention of COI.</td>
<td>7.0</td>
<td>N = 91 (53 female/20 male) with confirmed CTS via electrodiagnosis; Mean (±SD) age 44.2 (±5.4) for placebo group, 45.7 (±4.8) for diuretic group, 47.4 (±5.7) for NSAID-SR group and 45.4 (±5.2) for steroid group.</td>
<td>Trichlor-methiazide, 2mg daily (n = 16) vs. Tenoxicam-SR, 20mg daily (n = 18) vs. prednisolone at 20mg daily, followed by 2-week dose 10mg daily (n = 23) vs. Placebo or control group (n = 16). Assessments at baseline, 2 and 4 weeks.</td>
<td>No significant reduction from baseline GSS seen at 2nd and 4th weeks in placebo, NSAID-SR, and diuretic groups. However, mean score at 4 weeks in steroid group decreased significantly from a baseline of 27.9±6.9 to 10±7.54, (p &lt; 0.00001).</td>
<td>“For patients with mild to moderate CTS who opt for conservative treatment, corticosteroids are of greater benefit.”</td>
<td>Suggests oral steroids effective but diuretic and NSAID are not.</td>
</tr>
<tr>
<td>Pal 1988</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>6.0</td>
<td>N = 48 (43 female/5 male) with CTS diagnosed via nerve conduction tests; Mean (±SD) age 41 (±13) for Bendrofluazid group and 53 (±13) for placebo control group.</td>
<td>Bendrofluazide 5 mg a day (n = 23; 41 hands) vs. Placebo (N =25; 40 hands) for 4 weeks. Assessments at baseline, 4 weeks and 6 months.</td>
<td>No significant difference in clinical improvement outcomes between the two groups at follow-up assessments.</td>
<td>“Bendrofluazide 5mg daily for one month does not confer additional clinical benefit in the idiopathic CTS, but further trials with stronger diuretics and/or longer periods of treatment are warranted.”</td>
<td>Study suggests no short or long-term benefit.</td>
</tr>
</tbody>
</table>

**OPIOIDS – Oral, Transdermal, and Parenteral (includes Tramadol)**
Opioids have occasionally been used to treat pain for patients with CTS. Opioids are addressed in a separate Guideline. The treatment recommendations are summarized below. See Opioids Guideline for all supporting evidence.

**Acute Pain (Up to 4 Weeks)**

1. **Recommendation: Routine Use of Opioids for Treatment of Non-Severe Acute Pain**
Routine opioid use is strongly not recommended for treatment of non-severe acute pain (e.g., low back pain (LBP), sprains, or minor injury without signs of tissue damage).
2. Recommendation: Opioids for Treatment of Acute, Severe Pain

Opioids are recommended for treatment of acute, severe pain (e.g., crush injuries, large burns, severe fractures, injury with significant tissue damage) uncontrolled by other agents and/or with functional deficits caused by pain. They also may be indicated at the initial visit for a brief course for anticipated pain accompanying severe injuries (i.e., failure of other treatment is not mandatory). A Schedule IV opioid may be indicated if there is true allergy to NSAIDs and acetaminophen, other contraindication to an alternative medication, or insufficient pain relief with an alternative. Recommend to taper off opioid use in 1 to 2 weeks.

Indications – Patients should meet all of the following:

1) Severe injury with a clear rationale for use (objective functional limitations due to pain resulting from the medical problem, e.g., extensive trauma such as forearm crush injury, large burns, severe radiculopathy).

2) Other more efficacious treatments should have been instituted, and either: a) failed; and/or b) have reasonable expectations of the immediate need for an opioid to obtain sleep the evening after the injury.

3) Where available, prescription databases (usually referred to as Prescription Drug Monitoring Program (PDMP)) should be checked and not show evidence for conflicting opioid prescriptions from other providers or evidence of misreporting.

4) Non-opioid prescriptions (e.g., NSAIDs, acetaminophen) absent contraindication(s) should nearly always be the primary treatment and accompany an opioid prescription.

5) Low-dose opioids may be needed in the elderly who have greater susceptibility to the adverse effects of opioids. Those of lower body weight may also require lower opioid doses.

6) Dispensing quantities should be only what is needed to treat the pain. Short-acting opioids are recommended for treatment of acute pain. Long-acting opioids are not recommended.

7) Due to greater than 10-fold elevated risks of adverse effects and death, considerable caution is warranted among those using other sedating medications and substances including: i) benzodiazepines, ii) anti-histamines (H1-blockers), and/or iii) illicit substances. Patients should not receive opioids if they use illicit substances unless there is objective evidence of significant trauma or moderate to severe injuries. Considerable caution is also warranted among those who are unemployed as the reported risks of death are also greater than 10-fold.

Due to elevated risk of death and adverse effects, caution is also warranted when considering prescribing an opioid for patients with any of the following characteristics: depression, anxiety, personality disorder, untreated sleep disorders, substance abuse history, current alcohol use or current tobacco use, attention deficit hyperactivity disorder (ADHD), post-traumatic stress disorder (PTSD), suicidal control problems, thought disorders, psychotropic medication use, chronic obstructive pulmonary disease (COPD), asthma, or recurrent pneumonia. Considerable caution is also warranted among those with other comorbidities such as chronic hepatitis and/or cirrhosis, as well as coronary artery disease.

---

1USA classifies controlled substances that includes a classification system, ranging from Class I to Class V corresponding to lower risks of abuse and dependence. Class I includes substances with a high potential for abuse and without a recognized medical use (e.g., heroin, marijuana, LSD). Class II includes most opiates, amphetamines and cocaine. Class III includes buprenorphine, dihydromorphone, hydrocodone/codeine when compounded with an NSAID, Marinol. Class IV includes tramadol (in some states), carisoprodol, benzodiazepines, and long-acting barbiturates. Class V includes small amounts of codeine (e.g, 30mg, 60mg).

2Other indications beyond the scope of this guideline include acute myocardial infarction or agitation interfering with acute trauma management.

3Treatments to have tried generally include NSAIDs and acetaminophen. For LBP patients, additional considerations include muscle relaxants, progressive aerobic exercise, and directional exercise.

4Exceptions such as acute, severe trauma should be documented.
dysrhythmias, cerebrovascular disease, orthostatic hypotension, asthma, recurrent pneumonia, thermoregulatory problems, advanced age (especially with mentation issues, fall risk, debility), osteopenia, osteoporosis, water retention, renal failure, severe obesity, testosterone deficiency, erectile dysfunction, abdominal pain, gastroparesis, constipation, prostatic hypertrophy, oligomenorrhea, pregnancy, human immunodeficiency virus (HIV), ineffective birth control, herpes, allodynia, dementia, cognitive dysfunction and impairment, gait problems, tremor, concentration problems, insomnia, coordination problems, and slow reaction time. There are considerable drug-drug interactions that have been reported (see Appendices 2-3 of Opioids Guideline).

**Frequency/Duration** – Generally, opioids should be prescribed at night or while not working. (680) Lowest effective, short-acting opioid doses are preferable as they tend to have the better safety profiles, less risk of escalation, (681) less risk of lost time from work, (682) and faster return to work. (683) Short-acting opioids are recommended for treatment of acute pain and long-acting opioids are not recommended. Recommend opioid use as required by pain, rather than in regularly scheduled dosing.

If parenteral administration is required, ketorolac has demonstrated superior efficacy compared with opioids for acute severe pain, (684, 685) although ketorolac’s risk profile may limit use for some patients. Parenteral opioid administration outside of obvious acute trauma or surgical emergency conditions is almost never required, and requests for such treatment are clinically viewed as red flags for potential substance abuse.

**Indications for Discontinuation** – Resolution of pain, sufficient improvement in pain, intolerance or adverse effects, non-compliance, surreptitious medication use, consumption of medications or substances advised to not take concomitantly (e.g., sedating medications, alcohol, benzodiazepines), or use beyond 2 weeks.

**Harms** – Adverse effects are many (see section below on “Opioids Benefits and Harms”).

**Benefits** – Improved short-term pain control.

- **Strength of Evidence** – Recommended, Evidence (C)
- **Level of Confidence** – High

3. **Recommendation: Screening Patients Prior to Initiation of Opioids**

   Initial screening of patients is recommended with more detailed screening for: i) requiring continuation of opioids beyond 2 weeks for those with an acute severe injury, and ii) at consideration of initiation for severe pain but no objective evidence. Screening should include history(ies) of depression, anxiety, personality disorder, other psychiatric disorder, substance abuse, sedating medication use (e.g., anti-histamine/anti-H1 blocker(654)), benzodiazepine use, opioid dependence, alcohol abuse, current tobacco use, other substance use history, COPD, PTSD, other psychotropic medications, (severe) obesity, cognitive impairment, balance problems/fall risk, osteoporosis, and renal failure (see Appendix 1 of Opioids Guideline). Those who screen positive, especially to multiple criteria, are recommended to: i) undergo greater scrutiny for appropriateness of opioids (may include psychological evaluation), ii) consideration of consultation and examination(s) for complicating conditions and/or appropriateness of opioids, and iii) if opioids are prescribed, more frequent assessments for compliance, achievement of functional gains,(655, 686, 687) adverse effects, and symptoms and signs of aberrancy.

**Harms** – Negligible. If a consultation is needed, there are additional costs that are incurred.

**Benefits** – Improved identification of more appropriate candidates for opioids. Identification of patients at increased risk of adverse effects. In cases where someone has elevated, but potentially acceptable risk, may alert the provider to improve surveillance for complications and aberrant behaviors.

- **Strength of Evidence** – Recommended, Insufficient Evidence (I)
- **Level of Confidence** – High

4. **Recommendation: Opioid Dose Limits in Acute Pain**

Dispense only that which is required. The maximum daily oral dose recommended for opioid-naive, acute pain patients based on risk of overdose/death is 50mg morphine equivalent dose
In rare cases with documented functional improvement (see Appendix 1 of Opioids Guideline), higher doses may be considered, however, risks are substantially higher and greater monitoring is also recommended (see Subacute/Chronic Opioid recommendations below). Lower doses should be used for patients at higher risk of dependency, addiction and other adverse effects. Monitoring is also recommended and consultation may be considered for those patients on higher doses.

**Harms** – Theoretical potential to undertreat pain in some patients with increased pain sensitivity.

**Benefits** – Reduced risk for adverse physical and cognitive effects, dependency, addiction and opioid-related overdoses and deaths.

*Strength of Evidence – Recommended, Evidence (C)*

*Level of Confidence – Moderate*

---

**Figure 4. Death Rate (Hazard Ratio) vs. Morphine Equivalent Dosage (mg/d)**

Adapted from Dunn 2010 and Bohnert 2011.

*Statistical significance present for acute and chronic pain at and above 50 mg per day of oral morphine equivalent dose.

**Post-Operative Pain (Up to 4 Weeks) (After 4 weeks, see Subacute Pain)**

Oral opioids are commonly prescribed after sinus surgery, (689) major noncardiac surgical procedures, (690) mastectomy and immediate breast reconstruction (IBR), (691, 692) coronary artery bypass graft surgery, (693) major abdominal surgery (abdominal laparoscopic, abdominal hysterectomy, bowel resection or radical hysterectomy), (694-697) orthopedic surgery, (698) and molar extraction. (699)

1. **Recommendation: Limited Use of Opioids for Post-operative Pain**

   **Limited use of opioids is recommended for post-operative pain management as adjunctive therapy to more effective treatments.**

   **Indications** – For post-operative pain management, a brief prescription of short-acting opioids as adjunct to more efficacious treatments (especially Cox-2 NSAIDs such as celecoxib, non-selective NSAIDs after risk of bleeding is no longer a concern). (vi) A brief course of opioids is often needed for minor surgical procedures. However, minor wound laceration repairs often require no opioids. Evidence suggests perioperative pregabalin for 14 days and/or continuous femoral nerve catheter analgesia instead of solely using oral opioids results in superior knee arthroplasty functional outcomes with less venous thromboses. (700) Additional considerations include:

---

viStatistical significance present for acute and chronic pain at and above 50 mg per day of oral morphine equivalent dose.

viMore efficacious treatments also include therapeutic exercises, e.g., progressive ambulation especially for moderate to extensive procedures (e.g., arthroplasty, fusion).
1) Non-opioid prescriptions (e.g., NSAIDs, acetaminophen) should nearly always be the primary treatment and accompany an opioid prescription. Computerized programs may also assist in optimal management.(701)

2) The lowest effective dose of a short-acting opioid should be used,(681) as well as weaker opioids if possible.(682, 683)

3) Short-acting opioids are recommended for treatment of acute pain.

4) Dispensing should be only what is needed to treat the pain.\(^{vi}\)

5) Long-acting opioids are not recommended.

6) Low-dose opioids may be needed in the elderly who have greater susceptibility to the adverse risks of opioids. Those of lower body weight may also require lower opioid doses.

7) Where available, prescription databases (usually referred to as Prescription Drug Monitoring Program (PDMP)) should be checked for other opioid prescriptions. Due to greater than 10-fold elevated risks of adverse effects and death, considerable caution is warranted among those using other sedating medications and substances including: i) benzodiazepines, ii) anti-histamines (H\(_1\)-blockers), and/or iii) illicit substances.(653-656) Patients should not receive opioids if they use illicit substances unless there is objective evidence of significant trauma or moderate to severe injuries. Considerable caution is also warranted among those who are unemployed as the reported risks of death are also greater than 10-fold.(654, 655)

Due to elevated risk of death and adverse effects, caution is also warranted when considering prescribing an opioid for patients with any of the following characteristics: depression, anxiety, personality disorder, ADHD, PTSD, suicidal risk, impulse control problems, thought disorders, psychotropic medication use, substance abuse history, current alcohol use or current tobacco use, untreated sleep disorders, COPD, asthma, or recurrent pneumonia.\(^{654, 657-678}\) Considerable caution is also warranted among those with other comorbidities such as chronic hepatitis and/or cirrhosis,(679) as well as coronary artery disease, dysrhythmias, cerebrovascular disease, orthostatic hypotension, thermoregulatory problems, advanced age (especially with mentation issues, fall risk, debility), osteopenia, osteoporosis, water retention, renal failure, severe obesity, testosterone deficiency, erectile dysfunction, abdominal pain, gastroparesis, constipation, prostatic hypertrophy, oligomenorrhea, pregnancy, HIV, ineffective birth control, herpes, allodynia, dementia, cognitive dysfunction and impairment, gait problems, tremor, concentration problems, insomnia, coordination problems, and slow reaction time. There are considerable drug-drug interactions that have been reported (see Appendices 2-3 of Opioids Guideline). Inpatient management may moderate these recommendations provided there is careful monitoring, although these same management issues then apply post-discharge.

8) For patients taking opioids chronically prior to surgery, consultations with anesthesiology and/or pain management are generally needed as post-operative dosing may be very high and management is often quite challenging.

9) Ongoing prescriptions of opioids after the immediate post-operative period should generally be for patients who have undergone a major surgery or have other condition(s) necessitating opioids. Most patients should be making progress towards functional restoration, pain reduction and weaning off the opioids. Patients who have not progressed should be carefully evaluated for physical complications or psychiatric comorbidity, adherence to active treatments, and pending development of addiction or dependency.

\textit{Frequency/Duration} – For moderate and major surgeries, opioids are generally needed on a scheduled basis in the immediate post-operative period. Other post-operative situations may be sufficiently managed with an as needed opioid prescription schedule. Provision of opioids sufficient to participate in therapeutic exercise (e.g., progressive ambulation) and allow sleep may be needed. However, high dose use at night is not recommended due to respiratory depression and disruption of sleep architecture. Weaning should begin as soon as function is recovering and pain is subsiding. Subsequent weaning to as needed opioid use is recommended.

\(^{vi}\)Generally, this should be sufficient to cover two weeks of treatment. Prescriptions of 90-day supplies in the post-operative setting are not recommended.
Indications for Discontinuation – The physician should discontinue the use of opioids based on sufficient recovery, expected resolution of pain, lack of efficacy, intolerance or adverse effects, non-compliance, surreptitious medication use, self-escalation of dose, or use beyond 3 to 5 days for minor procedures, and 2 to 3 weeks for moderate/less extensive procedures. Use for up to 3 months may occasionally be necessary during recovery from more extensive surgical procedures (e.g., spine fusion surgery). However, with rare exceptions, only nocturnal use is recommended in months 2 to 3 plus institution of management as discussed in the subacute/chronic guidelines below. For those requiring opioid use beyond 1 month, the subacute/chronic opioid use recommendations below apply.

Harms – Adverse effects are many (see section on “Opioids Benefits and Harms”).

Benefits – Improved short-term, post-operative pain control. Some studies suggest this may modestly improve functional outcomes in the post-operative population.

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – High

2. Recommendation: Screening Patients Prior to Continuation of Opioids

Screening of patients is recommended for patients requiring continuation of opioids beyond the second post-operative week. Screening should include history(ies) of: depression, anxiety, personality disorder, pain disorder, other psychiatric disorder, substance abuse history, sedating medication use (e.g., anti-histamine/anti-H₁ blocker), benzodiazepine use, opioid dependence, alcohol abuse, current tobacco use, and other substance use history, COPD, PTSD, other psychotropic medications, (severe) obesity, cognitive impairment, balance problems/fall risk, osteoporosis, and renal failure (see Appendix 1 of Opioids Guideline). Those who screen positive, especially to multiple criteria, are recommended to: i) undergo greater scrutiny for appropriateness of opioids (e.g., may include psychological and/or pain evaluation); ii) compliance with active therapies (e.g., ambulation and other exercise after arthroplasty); iii) consider consultation examination(s) for complicating conditions and/or appropriateness of opioids; and iv) if ongoing opioids are prescribed, ensure more frequent assessments for treatment compliance, achievement of functional gains, (655, 686, 687) and symptoms and signs of aberrancy.

Harms – Negligible. If a consultation is needed, there are additional costs that are incurred.

Benefits – Identification of patients at increased risk of adverse effects. Improved identification of more appropriate and safe candidates for opioids compared with attempting post-operative pain control with non-opioids. This should reduce adverse effects. In cases where someone has elevated, but potentially acceptable risk, this may alert the provider to improve surveillance for complications and aberrant behaviors.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – High

3. Recommendation: Opioid Dose Limits in Post-operative Pain

The maximum daily oral dose recommended for opioid-naive, acute pain patients based on risk of overdose/death is 50mg morphine equivalent dose (MED)(688) (see Figure 4). Post-operative patients particularly require individualization due to factors such as the severity of the operative procedure, response to treatment(s) and variability in response. Higher doses beyond 50mg MED may be particularly needed for major surgeries in the first two post-operative weeks to achieve sufficient pain relief, however, greater caution and monitoring are warranted and reductions below 50mg MED at the earliest opportunity should be sought. Lower doses should be used for patients at higher risk of dependency, addiction and other adverse effects. In rare cases with documented functional improvement, ongoing use of higher doses may be considered, however, risks are substantially higher and greater monitoring is also recommended (see Subacute/Chronic Opioid recommendations below).

Harms – Theoretical potential to undertreat pain, which could modestly delay functional recovery.

Benefits – Reduced risk for adverse effects, dependency, addiction and opioid-related deaths.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Low

---

viStatistical significance present for acute and chronic pain at and above 50 mg per day of morphine equivalent dose.
Subacute (1-3 Months) and Chronic Pain (>3 Months)

1. **Recommendation: Routine Use of Opioids for Subacute and Chronic Non-malignant Pain**
   Opioid use is moderately not recommended for treatment of subacute and chronic non-malignant pain. Opioid prescription should be patient specific and limited to cases in which other treatments are insufficient and criteria for opioid use are met (see below).

   - **Harms** – May inadequately treat severe subacute or chronic pain.
   - **Benefits** – Less debility, fewer adverse effects, reduced accident risks, lower risks of dependency, addiction, overdoses, and deaths.

   - **Strength of Evidence** – Moderately Not Recommended, Evidence (B)
   - **Level of Confidence** – High

2. **Recommendation: Opioids for Treatment of Subacute or Chronic Severe Pain**
   The use of an opioid trial is recommended if other evidence-based approaches for functional restorative pain therapy have been used with inadequate improvement in function.\(^{(702, 703)}\)

   Opioids are then recommended for treatment of function impaired by subacute or chronic severe pain (e.g., inability to work due to any of the following: chronic severe radiculopathy, chronic severe peripheral neuropathies, complex regional pain syndrome (CRPS), and severe arthroses)\(^{(686)}\) (see Appendix 1 of Opioids Guideline).

   - **Indications** – Patients should meet all of the following:
     1) Reduced function is attributable to the pain. Pain or pain scales alone are insufficient reasons.\(^{(655, 686, 704-715)}\)
     2) A severe disorder warranting potential opioid treatment is present [e.g., CRPS, severe radiculopathy, advanced degenerative joint disease (DJD)].\(^{(706)}\)
     3) Other more efficacious treatments have been documented to have failed.\(^{(706)}\) Other approaches that should have been first utilized include physical restorative approaches, behavioral interventions, self-applied modalities, non-opioid medications (including NSAIDs, acetaminophen, topical agents, norepinephrine adrenergic reuptake blocking antidepressants or dual reuptake inhibitors; also antiepileptic medications particularly for neuropathic pain) and functional restoration. For LBP patients, this also includes\(^{ix}\) fear avoidant belief training and ongoing progressive aerobic exercise, and strengthening exercises. For CRPS patients, this includes progressive strengthening exercise. For DJD, this includes NSAIDs, weight loss, aerobic and strengthening exercises.
     4) An ongoing active exercise program is prescribed and complied with.
     5) Non-opioid prescriptions (e.g., NSAIDs, acetaminophen) absent a contraindication should nearly always be the primary pain medication and accompany an opioid prescription. Other medications to consider include topical agents, norepinephrine adrenergic reuptake blocking antidepressants or dual reuptake inhibitors; also antiepileptic medications particularly for neuropathic pain).
     6) The lowest effective dose should be used.\(^{(681)}\) Weaker opioids should be used whenever possible.\(^{(682, 683)}\) Meperidine is not recommended for chronic pain due to bioaccumulation and adverse effects.
     7) Low-dose opioids may be needed in the elderly who have greater susceptibility to the adverse risks of opioids. Those of lower body weight may also require lower opioid doses.
     8) Dispensing should be only what is needed to treat the pain.\(^{x}\)
     9) Extended-release/long-acting opioids are recommended to be used on a scheduled basis, rather than as needed.\(^{(706)}\) As needed opioids should generally be avoided for treatment of chronic pain, although limited use for an acute painful event (e.g., fracture, sprain) is reasonable. Sublingual fentanyl is not recommended for treatment of subacute or chronic pain. Caution is warranted with fentanyl patches due to unpredictable absorption.
     10) Where available, prescription databases (usually referred to as Prescription Drug Monitoring

---

\(^{ix}\)A previous trial of a muscle relaxant is generally recommended. However, if an opioid trial is contemplated, cessation of all depressant medications including muscle relaxants is advisable.

\(^{x}\)Generally, this should be sufficient to cover one week of treatment at a time during the trial phase. If a trial is successful at improving function, prescriptions for up to 90-day supplies are recommended.
Program (PDMP) should be checked for conflicting opioid prescriptions from other providers or evidence of misreporting.

11) Due to greater than 10-fold elevated risks of adverse effects and death, considerable caution is warranted among those using other sedating medications and substances including: i) benzodiazepines, ii) anti-histamines (H1-blockers), and/or iii) illicit substances. (653-656) Patients should not receive opioids if they use illicit substances unless there is objective evidence of significant trauma or moderate to severe injuries. Considerable caution is also warranted among those who are unemployed as the reported risks of death are also greater than 10-fold. (654, 655) Due to elevated risk of death and adverse effects, caution is also warranted when considering prescribing an opioid for patients with any of the following characteristics: depression, anxiety, personality disorder, untreated sleep disorders, substance abuse history, current alcohol use or current tobacco use, ADHD, PTSD, suicidal risk, impulse control problems, thought disorders, psychotropic medication use, COPD, asthma, recurrent pneumonia. (654, 657-678) Considerable caution is also warranted among those with other comorbidities such as chronic hepatitis and/or cirrhosis, (679) as well as coronary artery disease, dysrhythmias, cerebrovascular disease, orthostatic hypotension, asthma, recurrent pneumonia, thermoregulatory problems, advanced age (especially with mentation issues, fall risk, debility), osteopenia, osteoporosis, water retention, renal failure, severe obesity, testosterone deficiency, erectile dysfunction, abdominal pain, gastroparesis, constipation, prostatic hypertrophy, oligomenorrhea, pregnancy, HIV, ineffective birth control, herpes, allodynia, dementia, cognitive dysfunction and impairment, gait problems, tremor, concentration problems, insomnia, coordination problems, and slow reaction time. There are considerable drug-drug interactions that have been reported (see Appendices 2-3 of Opioids Guideline).

Frequency/Duration – Opioids use is generally initiated as a “trial” to ascertain whether the selected opioid produces functional improvement (see Appendix 1 of Opioids Guideline). Opioid use is generally prescribed on a regular basis, (716) at night or when not at work. (680) Only one opioid is recommended to be prescribed in a trial. More than one opioid should rarely be used. Lower opioid doses are preferable as they tend to have the better safety profiles, less risk of dose escalation, (681) less work loss, (682) and faster return to work. (683) Patients should have ongoing visits to monitor efficacy, adverse effects, compliance and surreptitious medication use. Opioid prescriptions should be shorter rather than longer duration. (717)

Indications for Discontinuation – Opioids should be discontinued based on lack of functional benefit (703) (see Appendix 1 of Opioids Guideline), resolution of pain, improvement to the point of not requiring opioids, intolerance or adverse effects, non-compliance, surreptitious medication use, medication misuse (including self-escalation and sharing medication), aberrant drug screening results, diversion, consumption of medications or substances advised to not take concomitantly (e.g., sedating medications, alcohol, benzodiazepines).

Harms – Adverse effects are many (see section on “Opioids Benefits and Harms”). May initiate path to opioid dependency.


Strength of Evidence – Recommended, Insufficient Evidence (I)

Level of Confidence – Low

3. Recommendation: Screening Patients Prior to Initiation of Opioids

Screening of patients is recommended prior to consideration of initiating a trial of opioids for treatment of subacute or chronic pain. Screening should include history(ies) of depression, anxiety, personality disorder and personality profile, (683, 718, 719) other psychiatric disorder, substance abuse history, sedating medication use (e.g., anti-histamine/anti-H1 blocker), (667) benzodiazepine use, opioid dependence, alcohol abuse, current tobacco use, and other substance use history, COPD, PTSD, other psychotropic medications, (severe) obesity, cognitive impairment, balance problems/fall risk, osteoporosis, and renal failure (see Appendix 1 of Opioids Guideline).
Those who screen positive, especially to multiple criteria, are recommended to: i) undergo greater scrutiny for appropriateness of opioids (may include psychological and/or psychiatric evaluation(s) to help assure opioids are not being used instead of appropriate mental health care); ii) consideration of consultation and examination(s) for complicating conditions and/or appropriateness of opioids; and iii) if opioids are prescribed, more frequent assessments for compliance, achievement of functional gains and symptoms and signs of aberrant use.

Harms – Negligible. If a consultation is needed, there are additional costs that are incurred. Benefits – Identification of patients at increased risk of adverse effects. Improved identification of more appropriate and safe candidates for treatment with opioids. This should reduce adverse effects. In cases where someone has elevated, but potentially acceptable risk, this may alert the provider to improve surveillance for complications and aberrant behaviors.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – High

4. Recommendation: Opioid Dose Limits in Subacute and Chronic Pain

The maximum daily oral dose recommended for subacute or chronic pain patients based on risk of overdose/death is 50mg Morphine Equivalent Dose (MED). In rare cases with documented functional improvements occurring with use above 50mg MED, subsequent doses up to 100mg may be considered, however, risks of death are much greater and more intensive monitoring is then also recommended. Lower doses should be considered in high-risk patients. Caution appears warranted in all patients as there is evidence the risk of dose escalation is present even among patients enrolled in a “hold the line (stable dose) prescribing strategy” treatment arm.

For those whose daily consumption is more than 50mg MED, greater monitoring is recommended to include: i) at least monthly to not more than quarterly appointments with greater frequencies during trial, dose adjustments and with greater co-morbid risk factors and conditions; ii) at least semiannual attempts to wean below 50mg MED if not off the opioid; iii) at least semiannual documentation of persistence of functional benefit; iv) at least quarterly urine drug screening (see drug screening section); and v) at least semiannual review of medications, particularly to assure no sedating medication use (e.g., benzodiazepine, sedating anti-histamines).

Harms – None in a short-term trial. For chronic pain patients, theoretical potential to undertreat pain and thus impair function. However, there is no quality literature currently available to support that position. Benefits – Reduced risk for adverse effects, dependency, addiction, and opioid-related deaths.

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – High


The use of an opioid treatment agreement (opioid contract, doctor/patient agreement, or informed consent) is recommended to document patient understanding, acknowledgement of potential adverse effects, and agreement with the expectations of opioid use (see Appendix 1 of Opioids Guideline). If consent obtained, it is recommended appropriate family members be involved in this agreement.

Harms – Negligible. Benefits – Educates the patient and significant others that these medications are high risk, with numerous adverse effects. It allows for a more informed choice. It provides a framework for initiation of a trial, monitoring, treatment goals, compliance requirement, treatment expectations, and conditions for opioid cessation. It should reduce risk of adverse events and opioid-related deaths, although that remains unproven to date.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate
6. Recommendation: Urine Drug Screening

Baseline and random urine drug screening, qualitative and quantitative, is recommended for patients prescribed opioids for the treatment of subacute or chronic pain to evaluate presence or absence of the drug, its metabolites, and other substance(s) use. In certain situations, other screenings (e.g., hair particularly for information regardingremote use or blood (for acute toxicity) may be appropriate.

Indications – All patients on opioids for subacute or chronic pain.

Frequency – Screening is recommended at baseline, randomly at least twice, and up to 4 times a year and at termination. More intensive screening is recommended for those consuming more than 50mg MED (see above). Federal guidelines recommend at least 8 tests a year among those utilizing opioid treatment programs.(739) Screening should also be performed “for cause” (e.g., provider suspicion of substance misuse including over-sedating, drug intoxication, motor vehicle crash, other accidents and injuries, driving while intoxicated, premature prescription renewals, self-directed dose changes, lost or stolen prescriptions, using more than one provider for prescriptions, non-pain use of medication, using alcohol for pain treatment or excessive alcohol use, missed appointments, hoarding of medications, and selling medications). Standard urine drug/toxicology screening processes should be followed (consult a qualified medical review officer).(740-742) If there is an aberrant drug screen result (either positive for unexpected drugs or unexpected metabolites or unexpectedly negative results), there should be a careful evaluation of whether there is a plausible explanation (e.g., drug not tested, drug metabolite not tested, laboratory cutpoint and dosing interval would not capture the drug/metabolite, laboratory error). In the absence of a plausible explanation, those patients with aberrant test results should have the opioid discontinued or weaned.(703)

Harms – No adverse clinical effects if properly interpreted.

Benefits – Identifies aberrant medication(s) and substance(s) use. Such uses are high-risk for opioid events including fatalities (see tables below). It provides objective evidence to cease an opioid trial or ongoing treatment. Identifies patients who may be diverting medication (those screening negative for prescribed medication).

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – High

Evidence for the Use of Opioids
See Opioids Guideline.

VITAMINS (Including Pyridoxine)

Treatment of CTS with pyridoxine (Vitamin B6) has been attempte(634, 650, 743-746) as there has been some association between pyridoxine deficiencies and peripheral neuropathies, as well as some reports of associations of deficiencies with CTS in some,(747) but not all studies.(748) Vitamin B12 has also been reported as a successful treatment for stroke patients with CTS.(749)

1. Recommendation: Pyridoxine for Treatment of Acute, Subacute, or Chronic CTS

Pyridoxine is not recommended for routine treatment of acute, subacute, or chronic CTS in patients without vitamin deficiencies.

Strength of Evidence – Not Recommended, Evidence (C)
Level of Confidence – Low

2. Recommendation: Other Vitamins for Treatment of Acute, Subacute, or Chronic CTS

There is no recommendation for or against the use of other vitamins for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low
Rationale for Recommendations
There are two quality studies that reviewed pyridoxine to treat CTS patients. However, benefits have not been shown in the highest quality study.(745) The moderate-quality crossover trial reported improvements in symptoms in 7 patients; however, 3 patients did not receive the placebo although their symptoms scores on pyridoxine were lower than in a control period.(743) While vitamin B-6 is relatively low risk and patients may use it without prescription, available evidence does not support its use for the routine treatment of CTS, thus it is not recommended. However, it may be a reasonable treatment option among patients with presumptive pyridoxine deficiency (e.g., malnutrition, alcoholism, malabsorption, especially jejunal disorders such as sprue, etc.).

Evidence for the Use of Pyridoxine for CTS
There is 1 high-quality RCT(745) and 1 moderate-quality randomized crossover trial(743) incorporated into this analysis. There is 1 low-quality RCT in Appendix 2.(746)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: vitamin B6, Vitamin B12, Pyridoxine, carpal tunnel syndrome, CTS, median nerve neuropathy, median neuropathy, median nerve disease, entrapment, neuropathy, nerve compression, burning, itching, numbness, tingling, randomized controlled trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 15 articles in PubMed, 3,114 in Scopus, 6 in CINAHL, 251 in Cochrane Library and 0 in other sources. We considered for inclusion 4 from PubMed, 0 from Scopus, 1 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 5 articles considered for inclusion, 3 randomized trials and 2 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spooner 1993 RCT</td>
<td>Sponsored by the Clinical Teaching and Research Fund of the College of Medicine at the University of Saskatchewan in Saskatoon. No mention of COI.</td>
<td>8.5</td>
<td>N = 35 (22 female/ 13 male) with CTS, EDS confirmed; mean age 42.5 years.</td>
<td>200mg pyridoxine once daily (n = 18) vs. Placebo (N = 17) for 12 weeks. Assessment at 6 and 12 weeks.</td>
<td>Mean score (SD) of night discomfort symptoms in treatment group: Entrance: 2.4 (1.4); 12 weeks: 1.9 (1.2) vs. control: Entrance 2.6 (1.3); 12 weeks: 2.4 (1.3), NS. Mean score of median palmar distal latency (ms) in treatment group: Entrance 2.5 (0.6); 12 weeks: 2.6 (0.4) vs. control: Entrance 2.8 (0.6); 12 weeks: 2.7 (0.4), NS. Mean (SD) swelling treatment: entrance 2.1 (1.6); 12 weeks: 1.3 (1.4) vs. control: entrance 2.6 (1.3); 12 weeks: 2.3 (1.2) (p &lt;0.05). Mean (SD) movement discomfort treatment: 3.1 (1.2); 1.7 (1.4) vs. control: 3.1 (1.3); 2.7 (1.3) (p &lt;0.001).</td>
<td>“Our findings do not support the use of pyridoxine for treating carpal tunnel syndrome.”</td>
<td>No statistical differences. Symptoms trended in favor of pyridoxine.</td>
</tr>
<tr>
<td>Ellis 1982 RCT Crossover Trial</td>
<td>Sponsored by Robert A Welch Foundation. No mention of COI.</td>
<td>6.5</td>
<td>N = 7 males with evidence of entrapment of median nerve, symptoms in ulnar nerve region with or without evidence of CTS, median nerve neuropathy, median neuropathy, median nerve disease, entrapment neuropathy, nerve compression, burning, itching, numbness, tingling</td>
<td>Pyridoxine 50mg vs Placebo for 12 weeks.</td>
<td>Aggregate mean symptom scores control 53 ± 10 (n = 4) vs. pyridoxine 11 ± 6 (n = 7), p &lt;0.001.</td>
<td>“Clinical improvements of the syndrome with pyridoxine therapy may frequently obviate hand surgery.”</td>
<td>Small sample size. Variable timeframes for measurements limit strength of conclusions.</td>
</tr>
</tbody>
</table>
TOPICAL MEDICATIONS: LIDOCAINE PATCHES
Topical lidocaine patches have been increasingly used to treat numerous pain conditions through transdermal application of topical anesthetic. (638, 750, 751)

Recommendation: Lidocaine Patches for Treatment of Acute, Subacute, or Chronic CTS
Lidocaine patches are recommended for treatment of select cases of acute, subacute, or chronic CTS with pain.

Indications – Moderate to severe CTS with pain as a central complaint when other treatable causes of the pain have been eliminated and after more efficacious treatment strategies, such as splinting and glucocorticosteroid injection(s), have been attempted.

Frequency/Duration – Usually 3 patches per day. Duration of use for chronic, localized pain may be as long as indefinitely, although most patients do not require indefinite treatment, as symptoms usually resolve, improve, or require surgery. Caution is warranted regarding widespread use of topical anesthetics for potential systemic effects from widespread administration. (752) For the hand this may require both patches and other applications or use in other body locations.

Indications for Discontinuation – Resolution, intolerance, adverse effects, lack of benefits, or failure to progress over a trial of at least 2 weeks.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
Topical lidocaine has been suggested to improve pain associated with CTS although the case diagnoses do not appear well substantiated in the available study as pain complaints as an overriding symptom among CTS patients raise concerns about alternate explanations for the symptoms. (751) In one moderate-quality study, lidocaine patches were suggested to be somewhat more effective than naproxen; (638) however, naproxen does not appear particularly effective and the study had a number of weaknesses. In the other study, injection was comparable to the patch, yet injections are likely a more effective strategy than naproxen, thus this body of evidence somewhat conflicts. Lidocaine patches are not invasive and have low adverse effects although some patients may experience local reactions such as skin irritation, redness, pain, or sores. These patches are also moderately or even high cost over time. While there are other lower cost topical treatments that provide analgesia (including heat, ice, and capsaicin), lidocaine patches may be a reasonable treatment option for pain related to CTS. Patients should be monitored to ensure that they are receiving benefit and to ascertain if there are any untoward local skin changes as a result of use.

Evidence for the Use of Topical Lidocaine Patches for CTS
There are 2 moderate-quality RCTs incorporated into this analysis. There are 2 low-quality RCTs in Appendix 2. (753, 754)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: lidocaine or lidocaine patch, carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, medean nerve, median neuropathy, burning, itching, numbness, tingling, hand, palm, finger, wrist, pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, retrospective, and prospective studies. We found and reviewed 56 articles in PubMed, 14 in Scopus, 2 in CINAHL, and 40 in Cochrane Library. We considered for inclusion 3 from PubMed, 1 from Scopus, 0 from CINAHL, Cochrane Library and other sources. Of the 4 articles considered for inclusion, 4 randomized trials and 0 systematic studies met the inclusion criteria.

Copyright© 2016 Reed Group, Ltd.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Compariso n Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nalamachu</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>4.5</td>
<td>N = 40 (28 female/12 male) neuropathic pain associated with CTS, Age 18-75.</td>
<td>Lidocaine patch 5%, up to 3, every 24 hours (n = 52) vs. Naproxen 500mg twice daily for 6 weeks (n = 48). Follow-up for 6 weeks.</td>
<td>Reductions in API scores between baseline and Week 6 for both lidocaine patch 5% (p &lt;0.0001) and naproxen (p = 0.0004), but no differences between treatments (p = 0.083). Significant difference in CGI-I for lidocaine patch 5% (51.1%) compared with naproxen 500mg 2x daily (24.3%) (p = 0.016); 71.8% lidocaine patch patients &quot;satisfied&quot; to &quot;very satisfied&quot; vs. 63.2% naproxen (NS).</td>
<td>This study demonstrates that the lidocaine patch 5% is effective in significantly relieving the pain associated with CTS and is well tolerated.</td>
<td>More diabetics in naproxen group (23.59% vs. 9.6%) suggest potential randomization failure and subsequent confounding. Severity (39.69% vs. 32.7%) and mean pain intensity somewhat worse in naproxen group (4.9±2.6 vs. 4.5±2.5). Excluded pain patch use, but not prior NSAID use. All appear to bias in favor of patch. Potentially, may have included treatment of other painful confounding diagnoses.</td>
</tr>
<tr>
<td>Nalamachu</td>
<td>RCT</td>
<td>Sponsored by Endo Pharmaceuticals. Dr. Nalamachu has served as consultant to Endo, Dr. Crockett is a statistician for B&amp;B Clinical Innovations.</td>
<td>4.5</td>
<td>N = 40 (28 female/12 male) electrodiagnostic evidence of CTS included median motor nerve distal latency &gt;4.10m sec. Mean age 48.</td>
<td>Lidocaine patch 5% (n = 20) vs methylprednisolone acetate 40mg depot injection (n = 20). Follow-up for 4 weeks.</td>
<td>Not significant between-group differences. Mean pain scores at 4 weeks: 2.2 patch vs. -2.1 injection (NS). Global improvements 88% patch vs. 74% injection.</td>
<td>“This pilot trial demonstrated that the lidocaine patch 5% was efficacious in reducing pain associated with CTS.”</td>
<td>Unclear whether patients had other painful diagnoses that explained the results.</td>
</tr>
</tbody>
</table>

**GABAPENTIN**

Gabapentin has been used to treat carpal tunnel syndrome.(755)

*Recommendation: Gabapentin for treatment of Carpal Tunnel Syndrome*

**Gabapentin is moderately not recommended for treatment of carpal tunnel syndrome.**

*Strength of Evidence – Moderately Not Recommended, Evidence (B)*

*Level of Confidence – Moderate*

*Rationale for Recommendation*

There is one high-quality, placebo-controlled study evaluating the use of gabapentin for treatment of CTS and finding it ineffective, thus gabapentin is moderately not recommended.

*Evidence for the Use of Gabapentin for CTS*

There is 1 high-quality RCT incorporated into this analysis.(755)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Gabapentin, Neurontin, Fanatrex, Gabarone, Neupentin, Neogab, Horizant, Gralise, carpal tunnel syndrome, CTS, median nerve neuropathy, median neuropathy, median nerve disease, entrapment, neuropathy, nerve compression, burning, itching,
numbness, tingling, wrist, hand, palm, finger, pain, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, and prospective studies. We found and reviewed 7 articles in PubMed, 627 in Scopus, 1 in CINAHL, 41 in Cochrane Library and 0 in other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 1 articles considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hui 2011</td>
<td>8.0</td>
<td>N = 140 (114 males/26 males) with diagnosed CTS lasting &gt;3 months; Mean (SD) age 52.3 (10.6) for gabapentin group and 51.0 (8.3) for placebo.</td>
<td>Gabapentin group receiving 300mg daily for first week, 600mg daily 2nd week and 900mg daily remaining treatment weeks (n = 71) vs. Placebo control group (n = 69). Assessments at baseline, 2 and 8 weeks.</td>
<td>During 2 and 8 weeks assessment, no significant difference reported between groups for global symptom scores reduction. Both groups showed improvement from baseline.</td>
<td>&quot;As gabapentin appears to have limited efficacy and would be required to be taken for a long time (because the majority of patients symptoms persist if left untreated), current evidence does not support its routine use for CTS&quot;</td>
<td>Gabapentin not effective.</td>
</tr>
</tbody>
</table>

**Physical Methods/Rehabilitation**

**Devices**

**MAGNETS**

Treatment of CTS and other hand, wrist, and forearm MSDs with magnets(756-758) and pulsed magnetic field therapy(759-761) has been attempted to manage pain.(612, 651, 762)

1. **Recommendation: Magnets for Management of Pain from of Acute, Subacute, or Chronic CTS**

MAGNETS are moderately not recommended for management of pain from acute, subacute, or chronic CTS.

*Strength of Evidence – Moderately Not Recommended, Evidence (B)*

*Level of Confidence – High*

2. **Recommendation: Pulsed Magnetic Field Therapy for Management of Pain from of Acute, Subacute, or Chronic CTS**

Pulsed magnetic field therapy is not recommended for management of pain from acute, subacute, or chronic CTS.

*Strength of Evidence – Not Recommended, Insufficient Evidence (I)*

*Level of Confidence – Moderate*

**Rationale for Recommendation**

Quality evidence suggests magnets(756, 757) are ineffective for treatment of CTS. Low-quality evidence suggests pulsed magnetic field therapy(759, 761) is not effective for treating CTS.(756) Magnets are not invasive, have no adverse effects, and are low cost, but other interventions have been shown effective. Thus, magnets are not recommended for treatment of CTS.

**Evidence for the Use of Magnets for CTS**

There are 1 high-(757) and 2 moderate-quality RCTs incorporated into this analysis.(756, 758) There are 3 low-quality RCTs in Appendix 2.(759-761)
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Magnet, pulsed magnetic field therapy, carpal tunnel syndrome, CTS, median nerve neuropathy, median nerve disease, entrapment, neuropathy, nerve compression, burning, itching, numbness, tingling, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 34 articles in PubMed, 33 in Scopus, 9 in CINAHL, and 865 in Cochrane Library. We considered for inclusion 8 from PubMed, 0 from Scopus, 2 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 10 articles considered for inclusion, 6 randomized trials and 4 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnet vs. Placebo</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carter 2002</td>
<td>RCT</td>
<td>Sponsored by The Oklahoma Center for Family Medicine Research. No mention of COI.</td>
<td>6.0</td>
<td>N = 30 (26 female/4 male) with wrist pain attributed to CTS. Mean age magnet 50.7±15.5 years, placebo 48.5±11.7 years.</td>
<td>Placebo magnet (N=15) vs. 1,000 gauss magnet (N=15); 45 minute treatment. Follow-up at 2 weeks.</td>
<td>Magnet mean (SD) vs. placebo mean (SD): Post-treatment pain: 3.6(3.1) vs. 2.6(2.7), NS; Pain at 2 weeks follow-up: 4.3(2.9) vs. 4.3(3.5), NS.</td>
<td>“The use of a magnet for reducing pain attributed to carpal tunnel syndrome was no more effective than use of the placebo device.”</td>
<td>Short-term study. Data suggest lack of efficacy.</td>
</tr>
<tr>
<td>Colbert 2010</td>
<td>RCT</td>
<td>Sponsored by National Institutes of Health and Oregon Clinical and Translational Research Institute. No COI.</td>
<td>8.5</td>
<td>N = 60 (45 female/15 male) with clinical evidence of carpal tunnel syndrome. Mean age: 50 years.</td>
<td>All magnets neodymium magnetized to deliver Static Magnetic Field (SMF). All devices applied at night. 15 mT (n = 20) vs. 45 mT (n = 20) vs. 0 mT aluminum disk (control) (n = 20). Outcomes measured after 6 week treatment period and 12 week no-treatment period.</td>
<td>No significant differences between groups for symptom severity or functional status at either 6 weeks (end treatment) or 12 weeks post-treatment.</td>
<td>“Participants in the active magnet groups and the control group experienced clinically relevant improvement after 6 weeks of treatment, but no significant between-group differences in outcome measures were shown.”</td>
<td>Data suggest lack of efficacy as groups (including sham) showed similar results.</td>
</tr>
<tr>
<td>Weintraub 2000</td>
<td>RCT</td>
<td>No sponsorship or COI.</td>
<td>5.0</td>
<td>N = 8 (4 females/1 male) hands from 6 patients with moderately severe carpal tunnel syndrome. Mean age: 62.5 years for females and 75 years for males.</td>
<td>Static (sub-maximal) magnetic field therapy applied 24hrs/day for 4 weeks (n = 8 hands) vs. Placebo device applied 24 hrs/day for 4 weeks (n = 8 hands). No long-term follow-up.</td>
<td>Magnet vs. Placebo – Mean neuropathic pain score improvement: 57% vs. 13% (p = 0.046).</td>
<td>“In conclusion, this novel treatment has the potential to positively influence mild cases of acropaesthesias of hands secondary to carpal tunnel syndrome and 57% of moderately advance cases.”</td>
<td>Small sample size (n=8). Pilot study</td>
</tr>
</tbody>
</table>
WRIST SPLINTING

Wrist splinting has been utilized to treat CTS. A precise mechanism of action is unclear, although it is believed to prevent hyperflexed postures, particularly while sleeping, that provoke symptoms. Placement of the wrist in functional neutral posture (approximately 15° of extension) is most typically performed; however, most studies do not specify the posture and at least one study utilized a neutral posture of 0° which actually is a modest degree of flexion. Whether those differences in postures are clinically meaningful is unknown.

**Recommendation: Nocturnal Wrist Splinting for Acute, Subacute, or Chronic CTS**

Nocturnal wrist splinting is moderately recommended for treatment of acute, subacute, or chronic CTS. (772)

**Indications** – Symptoms consistent with carpal tunnel syndrome.

**Frequency/Dose** – Wrist splints are recommended to be worn while sleeping.(387, 763-766) There is no recommendation for or against the use of splints during the daytime; however, splints theoretically increase force requirements needed to perform some jobs and have demonstrated alterations in other upper extremity postures; thus, they may have a relative contraindication to daytime use. However, one study testing nocturnal versus full-time use suggested modestly better results in electrodiagnostic parameters, but not symptoms, with full-time use. (765) There are numerous models and trials using different types of splints with all trials showing benefits and head-to-head trials, suggest there is, as yet, no identified optimal type of splint.(622, 774, 775)

**Discontinuation** – Splints should be re-adjusted if no response within 2 weeks of starting treatment, particularly to assure that the patient is wearing them properly as well as to assess fit. If there is only partial improvement and symptoms are sufficient for additional treatment, consideration of glucocorticosteroid injection and/or electrodiagnostic testing is indicated. If there is no improvement, splints should be discontinued and the accuracy of the diagnosis re-evaluated.

**Strength of Evidence** – Moderately Recommended, Evidence (B)

**Level of Confidence** – High

**Rationale for Recommendation**

Wrist splints have been shown to be effective compared to not splinting (764, 766) or to ergonomic education.(387) Splinting is also comparable to and in some measures superior to oral steroids.(647) One trial found splinting combined with NSAIDs comparable to glucocorticosteroid injection.(631) Both trials evaluating exercises and splinting used splinting for all subjects, precluding a comparison between those interventions. One trial suggested no superiority of a combination of tendon-gliding exercises combined with splinting compared with splinting alone.(610) Another trial suggested modest superiority of surgery over 18 months of follow-up; however, there may have been a slight bias in favor of surgery due to a baseline trend towards longer duration of symptoms in the splint group.(763) particularly in light of a subsequent report that those with shorter duration of symptoms had superior results with splinting. (776) Another trial compared splinting versus injection versus surgery and found few differences except for a modest trend favoring surgery over the long term. (777) A trial conducted in the Netherlands comparing splinting with surgery found little clinical difference, but concluded surgery was more cost-effective. (778) A recent report suggests splinting is more likely to be effective in those with milder symptoms of less than 1-year duration. (776) Wrist splints are not invasive, have no significant adverse effects, and are not costly. They are moderately recommended for treatment of CTS.

**Evidence for the Use of Wrist Splinting for CTS**

There is 1 high-(763) and 18 moderate-quality(387, 611, 622, 628, 631, 647, 764-766, 774, 775, 777-783) RCTs incorporated into this analysis. There are 9 low-quality RCTs and 1 prospective randomized blinded trial(614, 626, 767, 768, 784-789) in Appendix 2.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: wrist joint, wrist, wrists, splints, splint, splinting.
nocturnal splint; carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, burning, tingling, itching, numbness, hand, palm, finger, pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, and systematic review. We found and reviewed 71 articles in PubMed, 499 in Scopus, five in CINAHL, and 77 in Cochrane Library. We considered for inclusion 27 from PubMed, eight from Scopus, zero from CINAHL, zero from Cochrane Library and four from other sources. Of the 39 articles considered for inclusion, 23 randomized trials and five systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manente 2001 RCT</td>
<td>6.5</td>
<td>N = 83 (69 female/11 male) with CTS, EDS confirmed or signs, symptoms of CTS. Mean age splint group 46.10±12.94 years, control group 50.0±12.65 years.</td>
<td>Nocturnal hand brace called Manu every night for 4 weeks (N=41) vs. No treatment, observational period before starting any treatment (N=42), for 4 weeks. Assessments at 2 weeks and 4 weeks.</td>
<td>BCTQ symptomatic score (baseline/4 weeks): splint 2.75±0.7 to 1.54±0.4 at 4 weeks vs. controls 2.77±0.7 to 2.61±0.6 (p &lt;0.001). Sensory conduction velocities not different (p = 0.55). BCTQ function scores improved more in treated group from 1.89 to 1.48 vs. control from 2.02 to 2.03 (p &lt; 0.001).</td>
<td>“The study demonstrates that this hand brace is highly efficient in relieving symptoms and functional loss in CTS.”</td>
<td>Study evaluated a unique hand brace. Non-intervention controls may bias in favor of intervention.</td>
</tr>
<tr>
<td>Premoselli 2006 RCT</td>
<td>6.0</td>
<td>N = 50 (23 female/2 male) with unilateral or bilateral CTS, EDS confirmed. Mean age splint group 53.1±13.3 years, control group 46.5±13.8 years.</td>
<td>Nocturnal splint (custom molded) for a minimum of 6 hours (N=25) vs. No treatment (N=25) for 6 months. Assessments at baseline, 3 months, and 6 months.</td>
<td>Follow-up symptoms splint vs. control group (mean±SD): 3 months: 1.63±0.25 vs. 2.57±0.31 (p = 0.001); 6 months: 1.48±0.19 vs. 2.38±0.40 (p = 0.001); Sensory latency (ms): Recruitment: 2.74±0.28 vs. 2.79±0.38 (p = 0.63); 3 months: 2.59±0.39 vs. 2.85±0.336 (p = 0.02); 6 months: 2.61±0.37 vs. 2.71±0.43 (p = 0.50).</td>
<td>“Symptom relief and neurophysiological improvement after night-only splint wear therapy lasted up to the six-month follow-up visit.”</td>
<td>Dropout rate 28% over 6 month trial. Non-intervention controls may bias in favor of intervention.</td>
</tr>
<tr>
<td>Walker 2000 RCT</td>
<td>5.0</td>
<td>N = 21 (30 hands) with unilateral or bilateral CTS, EDS confirmed. Mean age 60±11.2 years.</td>
<td>Nocturnal splints (N=13) vs. Full-time splints (N=11). Follow-up for 6 weeks.</td>
<td>Symptoms severity (baseline/ follow-up): night only (2.89±0.96/3.0±0.93) vs. full-time (2.79±0.69/2.09±0.62) (NS). Functional deficits: night (2.75±1.01/2.14±0.87) vs. full time (2.27±0.73/1.93±0.77) (NS). Motor (p = 0.04) and sensory (p = 0.05) distal latencies improved more in full-time use.</td>
<td>“The study provides added evidence to support the efficacy of neutral wrist splints in CTS and suggests that physiologic improvement is best with full-time splint wear instructions.”</td>
<td>Symptoms/function data suggest no difference in efficacy. NCS data favor full-time use. High noncompliance with full-time use (27% completely compliant with daytime use) raises questions about validity of conclusions.</td>
</tr>
<tr>
<td>Werner 2005 RCT</td>
<td>4.5</td>
<td>N = 161 with signs/symptoms suggestive of CTS for &gt;1 week or &gt;3 times in last 6 months. No EDS used for inclusion but performed after entry. Mean age splint group</td>
<td>Nocturnal splints custom made that maintained wrist in neutral posture (n = 86) vs. Ergonomic education on line (n = 75); 6 week trial. Both groups given instruction on how to reduce ergonomic stressors in</td>
<td>Wrist, hand, finger discomfort in prior 30 days (baseline/follow-up): splints (7.24±2.08/ 4.43±3.71) vs. controls (6.60±2.51/5.58±3.30), p = 0.03. Splinted group had more visits to plant medical department (15.5±7.1 visits vs. 3.6±4.3 visits, p = 0.02)</td>
<td>“Benefit from a 6-weeks nocturnal splinting trial, and the benefits were still evident at the 1-year follow-up..”</td>
<td>High dropout rate (30.4%) and 50% questionnaires incomplete may sharply limit the value of the data.</td>
</tr>
<tr>
<td>Committee on Health and Safety. No COI.</td>
<td>44.74±1.02 years, ergonomic education group 43.77±1.44 years. work and home environments. Follow-up at 3, 6, and 12 months.</td>
<td>Boston Questionnaire for Assessment of Carpal Tunnel Symptom Severity (BQSS), mean±SD (pre-treatment/post-treatment): splint 2.80±0.63/ 2.38±0.77 vs. control 2.57±0.52/ 2.60±0.62 (p &lt;0.001). Boston Questionnaire for the Assessment of Carpal Tunnel Symptom Functional Status Scale (BQFSS), mean±SD (pre-treatment/post-treatment): splint 2.24±0.78/ 2.04±0.74 vs. control 2.00±0.71/ 2.08±0.70 (p = 0.015). VAS, mean±SD (pre-treatment/post-treatment): splint 5.84±2.46/ 4.26±2.67 vs. control 5.00±2.62/ 5.65±2.54 (p = 0.001). Phalen's test, mean±SD (pre-treatment/post-treatment): splint 24.43±17.41/24.59±18.89 vs. control 27.00±15.36/22.56±15.36 (p = 0.031). Grip strength, kg force, mean±SD (pre-treatment/post-treatment): splint 23.94±8.55/25.01±9.37 vs. control 22.05±8.37/23.90±8.88 (p = 0.020). Purdue Pegboard Test score, min, mean±SD (pre-treatment/post-treatment): splint 46.87±16.41/51.40±15.30 vs. control 40.81±17.27/53.72±11.29 (p = 0.021). Semmes-Weinstein Monofilaments (SWM) score, palmar side, mean±SD (pre-treatment/post-treatment): splint 100.91±90.92/89.78±78.98 vs. control 90.27±80.27/87.50±75.80 (p = 0.001).</td>
<td>&quot;A conservative treatment program including full-time splinting and formal education as key components can improve symptoms and hand function in patients with CTS.&quot;</td>
<td>Hall 2013 RCT No mention of sponsorship or COI.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Hall 2013 RCT No mention of sponsorship or COI. | 4.5 N = 62 age 18 and older with paresthesia in median nerve distribution in night or day, clumsiness, grasp weakness, sleep disturbances, not pregnant, and no medical (surgery or injections) and conservative (wearing hand splints) treatments in past 6 months. Mean age: 53.8 years. | Conservative treatment group: full-time wrist splint (neutral position with full finger and thumb motion) and education sessions (pathology of CTS, risk identification, goal setting for self-management of CTS symptoms) by an occupational therapist (2 treatment session in 1st week and between weeks 2 and 4 plus a 20 minute phone call at week 7) for 8 weeks (n = 31) vs. Control group: assessed and observed but given no intervention for 8 weeks (n = 31). Assessments at end of 8 weeks. | Conservative treatment group better than control group for symptom improvement and function. |
<table>
<thead>
<tr>
<th>Year</th>
<th>Study</th>
<th>Design</th>
<th>Duration</th>
<th>Sample Size</th>
<th>Intervention</th>
<th>Outcome Measures</th>
<th>Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>2012</td>
<td>MacDermid</td>
<td>RCT</td>
<td>9 weeks</td>
<td>N = 63</td>
<td>Experimental group: astaxanthin 4mg capsules after evening meals for 9 weeks followed by 3 week wash-out plus neutral wrist splint at night and during day when wrist in at-risk position (n = 32) vs. Control group: placebo capsules plus neutral wrist splint (n = 31). Assessments at 3 week intervals.</td>
<td>No significant differences between groups for primary outcomes, CTS Symptom Severity Scale (p=0.18) and CTS Functional Scale (p=0.40).</td>
<td>&quot;This study has not identified astaxanthin to be an effective adjunct to standard conservative management.&quot;</td>
<td>Comparable efficacy in groups. No benefit demonstrated for use of astaxanthin.</td>
</tr>
<tr>
<td>2002</td>
<td>Celiker</td>
<td>RCT</td>
<td>2 weeks &amp; 8 weeks</td>
<td>N = 23</td>
<td>Group A: (NSAID) acemetacine 120mg a day and nocturnal splint light-weight, neutral-positioned (n = 11) vs. Group B: 40mg methylprednisolone acetate injection (n = 12). Assessments at week 2 and week 8.</td>
<td>VAS pain scores (baseline/2nd week/8th week): NSAID plus splint 7.9±1.4/ 4.3±0.9/1.7±1.0 vs. injection 7.0±2.2/ 3.1±2.5/1.8±1.9 (p=0.05). Symptom severity scale results not different.</td>
<td>&quot;Both splinting combined with the use of a nonsteroidal anti-inflammatory drug and steroid injection into the carpal tunnel resulted in significant improvement in carpal tunnel syndrome.&quot;</td>
<td>No placebo control. Results suggest splinting and NSAID may be as effective as injection.</td>
</tr>
<tr>
<td>2006</td>
<td>Ucan</td>
<td>RCT</td>
<td>3 months &amp; 6 months</td>
<td>N = 67</td>
<td>Group A: Full-time splinting in neutral position with standard splint for 3 months (n = 23) vs. Group B: Single steroid injection (20mg triamcinolone acetate with 20mg lidocaine) and splinted for 3 months (n = 23) Group C: Surgery, open carpal tunnel release (n = 11). Assessments at baseline, 3 months, and 6 months after treatment.</td>
<td>Boston Questionnaire scores (baseline/3rd month/6th month): splinting 2.66±0.35/ 1.39±0.37/ 1.54±0.31 vs. splint plus steroid 2.79±0.63/1.41±0.32/1.96±0.63 vs. CTR 3.09±0.5/1.86±0.6/1.41±0.31 (p = 0.004). Palm-wrist median sensory nerve velocities: splint 27.26 ±5.3/29.6±7.16/ 29.56±4.83 vs. splint plus steroid 26.35±4.12/ 31.57±4.33/28.74±6.19 vs. CTR 23.98±4.28/32.20±4.17/ 33.15±4.1 (NS between groups). Those completely/almost satisfied 3rd/6th months splinting 69.6%/34.8% vs. splint plus</td>
<td>&quot;All treatment methods were effective, but (open) CTR was superior to conservative methods in the long term despite complications and longer recovery time.&quot;</td>
<td>Baseline differences. Appears to have targeted lower enrollment for surgery without stating such.</td>
</tr>
<tr>
<td>Study</td>
<td>RCT</td>
<td>Participants</td>
<td>Interventions</td>
<td>Outcomes</td>
<td>Conclusion</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>-----</td>
<td>--------------</td>
<td>---------------</td>
<td>----------</td>
<td>------------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mishra 2006</td>
<td>4.0</td>
<td>N = 66 with CTS, EDS confirmed for at least 1 month. Mean age splint group 42.91 years, steroid group 41.57 years.</td>
<td>Full-time splint use for 4 weeks with commercially available carpal tunnel splint (n = 20) vs. oral prednisolone 20mg a day for 2 weeks followed by 10mg a day for 2 weeks (n = 20). Follow-up at 1 and 3 months.</td>
<td>Mean±SD for splint vs. Steroid: Symptom severity score (SSS): SSS 0-1: 0.34±0.42 vs. 0.40±0.30 (p = 0.52); SSS 0-3: 0.30±0.54 vs. 0.49±0.44 (p = 0.42). Sensory distal latency (SDL): SDL 0-1: 0.16±0.63 vs. 0.13±0.71 (p = 0.86); SDL 0-3: 0.35±0.76 vs. 0.55±0.66 (p = 0.25).</td>
<td>“There was significant improvement in both groups clinically during follow-up at 1 and 3 months as well as electrophysiologically, at 3 months.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>No blinding. Suggests splinting is as effective as oral steroid, though function slightly better with splinting.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gerritsen 2002</td>
<td>8.5</td>
<td>N = 176 with CTS, EDS confirmed without previous splinting treatment or surgery. Age 18 years or older, mean age surgery group 49±11 years, splinting group 49±12 years.</td>
<td>Open surgical release (N=87) vs. splinting, custom made or prefabricated to immobilize wrist in a neutral position, at night for at least 6 weeks but could also wear it during the day (N=89) for 12 months. Assessments at 3, 6, 12, and 18 months.</td>
<td>Surgery success rates superior other than first month (1/3/6/12/18 months) surgery vs. splinting: 29 vs. 42% (p = 0.07); 80 vs. 54% (p &lt;0.001); 94 vs. 68% (p &lt;0.001); 92 vs. 72% (p = 0.002); 90 vs. 54% (p &lt;0.001); 94 vs. 68% (p &lt;0.001); 92 vs. 72% (p = 0.002); 92 vs. 72% (p = 0.002); 90 vs. 75% (p = 0.02). Nights awakening due to symptoms (1/3/6/12/18 months) surgery vs. splinting (mean±SD): 0.8±3.2 vs. 2.0±3.0 (p = 0.008); 2.6±3.5 vs. 2.2±3.1 (p = 0.49); 3.6±7.8 vs. 2.6±3.1 (p = 0.03); 3.6±2.9 vs. 2.9±3.0 (p = 0.13); 3.6±2.9 vs. 3.2±3.1 (p = 0.44). Severity of main complaint (1/3/6/12/18 months) surgery vs. splinting (mean±SD): 1.6±2.9 vs. 2.1±2.2 (p = 0.22); 5.1±3.3 vs. 3.2±2.7 (p &lt;0.001); 6.6±2.4 vs. 4.4±3.2 (p &lt;0.001); 6.4±2.7 vs. 5.1±3.1 (p = 0.005); 6.2±2.8 vs. 5.0±3.3 (p = 0.02). Paresthesia during day (1/3/6/12/18 months) surgery vs. splinting (mean±SD): 1.5±3.0 vs. 1.4±2.1 (p = 0.66); 4.8±3.2 vs. 2.2±3.2 (p &lt;0.001); 5.5±2.9 vs. 3.7±3.2 (p &lt;0.001); 5.5±2.9 vs. 4.0±3.4 (p = 0.004); 5.3±3.0 vs. 4.0±3.6 (p = 0.01). Paresthesia at night (1/3/6/12/18 months) surgery vs. splinting “Treatment with open carpal tunnel release surgery resulted in better outcomes than treatment with wrist splinting for patients with CTS.”</td>
<td>Duration of symptoms was somewhat worse in splinting group (median 52 vs. 40 weeks, NS). Both treatment arms document substantial improvement, which may reflect a good natural history.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Condition</td>
<td>Intervention</td>
<td>Outcome Measures</td>
<td>Results</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>---</td>
<td>-----------</td>
<td>--------------</td>
<td>------------------</td>
<td>---------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Korthals-de Bos 2006</td>
<td>4.0</td>
<td>N = 176 with CTS, EDS confirmed, 18 years of age or older.</td>
<td>Surgery, standard open carpal tunnel release (N=87) vs. nocturnal splinting with custom of prefabricated splint that immobilized wrist in neutral position for at least 6 weeks. Could wear splint during day if desired (n = 89). 1-year study. Assessments at baseline 3, 6, and 12 months.</td>
<td>Success rates higher at 12 months for surgery group, surgery 92% vs. splint 72% (95% CI 8.3-31). Nights awakening due to complaints not different (surgery 3.6±2.9 vs. splint 2.9±3.0), 95% CI -0.2-1.7. Severity of main complaint higher in surgery (6.4±2.7 vs. 5.1±3.1) 95% CI 0.4-2.2. Paraesthesia during the day: surgery 5.5±2.9 vs. splint 4.0±3.4 (95% CI 0.5-2.5). Paraesthesia at night: surgery 5.2±3.6 vs. splint 4.5±3.4 (95% CI -0.4-1.8). Mean aggregate costs 2,126€ surgery vs. 2,111€ splint, NS. Absenteeism comparable (50 vs. 52 days).</td>
<td>“In the Netherlands, surgery is more cost-effective compared with splinting, and recommended as the preferred method of treatment for patients with CTS.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Garfinkel 1998</td>
<td>6.0</td>
<td>N = 52 with CTS signs and symptoms (at least 2 of 5 – positive Tinel sign, positive Phalen sign, pain in median nerve distribution, sleep disturbances resulting from hand symptoms, and numbness or paresthesias in median nerve distribution) EDS confirmed. Mean age yoga group 48.9, splint group 48.7 years.</td>
<td>Standard splint with metal insert to supplement current treatment (n = 25) vs. Iyengar yoga (1-1.5 hour, 2x a week for 8 weeks focused on upper body postures, improving flexibility, correcting alignment of hands, wrists, arms, and shoulders, stretching, increasing awareness of optimal joint position during use (n = 26). Current treatment not described. Assessments at baseline and 8 weeks.</td>
<td>Grip strength (pretest/posttest) mean±SD: 161.6±70.4/187.4±68.8 vs. splint 183.9±69.5/190.5±68.2mmHg (p=0.37), Pain reduced (pre-/post-test) mean±SD: yoga 5.0±2.8/2.9±2.2 (p = 0.02) vs. splint 5.2±2.1/4.3±2.2 (p = 0.16). Median nerve sensory conduction (pretest/posttest) mean±SD: yoga 4.40±1.5ms/3.97±1.5 (p = 0.18) vs. splint 4.66±1.4/4.36±1.6ms (p = 0.28).</td>
<td>“In this preliminary study, a yoga-based regimen was more effective than wrist splinting or no treatment in relieving some symptoms and signs of carpal tunnel syndrome.”</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Splints vs. Other Treatments including Exercise and Yoga**

<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Condition</th>
<th>Intervention</th>
<th>Outcome Measures</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Korthals-de Bos 2006</td>
<td>4.0</td>
<td>N = 176 with CTS, EDS confirmed, 18 years of age or older.</td>
<td>Surgery, standard open carpal tunnel release (N=87) vs. nocturnal splinting with custom of prefabricated splint that immobilized wrist in neutral position for at least 6 weeks. Could wear splint during day if desired (n = 89). 1-year study. Assessments at baseline 3, 6, and 12 months.</td>
<td>Success rates higher at 12 months for surgery group, surgery 92% vs. splint 72% (95% CI 8.3-31). Nights awakening due to complaints not different (surgery 3.6±2.9 vs. splint 2.9±3.0), 95% CI -0.2-1.7. Severity of main complaint higher in surgery (6.4±2.7 vs. 5.1±3.1) 95% CI 0.4-2.2. Paraesthesia during the day: surgery 5.5±2.9 vs. splint 4.0±3.4 (95% CI 0.5-2.5). Paraesthesia at night: surgery 5.2±3.6 vs. splint 4.5±3.4 (95% CI -0.4-1.8). Mean aggregate costs 2,126€ surgery vs. 2,111€ splint, NS. Absenteeism comparable (50 vs. 52 days).</td>
<td>“In the Netherlands, surgery is more cost-effective compared with splinting, and recommended as the preferred method of treatment for patients with CTS.”</td>
</tr>
<tr>
<td>Garfinkel 1998</td>
<td>6.0</td>
<td>N = 52 with CTS signs and symptoms (at least 2 of 5 – positive Tinel sign, positive Phalen sign, pain in median nerve distribution, sleep disturbances resulting from hand symptoms, and numbness or paresthesias in median nerve distribution) EDS confirmed. Mean age yoga group 48.9, splint group 48.7 years.</td>
<td>Standard splint with metal insert to supplement current treatment (n = 25) vs. Iyengar yoga (1-1.5 hour, 2x a week for 8 weeks focused on upper body postures, improving flexibility, correcting alignment of hands, wrists, arms, and shoulders, stretching, increasing awareness of optimal joint position during use (n = 26). Current treatment not described. Assessments at baseline and 8 weeks.</td>
<td>Grip strength (pretest/posttest) mean±SD: 161.6±70.4/187.4±68.8 vs. splint 183.9±69.5/190.5±68.2mmHg (p=0.37), Pain reduced (pre-/post-test) mean±SD: yoga 5.0±2.8/2.9±2.2 (p = 0.02) vs. splint 5.2±2.1/4.3±2.2 (p = 0.16). Median nerve sensory conduction (pretest/posttest) mean±SD: yoga 4.40±1.5ms/3.97±1.5 (p = 0.18) vs. splint 4.66±1.4/4.36±1.6ms (p = 0.28).</td>
<td>“In this preliminary study, a yoga-based regimen was more effective than wrist splinting or no treatment in relieving some symptoms and signs of carpal tunnel syndrome.”</td>
</tr>
</tbody>
</table>
Brininger 2007
RCT
Sponsored by the School of Health and Rehabilitation Science Development Fund, School of Health and Rehabilitation Sciences, University of Pittsburg, PA. No COI.

6.0
N = 61 at least 18 years of age with a positive Tinel sign or Phalen maneuver and complaints of nocturnal numbness and tingling. Mean age 50 years.

Neutral wrist and metacarpophalangeal (MCP) splint, custom splint positioning MCP joints 0°-10° flexion, NW/MCP (n = 17) vs. neutral wrist and MCP exercise group (tendon and nerve gliding exercises 3-5x a day with 10 reps in each position held for 5 seconds), NW/MCP-X (n = 16) vs. wrist cock-up splint prefabricated that immobilized wrist in 20° of extension, WCU (n = 12) vs. wrist cock-up splint and exercise, WCU-X (n = 16). All groups wore splint during sleep for 4 weeks and received educational brochure on CTS. Assessments at baseline, 4 and 8 weeks.

All groups saw significant decrease in CTS symptoms (no p-value reported).

"Our results provide further evidence of the effectiveness of splinting, designed to target an underlying anatomic problem, for reducing symptoms and improving functional status in patients with mild-to-moderate CTS."

Small group numbers. No table or graphic for results. Baseline comparability for group strength different between groups.

Baysal 2006
RCT
No mention of sponsorship or COI.

5.5
N = 36 (72 wrists) females with bilateral CTS, EDS confirmed. Mean age Group 1 47.8±5.5 years, Group 2 50.1±7.3, Group 3, 51.4±5.2 years.

Group 1: tendon- and nerve-gliding exercises 5 daily sessions, each exercise repeated 10x each session for 3 weeks plus splinting with custom made neutral volar splint for 3 weeks all night and during day (n = 12) vs. Group 2: ultrasound 15 minutes a session to palmar carpal tunnel area, frequency 1 MHz, intensity 1.0 W/cm², 15 treatments 1x a day, 5x a week for 3 weeks plus splinting (n =12) vs. Group 3: ultrasound, splinting and exercises n = 12). Full-time splint use; 8 week treatment.

Pain score before treatment/after treatment/after 8 weeks follow-up: Group I: 4.8±2.3/3.3±2.9/ 2.6±2.8; Group II: 5.7±2.7/2.2±1.9/ 2.5±2.8; Group III: 5.6±3.5/1.3±1.8/ 0.8±0.9. Functional status score: Group I: 20.6±7.8/14.8±7.5/ 14.9±6.6; Group II: 21.9±9.1/16.1±8.5/ 16.1±8.7; Group III: 20.5±7.1/11.7±3.6/ 12.6± 3.4. NS between groups for study outcomes.

"The result of this study emphasizes the efficacy of conservative treatment in CTS. In all patient groups, the treatment combinations were significantly effective immediately and 8 weeks after the treatment."

Small group numbers. No table or graphic for results. Baseline comparability for group strength different between groups.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Intervention</th>
<th>Outcome Measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fusakul 2014</td>
<td>RCT</td>
<td>5.5</td>
<td>N = 66 (126 hands) aged 18 and older with CTS symptoms and a mild-to-moderate diagnosis made with clinical exams and electrodiagnosis. Mean age Group I – 50.70±1.39 years, Group II – 50.79±1.38 years. Group I: low-level laser therapy (LLLT), 18J per session over carpal tunnel area, 15 sessions for 5 weeks plus neutral wrist splint at night and during day for 12 weeks (n = 63) vs. Group II: placebo treatment, red light without laser power output over carpal tunnel, 15 sessions for 5 weeks plus neutral wrist splint at night and during day for 12 weeks (n = 63). Both groups encouraged to perform tending gliding exercises. Follow-up 5 and 12 weeks after treatment.</td>
<td>Symptom Severity Scale (SSS) mean±SD (baseline/week 5/week 12): Group I 2.10±0.68/1.68±0.66/1.49±0.58 vs. Group II 1.68±0.56/1.43±0.49/1.35±0.51 (p=0.031 at week 5). Distal motor latency (DML) mean ±SD (baseline/week 12): Group I 4.84±0.15/4.73±0.13 vs. Group II 5.20±0.18/6.63±1.10 (p=0.015). <strong>Both LLLT and splints improved the clinical parameters of our study, but LLLT was electroneurophysiologically superior to splints with regard to the conduction of the median motor nerve fibers.</strong></td>
</tr>
<tr>
<td>Soyupek 2012</td>
<td>RCT</td>
<td>4.5</td>
<td>N = 52 (81 wrists) with CTS, EDS confirmed. Mean age splinting, PCS, PNSAI: 47.95±6.93 years, 50.50±8.71 years, 53.79±10.40 years. Phonophoresis with corticosteroid (betamethasone valerate 0.1% cream), CS (PCS) over carpal tunnel for 10 minutes/session at frequency 3 MHz, intensity 1.5 W/cm² 5x a week for 3 weeks (n = 28) vs. phonophoresis with non-steroidal anti-inflammatory drug (diclofenac diethyl ammonium gel), NSAI (PNSAI) over carpal tunnel for 10 minutes/session frequency 3 MHz, intensity 1.5 W/cm² 5x a week for 3 weeks (n = 23) vs. wrist splinting in neutral position during</td>
<td>VAS difference from baseline to after 3 months, mean±SD (baseline/after 3 months): splinting group 50.69±23.45/37.91±23.94 (NS); PCS 60.35±18.95/30.35±18.15 (p&lt;0.017); PNSAI 69.13±16.21/45.65±23.65 (p&lt;0.017). Boston Questionnaire total difference from baseline to after 3 months, mean±SD (baseline/after 3 months): splinting group 43.34±10.89/39.26±10.03 (NS); PCS 54.21±11.34/39.14±10.33 (p&lt;0.017); PNSAI 53.69±41.86/41.86±10.03 (p&lt;0.017). Tinel’s sign, %, difference from baseline to after 3 months (baseline/after 3 months): splinting group</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Study</th>
<th>Methodology</th>
<th>Study Population</th>
<th>Intervention</th>
<th>Primary Outcomes</th>
<th>Secondary Outcomes</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kumnerddee 2010</td>
<td>RCT</td>
<td>N = 61 with mild-to-moderate CTS, EDS confirmed. Mean age 50.37±9.01 years; night splinting 51.73±8.92 years.</td>
<td>Acu group: 10 sessions of electro-acupuncture 2x a week on meridian of affected area (n = 30) vs. NS group: prefabricated volar neutral wrist splint worn at night for 5 weeks (n = 31). Assessments at baseline and end of treatment.</td>
<td>Mean±SD VAS (baseline/end of treatment): acupuncture 22.57±22.27/7.97±14.99 vs. night splinting 22.59±26.09/17.60±22.37 (p = 0.028). NS between groups for Symptom Severity Scale (p = 0.295) and Functional Status Scale (p=0.663).</td>
<td>“Electro-acupuncture provides more pain attenuating effect than night splinting in mild-to-moderate degree CTS.” Comparable efficacy, but pain symptoms relieved slightly better with acupuncture group. Study susceptible to significant contact time bias.</td>
<td></td>
</tr>
<tr>
<td>Storey 2013</td>
<td>RCT</td>
<td>N = 49 diagnosed with CTS from history and clinical exam confirmed with nerve conduction studies. Mean age C-Trac splint 47 years, BWB 39 years.</td>
<td>C-Trac splint (C-shaped, tubular, semirigid frame contoured around dorsum of wrist and hand with air pressure bladder to control pressure to 180-190mmHg for 2 minutes) 3x a week first 4 weeks then as necessary (n = 25) vs. Beta Wrist Brace (BWB) resting splint at night and during activities that provoke symptoms first 4 weeks then as necessary (n = 24). Follow-up at 4, 8, 26, and 52 weeks.</td>
<td>No significant differences between groups for primary outcomes, Levine symptom (p = 0.213) and function (p = 0.308) scores by week 8. No significant differences between groups for secondary outcomes at 8 weeks, Semmes-Weinstein monofilament scores (p = 0.0567), grip strength (p = 0.568), lateral pinch (p = 0.728), tripod pinch (p = 0.183).</td>
<td>“These results suggest that C-Trac splint is not dissimilar in efficacy to a resting Beta Wrist Brace.” Pilot study showing similar efficacy between C-Trac splints compared to Beta wrist braces at 8 weeks, 6 months and 12 months.</td>
<td></td>
</tr>
<tr>
<td>De Angelis 2009</td>
<td>RCT</td>
<td>N = 120 age 18 or older with possible CTS, pain, numbness, and paresthesias</td>
<td>Hand brace MANU® that does not impede thumb-index finger pinch, thumb-little finger opposition, and wrist</td>
<td>No significant differences between groups for the primary study outcomes (p = 0.097-0.821).</td>
<td>“Our findings demonstrate that a conservative treatment by the hand brace or a splint is effective as High dropout rate. At 3 months, comparable efficacy</td>
<td></td>
</tr>
<tr>
<td>Sponsored by the AGF Orthopaedic Devices s.r.l. company. No COI.</td>
<td>and/or hypoesthesia in the median nerve distribution, positive Phalen test, exclusive or predominant in one, and electrophysiologic al diagnosis of CTS. Mean age MANU® 46.0±11.8 years, CAMP TIHELLE® 46.3±7.9 years.</td>
<td>flexion and extension worn every night for 3 months (n = 59) vs. wrist splint CAMP TIHELLE® that immobilizes wrist in dorsiflexion position with external angle of 30° and internal angle of 16° worn every night for 6 months (n = 61). Follow-up at 3 months and 6 months after treatment.</td>
<td>long as they are employed as already shown in other studies.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Allied Health Therapies
ACUPUNCTURE
Acupuncture has been used to treat CTS and other hand, wrist, and forearm MSDs. (790, 791) There is evidence of its efficacy for treatment of chronic spine disorders, although the evidence suggests traditional acupuncture is not superior to other acupuncture methods (see Chronic Pain and Low Back Disorders Guidelines).

Recommendation: Acupuncture for Acute, Subacute, or Chronic CTS
Acupuncture is not recommended for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – Not Recommended, Evidence (C)
Level of Confidence – Low

Rationale for Recommendation
There are quality trials of acupuncture compared with placebo or sham acupuncture and they have failed to show benefit of acupuncture for treatment of CTS. (792) One trial found no differences between acupuncture and oral steroid. (793, 794) Another trial susceptible to contact time bias found minimal differences between acupuncture and nocturnal wrist splinting. (781) Thus, the highest quality evidence suggests acupuncture is ineffective for treatment of CTS and acupuncture is not recommended.

Evidence for the Use of Acupuncture
There are 4 moderate-quality RCTs incorporated into this analysis. (781, 792-794) There are 3 low-quality RCTs in Appendix 2. (795-797)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Acupuncture, Acupuncture Therapy, carpal tunnel syndrome, CTS, median nerve neuropathy, median nerve disease, entrapment, neuropathy, nerve compression, burning, itching, numbness, tingling, wrist, hand, palm, finger, pain, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random," randomized, randomization, randomly; systematic, systematic review, retrospective studies, and prospective studies. We found and reviewed 40 articles in PubMed, 411 in Scopus, 83 in CINAHL, 46 in Cochrane Library and 0 in other sources. We considered for inclusion 7 from PubMed, 2 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 9 articles considered for inclusion, 8 randomized trials and 2 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yao 2012</td>
<td>RCT</td>
<td>Sponsored by the Department of Physical Medicine and Rehabilitation, university of California and by the National Institute of Disability Research grant. No COI.</td>
<td>7.0</td>
<td>N = 41 (gender not specified); acupuncture-naïve adult patients with mild to moderate CTS. Found through electrodiagnostic testing; mean age: Group 1 48.5±10.5; Group 2 – 53.6±7.65.</td>
<td>Acupuncture group given treatment during 6 weekly sessions for 20 minutes. Group asked to feel a de-qi sensation; heaviness (n = 21) vs. Placebo acupuncture group acupuncturists stopped manipulate needle for 2 seconds. Both groups given wrist splints for sleeping (n = 20). Follow-up baseline, immediately after 6 weeks treatment, 2 weeks and 3 months after last treatment.</td>
<td>Comparing baseline to three months after the last treatment carpal Tunnel Self-Assessment Questionnaire (CTSAQ) scores improved in both groups. Group 1, 0.58 improvement (p = 0.03), Group 2 improved by 0.81 (p = 0.001). Analyzing CTSAQ hand function 3 months after last treatment group 1, improvement by 0.45 (p = 0.17) and group 2, improvement by 0.48 (p = 0.02) both improved significantly.</td>
<td>“Both treatment and placebo groups demonstrated improvements from baseline.”</td>
<td>Splints given to all participants. Small sample size with 20% dropout in 1 arm. Acupuncture not superior to placebo acupuncture.</td>
</tr>
<tr>
<td>Yang 2009</td>
<td>RCT</td>
<td>Sponsored by Kuang Tien General Hospital grant. No COI.</td>
<td>5.5</td>
<td>N = 77 (63 females/14 males) consecutive and prospective patients with mild to moderate CTS and naive to acupuncture treatment (confirmed by NCS); mean age: Group 1 – 9.3±8.9; Group 2 – 49.9±10.3.</td>
<td>Acupuncture 8 sessions of 30 minutes duration for 4 weeks (2x a week) (n = 38) vs. Steroid treatment group: 20mg daily of prednisolone for 2 weeks and 10mg daily for following 2 weeks. 4 weeks total (n = 39). Follow-up baseline, 2 weeks, 4 weeks for Global Symptom Score and nerve conduction study (NCS) scores at baseline and 4 weeks.</td>
<td>At study end, there was a high percentage of improvement in both acupuncture and steroid groups at 2 weeks and 4 weeks (p &lt; 0.01). Although there was no statistical significance between the two group at these follow ups. Nocturnal awakening week 4, acu group 3.5 ± 3.8 vs steroid group 1.5 ± 1.9, (p &lt; 0.03).</td>
<td>“Despite the limitations, this randomized, controlled study indicates that short-term acupuncture treatment is as effective as short-term low-dose steroid for mild-to-moderate CTS.”</td>
<td>Minimal differences between groups observed. Population poorly described.</td>
</tr>
<tr>
<td>Yang 2011</td>
<td>RCT</td>
<td>Sponsored by Kuang Tien General Hospital grant. No COI.</td>
<td>5.0</td>
<td>N = 70 whom had not done any other type of intervention since the other study. (Yang 2009); Mean age: Group 1 – 49.3±8.9;</td>
<td>Acupuncture consisted of 8 sessions of 30 minute duration administered for 4 weeks (twice a week) (n = 38) vs. Steroid treatment group prescribed 20mg daily of prednisolone for 2</td>
<td>Global Symptom Score (GSS) month 7, group 1 3.4±5.8 vs group 2 7.2±5.4 (p &lt;0.01). GSS at month 13 group 1, 4.5±7.7 vs group 2, 11±8.6 (p &lt;0.01). Month 13 – Baseline improvement in GSS group 1, -11.53±7.63 vs group 2, 3.28±10.64 (p</td>
<td>“[T]herefore, we conclude that acupuncture treatment can be considered as an alternative therapy to other conservative treatments for those who do not opt for early</td>
<td>Long term follow up of prior study. No statistical difference between groups at any time point.</td>
</tr>
<tr>
<td>Kumnerddee 2010 RCT</td>
<td>4.0</td>
<td>N = 61 with mild to moderate CTS who have not participated in surgical treatment, steroid injections, or were pregnant, all patients asked to discontinue use of NSAIDs during study; age: Group 1: 50.37±9.01; Group 2: 51.73±8.92</td>
<td>Acupuncture group, 10 sessions 2x a week, needles placed around median nerve and received 1 Hz current for 30 minutes (n =30) vs. Night Splinting group for 5 weeks, use of metal bar splint to restrict wrist flexion during sleep (n = 30). Follow-up at baseline and immediately after treatment period (5 weeks).</td>
<td>Boston Carpal Tunnel Outcome Scale (BCTS) decreased significantly, 1.92± 0.54 (baseline) to 1.53±0.34 (treatment end) Acu group (p &lt;0.001) vs. 1.88±0.48 (baseline) to 1.61±0.43 (end) (p &lt;0.007) splint group. Acu group Symptom Severity Scale (SSS) 2.03±0.61 (baseline) to 1.57±0.39 (end), Functional Status Scale (FSS) 1.76±0.63 (baseline) to 1.50±0.39 (end) and VAS 22.57±22.67 (baseline) to 7.97±14.99 (end) scores all decreased significantly (p &lt;0.05) vs. night splinting for “Electro-acupuncture provides more pain attenuating effect than night splinting in mild-to-moderate degree CTS.”</td>
<td>Comparable efficacy, but pain symptoms relieved slightly better with acupuncture group. Study susceptible to significant contact time bias.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>-------</td>
<td>-------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>which only SSS decreased significantly (p = 0.008) at 5 weeks. Comparing groups: VAS reduction Acu group 14.60±19.31 vs 4.97±24.37 NS group (p = 0.028).</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
BIOFEEDBACK
Biofeedback is a behavioral medicine method of providing automated information and training to improve control of certain physiologic processes which are normally inaccessible to a subject’s perception. Audible electromyographic (EMG) biofeedback has been used to treat CTS.(798)

Recommendation: Biofeedback for Acute, Subacute, or Chronic CTS
There is no recommendation for or against the use of biofeedback for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There are no quality studies regarding the utilization of biofeedback for treating CTS patients. Biofeedback is not invasive, has no adverse effects, and is moderate cost. However, in the absence of quality evidence, there is no recommendation for or against its use.

Evidence for the Use of Biofeedback
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Biofeedback or psychology; carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, burning, itching, numbness, tingling, hand, palm, finger, wrist, pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 14 articles in PubMed, 92 in Scopus, 3 in CINAHL, and 1 in Cochrane Library. We considered for inclusion 3 from PubMed, 0 from Scopus, CINAHL, Cochrane Library or other sources. Of the 3 articles considered for inclusion, 0 randomized trials and 0 systematic studies met the inclusion criteria.

LOW LEVEL LASER THERAPY
Low level laser treatment (LLLT) has been used to treat MSDs including CTS.(790, 799, 800) It usually involves laser energy that does not induce significant heating (the theory is that the mechanism of action is through photoactivation of the oxidative chain).(801)

Recommendation: Low Level Laser Therapy for Acute, Subacute, or Chronic CTS
Low level laser therapy is not recommended for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – Not Recommended, Evidence (C)
Level of Confidence – Low

Rationale for Recommendation
There are multiple moderate-quality studies evaluating LLLT with all of the higher quality studies demonstrating lack of efficacy. There are 5 trials comparing LLLT with sham/placebo laser and the 3 highest quality studies found lack of benefit.(802-804) One trial found no differences when compared with ultrasound(805) and a second trial found ultrasound superior.(806) Another study found no additive benefits of LLLT over splinting.(807) Thus, higher quality evidence indicates LLLT is not effective for treatment of CTS. Low-level laser is not invasive, has low adverse effects, but is costly. It is not recommended for the treatment of CTS.

Evidence for the Use of Low-Level Laser Therapy for CTS
There are 11 moderate-quality RCTs and 1 moderate-quality crossover trial incorporated into this analysis.(779, 799, 802-811) There is 1 low-quality RCT in Appendix 2.(812)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: laser or low-level laser therapy, carpal tunnel, medial...
We found and reviewed 41 articles in PubMed, 541 in Scopus, 29 in CINAHL, 38 in Cochrane Library and. We considered for inclusion 9 from PubMed, 5 from Scopus, 0 from CINAHL, and Cochrane Library. Of the 14 articles considered for inclusion, 13 randomized trials and 0 systematic review met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Irvine 2004</td>
<td>RCT</td>
<td>7.5</td>
<td>N = 15 (12 female/3 male) with CTS. Ranging in age from 34 to 67 years, (46 ± 11).</td>
<td>Gallium/aluminum/arsenide laser treatment (n = 8) vs. Control group or treatment with a sham laser (n = 7). Follow-up for 4 weeks.</td>
<td>Improvement in sham laser (p = 0.034) and LLLT treatment groups, (p = 0.043). NS between group differences, (p = 0.69).</td>
<td>&quot;[L]LLT is no more effective in the reduction of symptoms of CTS than is sham treatment.&quot;</td>
<td>No difference between groups.</td>
</tr>
<tr>
<td>Tascioglu 2012</td>
<td>RCT</td>
<td>7.5</td>
<td>N = 60 (46 female/14 male) with CTS symptoms shorter than 6 months. Aged between 28 and 68 years.</td>
<td>First group received Ga-Al-As laser irradiation at each point, once daily, 5 days a week (N = 20) vs. Second group treated with same low-power laser, but painful points irradiated with duration of 1 minute, once daily, 5 days a week (n = 20) vs. Third group received placebo laser, with duration of 2 minutes irradiation, 1x daily, 5 days a week (n = 20). Follow-up for 15 days.</td>
<td>Pain scores decreased significantly in all groups at Study end for group I, II and III, (p &lt; 0.001, p &lt;0.001, and p &lt; 0.01). FSS scores improved in all groups, (p &lt;0.05).</td>
<td>&quot;In conclusion, the results of this study indicate that low level laser, given at two different dosages, was no more effective than placebo in the treatment of CTS.&quot;</td>
<td>Comparable results showing LLL not superior to placebo.</td>
</tr>
<tr>
<td>Bakhtiary 2004</td>
<td>RCT</td>
<td>7.0</td>
<td>N = 40 and 10 (gender not specified) with bilateral and unilateral CTS confirmed by electromyography or 90 wrists. Age means for laser/ultrasound groups: 48</td>
<td>Ultrasound,15 minute sessions with frequency of 1 MHz and intensity of 1.0W/cm², pulsed mode duty cycle of 1:4 and transducer area 5cm² (N = 45) vs. Low-level laser therapy, applied low intensity 9J, infrared laser diode, 830nm at 5 points, 1.8J/point, daily 15</td>
<td>Thumb sensory latencies favored ultrasound: -0.7 vs-0.2, (p = 0.003). Other electrodiagnostic measures all favored ultrasound. VAS pain scores -6.3 in the ultrasound group vs. -2.0 in laser group, (p &lt; 0.001) at 4 weeks after completion of treatment.</td>
<td>&quot;[U]ltrasound treatment is more effective than low level laser therapy in patients with mild to moderate carpal tunnel syndrome.&quot;</td>
<td>Suggests laser not effective compared with ultrasound.</td>
</tr>
<tr>
<td>Year</td>
<td>Study</td>
<td>Design</td>
<td>Duration</td>
<td>Participants</td>
<td>Intervention</td>
<td>Outcome Measures</td>
<td>Findings</td>
</tr>
<tr>
<td>-------</td>
<td>-------</td>
<td>--------</td>
<td>----------</td>
<td>--------------</td>
<td>--------------</td>
<td>-----------------</td>
<td>----------</td>
</tr>
<tr>
<td>Naeser 2002</td>
<td>RCT</td>
<td>Double-blind</td>
<td>Crossover</td>
<td>N = 11 (2 female/9 male) with mild to moderate CTS, EDS confirmed. Age range from 40 to 68 years (mean 53.5 y).</td>
<td>Device 1: Red-beam laser, continuous 15-mW, applied to shallow acupuncture points located on the fingers and hand, 3 times weekly (n = 11) vs. Device 2: Infrared pulsed laser, 180ns, 9.4W, located on the elbow, shoulder, upper back, and cervical paraspinal areas, 3 times weekly (n = 11) Device 3: Microamps TENS 580µA-3.5mA device, applied to the affected wrist, 3 times weekly (n = 11). Follow-up for 3 to 4 weeks.</td>
<td>McGill Pain Questionnaire scores were significantly lower with real treatment, (p = 0.0035). Sensory latencies were improved with real treatment, (p = 0.009), but not motor latencies, (p = 0.27).</td>
<td>&quot;[LLLT] appears to be an affective substitute for surgery…especially when this new conservative treatment is applied in the early stages of CTS (preferably within 1y of symptom onset) and with middle to moderate cases (as defined with NCSs and where there is no abnormality on needle electromyography).&quot;</td>
</tr>
<tr>
<td>Ercik 2007</td>
<td>RCT</td>
<td>Placebo-controlled</td>
<td>Double-blind</td>
<td>N = 81 (70 female/11 male) with CTS diagnosis, on both clinical examination and electromyographic (EMG) study. Age range, 26-78.</td>
<td>Group 1 or laser group received 7 joules/per point over carpal tunnel area at wrist (n = 41) vs. Group 2 or placebo laser therapy group (n = 40). Follow-up at 4 and 12 weeks.</td>
<td>VAS scores for day and night showed significant decrease in both groups at end of therapy, (p &lt; 0.001). Statistically significant improvement in sensory nerve velocity, and sensory and motor distal latencies in laser group, (p &lt;0.001), and sensory nerve velocity meaningful in placebo group, (p &lt;0.05).</td>
<td>&quot;In using LLLT, (1) there was no difference relative to pain relief and functional capacity during the follow-up in CTS patients; (2) there were positive effects on hand and pinch grip strengths.&quot;</td>
</tr>
<tr>
<td>Yagci 2009</td>
<td>RCT</td>
<td></td>
<td></td>
<td>N = 45 (hands) with symptoms and signs of suspected</td>
<td>Splinting or S group splinted in neutral position with standard cotton-polyester splints</td>
<td>No differences at baseline and third month, (p &gt;0.05). Symptom severity score of SLLLT group statistically</td>
<td>&quot;As a conclusion, both SLLLT and splinting provided improvements in clinical parameters but</td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Randomization</td>
<td>Funding</td>
<td>Setting</td>
<td>Sample Size</td>
<td>Intervention</td>
<td>Comparator</td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>---------------</td>
<td>---------</td>
<td>---------</td>
<td>-------------</td>
<td>-------------</td>
<td>------------</td>
</tr>
<tr>
<td>Chang 2008</td>
<td>RCT</td>
<td>Placebo-controlled</td>
<td>Double-blind</td>
<td>Sponsored by the National Science Council of the Republic of China</td>
<td>5.5</td>
<td>N = 36 with mild to moderate degree of CTS. Age mean for laser/ and placebo groups; 46.01 ± 11.65 / 49.07 ± 11.28.</td>
<td>Laser group received laser treatment (10 Hz, 50% duty cycle, 60 mW, once daily for two weeks (N = 20 wrists)) vs. Placebo group received sham laser treatment (N = 20 wrists). Follow-up after 2 weeks of treatment for 18 week.</td>
</tr>
<tr>
<td>Saeed 2012</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td></td>
<td></td>
<td>5.5</td>
<td>N = 100 with unilateral CTS diagnosed clinically and electrophysiologically. The mean age was 35.59 ± 6.1.</td>
<td>Group A, treated by Ultrasound therapy 1MHz, 1.0 Watt/cm2, 5x a week for 4 weeks (n = 50) vs. Group B, treated with LLLT or 830 nm infrared, 5x a week for 4 weeks (n = 50). Follow-up for 4 weeks.</td>
</tr>
<tr>
<td>Fusakul 2014</td>
<td>RCT</td>
<td>Double-blind</td>
<td>Sponsored by grant from Research Support Funding of the Faculty of Medicine at Vajira Hospital, Navamindradhiriwaj University, Thailand</td>
<td>No COI</td>
<td>5.5</td>
<td>N = 66 with mild to moderate carpal tunnel syndrome (CTS). Mean age for group I / II: 50.70 ± 1.39 / 50.79 ± 1.38.</td>
<td>Group I, LLLT with a splint of 15 sessions, 3 times weekly for 5 weeks (n = 63 hands) vs. Group II, placebo treatment with splint for 15 sessions, 3x a week for 5 weeks (n = 63 hands). Follow-up for 5 weeks.</td>
</tr>
</tbody>
</table>

Copyright© 2016 Reed Group, Ltd.
<table>
<thead>
<tr>
<th>Studied</th>
<th>Year</th>
<th>Design</th>
<th>Sponsorship/COI</th>
<th>N</th>
<th>Selection Criteria</th>
<th>Intervention</th>
<th>Main Findings</th>
<th>Study Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shooshtari 2008</td>
<td>4.0</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>80</td>
<td>CTS based on clinical examination and electromyographic (EMG) findings. Age range 30-70.</td>
<td>Group A received low power laser waves by physiolaser Olympic with multiple probe five times weekly (n = 40) vs. Group B received flash laser (n = 40).</td>
<td>Median transcarpal sensory NCV after/before treatment, (p &lt;0.001). Hand grip power increased 15.39% Group A with no meaningful improvements in Group B. NCV of median nerve in Group A improved about 3.25% ms, 1.99% ms, 6.43 m/s, with no meaningful changes in Group B.</td>
<td>“Laser therapy as a new conservative treatment is effective in treating CTS paresthesia and numbness and improved the subjects' power of hand grip and electrophysiological parameters.”</td>
</tr>
<tr>
<td>Raeissadat 2010</td>
<td>4.0</td>
<td>RCT</td>
<td>Single-blind</td>
<td>65 (hands) with mild or moderate CTS. The mean age of patients was 43.9 years.</td>
<td>Group I received local corticosteroid injection or Hydrocortisone 50mg (n = unknown) vs. Group II, received low level laser therapy or 20J/cm² in 11 seconds/session for each of 5 points, 775nm, 10 sessions and 3sessions / week (n = unknown). Follow-up for 10 months.</td>
<td>Severity of disease in injection group based on electrodiagnostic findings; mild in 41.2%, moderate in others. After 10 months, electrodiagnostic studies normal in 32.4% (38.7% before treatment), mild in 23.5% (22.6%), moderate in 41.2% (35.5%), severe in 2.9% (3.2%). Median nerve distal sensory latency before (DSL1) and 10 months after accomplishing treatment and comparison of 2 groups: injection therapy vs laser therapy: 4.28±0.36 vs 4.25±0.43 DSL1, and 3.9±0.5 vs 4±0.6, DSL2, (p &gt;0.05). Distal motor latency: 4.3±0.6 vs 4.33±0.65 (MDL1) and 4.2±0.7 vs 4.17±0.8 (DML2), (p &lt;0.05). Before vs. 10 months after treatment severity of disease: mild 45.2% vs 22.6%.</td>
<td>“Low level laser therapy can be as effective as local injection in reducing pain and severity of disease (based on electrodiagnostic medicine classification) in patients with mild and moderate CTS even in long term (after 10 months).”</td>
<td></td>
</tr>
</tbody>
</table>

Comparable efficacy. Patient blinding not possible due to different treatments (injection vs. laser).
MANIPULATION AND MOBILIZATION

Manipulation and mobilization are two types of manual therapy which have been used for treatment of CTS. These include wide arrays of different techniques and schools of thought. Some consider these two interventions to be on a spectrum of velocity and applied force. In general, mobilization involves assisted, low-force, low-velocity movement. Manipulation involves high-force, high-velocity, and low-amplitude action with a focus on moving a target joint (see Chronic Pain and Low Back Disorders Guidelines for more details).

1. Recommendation: Manipulation of the Wrist Acute, Subacute, or Chronic CTS

There is no recommendation for or against the use of manipulation of the wrist for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

2. Recommendation: Manipulation of the Spine for Acute, Subacute, or Chronic CTS

Manipulation of the spine is not recommended for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – Not Recommended, Insufficient Evidence (I)
Level of Confidence – High

Rationale for Recommendations

There are two moderate-quality studies that evaluate manipulation for treatment of CTS. However, both have considerable methodological problems. One study compared manipulation plus ultrasound versus ibuprofen. Exclusion criteria did not exclude prior ibuprofen use, which is may well have been widespread, resulting in a comparison analogous to no treatment, which biases towards the other treatment arm, ibuprofen use was PRN after 2 weeks, subject contact time differed between groups, all biasing in favor of manipulation plus ultrasound. That study failed to find improvements compared with ibuprofen(637) which as noted previously appear ineffective. The other moderate-quality study had two active-treatment arms.(819) Thus, there is no quality study showing manipulation is effective as a treatment for CTS. Manipulation is not invasive, is moderately costly, but does have rare adverse effects from cervical manipulation. Cervical manipulation is not recommended for treatment of CTS. There is no recommendation for or against manipulation of the wrist as there is an absence of quality evidence.

Evidence for the Use of Manipulation and Mobilization for CTS

There are 2 moderate-quality RCTs incorporated into this analysis.(637, 819) There are 3 low-quality RCTs in Appendix 2.(625, 820, 821)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: manipulation or mobilization / carpal tunnel, median nerve, median, carpal, disease, entrapment, neuropathy, syndrome, compression, CTS, burning, itching, numbness, tingling, hand, palm, finger, wrist, and pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 38 articles in PubMed, 172 in Scopus, 26 in CINAHL, and 10 in Cochrane Library. We considered for inclusion 3 from PubMed, 8 from Scopus, 3 from CINAHL, 1 from Cochrane Library and 0 from other sources. Of the 15 articles considered for inclusion, 3 randomized trials and 8 systematic studies met the inclusion criteria.
1. Recommendation: Massage for Acute, Subacute, or Chronic CTS

Massage is not recommended for most patients for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – Not Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

2. Recommendation: Massage for Acute, Subacute, or Chronic CTS with Forearm Myofascial Pain

Massage is recommended for treatment of select patients with acute, subacute, or chronic CTS who have significant myofascial pain.

Indications – Symptoms of carpal tunnel syndrome combined with forearm myofascial pain sufficient for the patient to require treatment. Generally, the patient should have failed other treatments including splints and glucocorticosteroid injection.

Frequency/Dose – Three to 4 appointments. Objective evidence of improvement should be followed. Additional 3 or 4 treatments should be based on improvement in objective measures.
Discontinuation – Resolution, failure to objectively improve, or intolerance.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendations
There is no quality evidence of efficacy for massage as a treatment for CTS. There is one moderate-quality trial that suggested Madenci hand massage (author same as the named massage technique) was effective as a combined therapy, however, the study design includes significant contact time biases and multiple unquantified co-interventions.(822) Regardless, massage is not thought to be helpful for typical CTS patients. However, some patients with forearm myofascial pain are thought to potentially derive some benefits. Objective measures should be followed documenting improvement in order for additional treatments to be added. Massage is not invasive, has few adverse effects, but is moderately costly over time.

Evidence for the Use of Massage
There is 1 moderate-quality RCT incorporated into this analysis.(822) There are 2 low-quality RCTs in Appendix 2.(823, 824)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Massage, soft tissue massage and carpal tunnel syndrome, CTS, median nerve neuropathy, median neuropathy, median nerve disease entrapment, neuropathy nerve compression, burning, itching, numbness, tingling, and pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 22 articles in PubMed, 209 in Scopus, 13 in CINAHL, 128 in Cochrane Library and 0 in other sources. We considered for inclusion 3 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 3 articles considered for inclusion, 3 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Madenci 2012 RCT</td>
<td>4.5</td>
<td>N = 80 (76 females/4 males) with CTS with symptoms for longer than 6 weeks and at least 1 positive test of following: Tinel, Phalen, Buda, and Carpal compressio</td>
<td>Group I, splint plus massage; Madenci hand massage technique (MHMT) self-applied for 6 weeks with weekly follow-up visits (n = 40) vs. Group II, splint (n = 40). Both groups received tendon and nerve gliding exercises and analgesic drugs. All wore wrist-hand resting splint during sleep at night for 6 months.</td>
<td>Patient global assessment (PGA, pre-treatment/post-treatment, mean±SD): Group I (8.5±1.2/2.3±0.8) v. Group II (8.2±1.2/4.1±0.7), p = 0.001. Physician global assessment (MDPGA, pre-treatment/post-treatment, mean±SD): Group I (5.9±0.8/1.2±0.5) v. Group II (5.1±0.9/2.7±0.8), p = 0.002. Grip strength right: Group I (25.4±6.3/30.3±5.2) vs. Group II (25.7±5.9/28.2±3.2), p = 0.042. Grip strength left: Group I (21.2±3.2/26.9±2.6) vs. Group II (20.5±3.3/24.1±2.3), p = 0.041. Boston symptom severity scale: Group I (3.9±1.1/1.8±0.4) v. Group II (3.7±1.0/2.5±0.5), p = 0.001. Boston functional capacity scale: Group I (3.2±0.8/2.0±0.4) v. Group II (3.2±0.6/2.6±0.6), p = 0.001. “Statistically more significant improvement was observed in PGA, MDPGA, hand grip strength scores, and electrophysiolo gical parameters in the group applied MHMT as compared to the group applied splint therapy only.”</td>
<td>Data suggest “splint+massage” treatment superior to splint alone for global score outcome but not for any other outcomes including objective electrophysiologic measures. Study susceptible to significant contact time bias. Both groups also provided exercises and analgesics.</td>
<td></td>
</tr>
</tbody>
</table>
THERAPEUTIC TOUCH
Therapeutic touch, considered an alternative healing technique, involves the use of the practitioner’s hands to focus and facilitate healing. (825)

Recommendation: Therapeutic Touch for Acute, Subacute, or Chronic CTS
Therapeutic touch is not recommended for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – Not Recommended, Evidence (C)
Level of Confidence – Low

Rationale for Recommendation
There are no quality studies suggesting therapeutic touch is effective for treatment of CTS. (825) Therapeutic touch is not invasive, has no adverse effects, and is low cost. However, it has not been shown to be efficacious and other treatments have documented benefit, thus therapeutic touch is not recommended for the treatment of CTS.

Evidence for the Use of Therapeutic Touch for CTS
There are no quality studies. There is 1 low-quality RCT in Appendix 2. (825)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Therapeutic touch and carpal tunnel syndrome, CTS, median nerve neuropathy, median neuropathy, median nerve disease entrapment, neuropathy nerve compression, burning, itching, numbness, tingling, and pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, random; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 22 articles in PubMed, 209 in Scopus, 13 in CINAHL, 128 in Cochrane Library and 0 in other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 1 article considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

ICE/SELF-APPLIED ICE
Ice has been rarely used to treat CTS.

Recommendation: Ice for Treatment of Acute, Subacute, or Chronic CTS
There is no recommendation for or against use of ice for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There are no quality studies suggesting ice is effective for treatment of CTS. Ice is not invasive, has no adverse effects, and is low cost when self-applied. However, it has not been shown to be efficacious and other treatments have documented benefit, thus it is suggested other treatments should be used in preference.

Evidence for the Use of Ice
There are no quality studies incorporated into this analysis. A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: ice; self-applied ice, cold therapy, carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, burning, itching, numbness, tingling, hand, palm, finger, wrist, pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, random; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and
HEAT/SELF-APPLIED HEAT
Various forms of heat treatment have sometimes been used to treat CTS. (626)

Recommendation: Heat for Treatment of Acute, Subacute, or Chronic CTS
There is no recommendation for or against use of heat for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There are no quality studies suggesting heat is effective for treatment of CTS. There is one trial with paraffin as a cointervention. (626) Heat is not invasive, has no adverse effects, and is low cost when self-applied. However, it has not been shown to be efficacious and other treatments have documented benefit, thus it is suggested other treatments should be used in preference.

Evidence for the Use of Heat
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, and Cochrane Library without date limits using the following terms: Self applied heat, heat therapy, electrical induced heat, dielectric heating, self-applied heat therapy, carpal tunnel syndrome, CTS, median nerve neuropathy, median neuropathy, median nerve disease, entrapment, neuropathy, nerve compression, burning, itching, numbness, tingling, and pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 44 articles in PubMed, 34 in Scopus, 2 in CINAHL, and 38 in Cochrane Library. We considered for inclusion 1 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 2 articles considered for inclusion, 0 randomized trials and 0 systematic studies met the inclusion criteria.

DIATHERMY
Diathermy is a type of heat treatment that has been used clinically to heat tissue. (826, 827) There are two forms of diathermy – short wave and microwave. High-dose diathermy is also used to coagulate tissue. Proponents of diathermy utilize it to treat a wide range of conditions, believing it penetrates deeper than hot packs or heating pads and stimulates healing. (827, 828)

Recommendation: Diathermy for Treatment of Acute, Subacute, or Chronic CTS
There is no recommendation for or against use of diathermy for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There are no quality studies clearly demonstrating that diathermy is effective for treatment of CTS. The two available trials have considerable methodological flaws (e.g., represented as double blinded). Diathermy is not invasive, has no adverse effects, but becomes moderately costly with repeated applications. It has not been clearly shown to be efficacious and other treatments have documented benefit, thus it is suggested other treatments should be used in preference.

Evidence for the Use of Diathermy
There are 2 moderate-quality RCTs incorporated into this analysis. (829, 830)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, and Cochrane Library without date limits using the following terms: diathermy; carpal tunnel syndrome, median.
neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 33 articles in PubMed, 153 in Scopus, 0 in CINAHL, and 3 in Cochrane Library. We considered for inclusion 0 from PubMed, 2 from Scopus, 0 from CINAHL, 1 from Cochrane Library and 0 from other sources. Of the 3 articles considered for inclusion, 2 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frasca 2011</td>
<td>RCT</td>
<td>Double-blind, No sponsorship or COI</td>
<td>5.0</td>
<td>N = 22 (19 females/3 males) with idiopathic unilateral or bilateral, mild to moderate carpal tunnel syndrome (CTS). Mean age HT group 50.8±13.8 and for SC group 56.4±13.8</td>
<td>Hyperthermia treatment or HTG for 8 sessions, 20 minutes each (n = 11) vs. sham-controlled groups or SCG for 8 sessions, 20 minutes each (n = 11). Follow-up at baseline and 3 weeks.</td>
<td>At final visit of HTG improvement in pain severity vs. baseline (VAS: p = 0.002, Levine-Boston I p &lt;0.0001) and functional impairment (Levine-Boston II p = 0.002) No significant difference in SCG vs. baseline value (VAS p = 0.713 Levine-Boston I p = 0.14). Comparisons of changes in outcome measures for HTG pain severity (VAS p = 0.004, Levine-Boston I p = 0.009) No significant difference for SCG. VAS for HTG 17.9mm.</td>
<td>“Hyperthermia produced short-term improvements in pain and function in patients with mild to moderate carpal tunnel syndrome in the absence of any sizeable change in neurophysiologic parameters.”</td>
<td>Small sample size. Study represented as double blinded, but cannot blind this type of study design using heat.</td>
</tr>
<tr>
<td>Incebiyik 2014</td>
<td>RCT</td>
<td>Double-blind, No mention of sponsorship or COI</td>
<td>4.5</td>
<td>N = 31 females with mild and moderate CTS. Mean age for Group 1 51±10.07 and for Group 2 44.9±10.84.</td>
<td>Group 1 hot pack, Short-wave diathermy or SWD, and gliding exercises for 15 sessions, 5 times weekly (n = 15) vs. Group 2 hot pack, placebo for SWD, and gliding exercises for 15 sessions, 5 times weekly (n = 13). Follow-up at baseline and at 3 weeks.</td>
<td>At baseline vs. 3 weeks, between-group comparison: Tinel test/Phalen test/Reverse Phalen test/Carpal compression test/VAS/Levine-Boston Symptom Severity Scale or SSS/Functional Status Scale or FSS; p &lt;0.001 group 1 vs. p = 0.500 group 2/p &lt;0.001 vs p = 1.000/p &lt; 0.001 vs p=1.000/p &lt; 0.001 vs p = 1.000/p &lt; 0.001 vs p = 0.234/p &lt; 0.001 vs p = 0.204.</td>
<td>“SWD provided short-term improvements in pain, clinical symptoms, and hand function in patients with mild and moderate CTS.”</td>
<td>Data suggest treatment superior to placebo. Many cointervention s poorly tracked. Trial susceptible to contact time bias.</td>
</tr>
</tbody>
</table>

ULTRASOUND (Therapeutic)
Ultrasound has been used to treat many MSDs including CTS.(805, 831-834)

Recommendation: Ultrasound for Acute, Subacute, or Chronic CTS in Select Patients Who Fail Splint Use or Decline Injection

There is no recommendation for or against ultrasound for treatment of acute, subacute, or chronic CTS.
Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
The highest quality trial found ultrasound to be ineffective compared with sham ultrasound where both groups were treated with splinting. (640) One moderate-quality study found modest efficacy comparing ultrasound with placebo. (831) Another study had no placebo control and found ultrasound superior to low level laser therapy. (805) One trial found ultrasound comparable to glucocorticosteroid injection. (835) The remaining quality studies included co-interventions (611, 637) or had a lower quality rating and mostly suggested lack of efficacy. (832)

Ultrasound is not invasive, has few adverse effects, and is moderate to high cost depending on the number of treatments (which were numerous in the quality studies). As the available studies substantially conflict, there is no recommendation for or against therapeutic ultrasound. However, some evidence suggests possible efficacy of phonophoresis (see phonophoresis).

Evidence for the Use of Ultrasound for CTS
There are 1 high- (640) and 7 moderate-quality (611, 637, 805, 831, 833, 835, 836) RCTs incorporated into this analysis. There are 4 low-quality RCTs in Appendix 2. (785, 832, 837, 838)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: ultrasound therapy, carpal tunnel syndrome, median nerve neuropathy, median neuropathy, median nerve disease entrapment, neuropathy nerve compression, burning, itching, numbness, tingling, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 56 articles in PubMed, 6329 in Scopus, 8 in CINAHL, 43 in Cochrane Library and 0 in other sources. We considered for inclusion 11 from PubMed, 0 from Scopus, 2 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 18 articles considered for inclusion, 13 randomized trials and 1 systematic review met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ultrasound vs. Placebo</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yildiz 2011 RCT</td>
<td>8.0</td>
<td>N = 51 (25 median nerves; 43 female/8 male) with signs and symptoms of CTS for more than a month and mild-to-moderate CTS after electrodiagnostic test confirmation. Age range 39-66 years.</td>
<td>Group 1: sham ultrasound (US), ultrasound system in off mode. 15 minute sessions once a day 5 times a week for 2 weeks plus splinting with a neutral custom-molded thermoplastic volar wrist splint at night and during the day for 8 weeks (n = 17, 25 median nerves) vs. Group 2: US, pulse mode (1:4) with gel without medication at 1 MHz frequency and 1 W/cm² intensity plus splinting (n = 17, 26 median nerves) vs. Group 3: ketoprofen phonophoresis (PH), US pulse mode (1:4) with 2.5% ketoprofen gel at 1 MHz frequency and 1 W/cm² intensity plus splinting (n = 17, 25 median nerves). Follow-up for 8 weeks.</td>
<td>Mean±SD VAS (baseline/2 week/8 week): Group 1, 5.76±2.45/2.72±2.07/3.28±2.74 vs. Group 2, 4.96±2.50/2.43±2.77±2.74 vs. Group 3, 6.04±2.40/3.03±1.96/0.98±1.65 (p = 0.002. Group 3 &gt; Group 1; p = 0.004. Group 3 &gt; Group 2).</td>
<td>“Our results suggest that ketoprofen PH in addition to splinting is superior to the combination of US and splinting with respect to pain only in middle term patients with CTS.”</td>
<td>Ultrasound plus splinting was not superior to splinting alone. Ketoprofen plus splinting was associated with a reduction in pain at 8 weeks.</td>
</tr>
<tr>
<td>Ebenbiche 1998 RCT</td>
<td>6.5</td>
<td>N = 45 (gender not specified) with mild to moderate CTS. Mean age51.</td>
<td>Ultrasound daily 15 minute sessions, 5x a week for 2 weeks then twice a week for 5 more weeks, 1MHz with intensity 1.0W/cm², pulsed mode duty cycle of 1:4 and transducer area of 5cm² (n = 45 wrists) vs. sham ultrasound (n = 45 wrists). Follow-up period 6 months.</td>
<td>Main changes in symptom complaints were (active/sham): Week 2 (-1.05/0.05, p = 0.015), end of therapy (-0.17/-2.14, p = 0.001) and 6 months (-0.08/-2.76, p &lt;0.005). Grip strength measures improved (p&lt;0.0005). EDS measures improved (p&lt;0.05).</td>
<td>“There are satisfying short to intermediate term effects due to ultrasound treatment in patients with mild to moderate idiopathic carpal tunnel syndrome.”</td>
<td>Suggests ultrasound efficacious. High numbers of treatments (20). No assessment of blinding provided.</td>
</tr>
<tr>
<td><strong>Ultrasound vs. Other Treatments or in Combination(s)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bakhtiyari 2004 RCT</td>
<td>7.0</td>
<td>N = 40 (age not specified) and</td>
<td>Ultrasound, 15 minute sessions with frequency of 1 MHz and</td>
<td>VAS pain / severity of symptoms / functional status / grip strength, (p &lt; 0.001) and two point discrimination (p &lt;0.016). Group A, improved for all clinical outcomes, (p &lt;0.001), except the grip strength.</td>
<td>“Both ultrasound treatment and corticosteroid injection plus splinting were effective on the clinical symptoms and the electrophysiological findings of CTS.”</td>
<td>Both groups improved meaningfully over time, but differences between groups minimal with only one significant difference.</td>
</tr>
</tbody>
</table>

**Notes:**
- **RCT:** Randomized Controlled Trial
- **COI:** Conflicts of Interest
- **Score:** Evaluation of the study using a specific scoring system (5-11)
- **Sample Size:** Description of the sample size and demographic details
- **Results:** Summary of the study results
- **Conclusion:** Summary of the study conclusions
- **Comments:** Additional comments or considerations

**Copyright© 2016 Reed Group, Ltd.**
<table>
<thead>
<tr>
<th>RCT</th>
<th>Sponsored by grant from Semnan Medical Sciences University. No mention of COI.</th>
<th>10 with bilateral and unilateral CTS confirmed by electromyograph y or 90 wrists. Age means for laser/ultrasound groups: 48 (13.4)/45 (17.1). intensity of 1.0W/cm², pulsed mode duty cycle of 1:4 and transducer area 5cm² (N = 45) vs. low-level laser therapy, applied low intensity 9J, infrared laser diode, 830nm at 5 points, 1.8J/poinpt, daily 15 minute sessions 5 times a week (n = 45). Follow-up for 3 weeks. 0.2, p = 0.003. Other electrodiagnostic measures all favored ultrasound. VAS pain scores were -6.3 vs. -2.0, p &lt;0.001 at 4 weeks after treatment completion. therapy for treatment of carpal tunnel syndrome. treatments (15) in protocol is high.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baysal 2006</td>
<td>No mention of sponsorshi p and COI.</td>
<td>N = 36 (72 wrists) females with bilateral CTS, EDS confirmed. Mean age Group 1 47.8±5.5 years, Group 2 50.1±7.3, Group 3, 51.4±5.2 years. Group 1: tendon- and nerve-gliding exercises 5 daily sessions, each exercise repeated 10 times at each session for 3 weeks plus splinting with custom made neutral volar splint for 3 weeks all night and during the day (n = 12) vs Group 2: ultrasound administered 15 minutes per session to the palmar carpal tunnel area at frequency of 1 MHz and intensity of 1.0 W/cm², 15 treatments once a day, five time a week for 3 weeks plus splinting (n = 12) vs Group 3: ultrasound, splinting and exercises (n = 12). Full-time splint use; 8 week treatment. Assessments at first treatment, end of therapy, and after 8 weeks follow-up. Pain score before treatment/after treatment /after 8 weeks follow-up: Group I: 4.8±2.3/3.3±2.9/ 2.6±2.8; Group II: 5.7±2.7/2.2±1.9/ 2.5±2.8; Group III: 5.6±3.5/1.3±1.8/ 0.8±0.9. Functional status score: Group I: 20.6±7.8/14.8±7.5/ 14.9±6.6; Group II: 21.9±9.1/16.1±8.5/ 16.1±8.7; Group III: 20.5±7.1/11.7±6.6/ 12.6± 3.4. NS between groups for study outcomes. &quot;The result of this study emphasizes the efficacy of conservative treatment in CTS. In all patient groups, the treatment combinations were significantly effective immediately and 8 weeks after the treatment.&quot; Results suggest ultrasound may have some benefits, although it was not compared to a sham, placebo or no treatment. All groups were splinted.</td>
</tr>
<tr>
<td>Davis 1998</td>
<td>RCT</td>
<td>N = 91 with self-reported symptoms of CTS and EDS confirmed CTS. Mean age ibuprofen group 38±5 year, manipulation group 36±6 years. Ibuprofen (800mg 3x a day for 1 week, then 2x a day for 1 week, then PRN 7 weeks) and nocturnal cock-up wrist supports (n = 46) vs. high velocity, low amplitude manual thrust procedures: manipulation to upper extremity and spine (3 treatments a week for 2 weeks; 2 treatments a week for 3 weeks; 1 treatment a week for 4 weeks) plus ultrasound applied over the carpal tunnel for half of chiropractic treatment visits, 1 MHz and 1.0-1.5 W/cm² at 50% duty cycle for 5 minutes plus CTS outcome assessment physical distress (mean±SD) baseline to end of study: IBU and splint 14.66±9.89 to 5.7±6.28 vs. ultrasound and manipulation 12.47±8.07 to 9.25±8.14 (p = 0.0132). CTS outcome assessment mental distress (mean±SD) baseline to end of study: IBU and splint 33.61±12.02 to 14.94±11.33 vs. ultrasound and manipulation 28.94±11.69 to 17.29±13.24 (p = 0.0085). &quot;Carpal tunnel syndrome associated with median nerve demyelination but not axonal degeneration may be treated with commonly used components of conservative medical or chiropractic care.&quot; Baseline did not exclude prior ibuprofen use or manipulation, but prior use of these treatments is likely differential between 2 groups and potentially fatal study flw. Ibuprofen use PRN after 2 weeks and subject contact differed between groups bias in favor of manipulation/ ultrasound. High dropout rates. Study mainly compares variable dose ibuprofen vs. manipulation plus ultrasound as both</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Diagnosis</td>
</tr>
<tr>
<td>-------</td>
<td>----</td>
<td>-----------</td>
</tr>
<tr>
<td>Chang 2014 RCT</td>
<td>60</td>
<td>Nocturnal wrist supports (n = 45). Study duration: 9 weeks. Assessments at baseline and end of study.</td>
</tr>
<tr>
<td>Piravej 2004 RCT</td>
<td>18</td>
<td>N = 18 females (30 hands) with mild to moderate CTS for less than 12 months (mean 6.53±4.33 months), no treatment for at least 1 month and no steroid injections in last 3 months. Age range 33-68</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Diagnosis</th>
<th>Age</th>
<th>Treatment</th>
<th>Follow-up</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chang 2014 RCT</td>
<td>60</td>
<td>Nocturnal wrist supports (n = 45). Study duration: 9 weeks. Assessments at baseline and end of study.</td>
<td>45.9 years</td>
<td>Paraffin therapy, Twice per week. (N = 30) vs. Group 2: ultrasound + splint only, twice per week. (n = 30) Follow up period: 8 weeks after treatment.</td>
<td>No significance between group difference in EDS.</td>
<td>Minimal differences seen between groups. Data suggests ultrasound and splint not superior to paraffin and splint.</td>
</tr>
<tr>
<td>Piravej 2004 RCT</td>
<td>18</td>
<td>N = 18 females (30 hands) with mild to moderate CTS for less than 12 months (mean 6.53±4.33 months), no treatment for at least 1 month and no steroid injections in last 3 months. Age range 33-68</td>
<td>4.5</td>
<td>Ultrasound 0.5 W/cm² for 10 minutes, 5 days a week for 4 weeks plus placebo (n = 15 hands) vs ultrasound 0.0 W/cm² plus diclofenac 75mg a day (n = 15 hands). Follow-up within 5 days after 4 weeks of treatment.</td>
<td>Significant improvements in symptom severity scores seen in both groups. The effect size (ES) of the symptom severity scores was 0.63 for both groups. However, significant improvements in functional status scores (ES 0.38) and pain scales (ES 0.74) only seen in US therapy group. An effect size of 0.3 to 0.8 is considered a &quot;moderate&quot; effect.</td>
<td>“To improve the functional status of CTS patients, a combination of ultrasound therapy and a wrist orthosis may be more effective than a combination of paraffin therapy and a wrist orthosis. Since this is an exploratory trial, further confirmatory testing is suggested to justify the efficacy of these two treatments.”</td>
</tr>
</tbody>
</table>

**Ultrasound vs. Ultrasound plus NSAID**

<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Diagnosis</th>
<th>Age</th>
<th>Treatment</th>
<th>Follow-up</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Piravej 2004 RCT</td>
<td>18</td>
<td>N = 18 females (30 hands) with mild to moderate CTS for less than 12 months (mean 6.53±4.33 months), no treatment for at least 1 month and no steroid injections in last 3 months. Age range 33-68</td>
<td>4.5</td>
<td>Ultrasound 0.5 W/cm² for 10 minutes, 5 days a week for 4 weeks plus placebo (n = 15 hands) vs ultrasound 0.0 W/cm² plus diclofenac 75mg a day (n = 15 hands). Follow-up within 5 days after 4 weeks of treatment.</td>
<td>Significant improvements in symptom severity scores seen in both groups. The effect size (ES) of the symptom severity scores was 0.63 for both groups. However, significant improvements in functional status scores (ES 0.38) and pain scales (ES 0.74) only seen in US therapy group. An effect size of 0.3 to 0.8 is considered a &quot;moderate&quot; effect.</td>
<td>“To improve the functional status of CTS patients, a combination of ultrasound therapy and a wrist orthosis may be more effective than a combination of paraffin therapy and a wrist orthosis. Since this is an exploratory trial, further confirmatory testing is suggested to justify the efficacy of these two treatments.”</td>
</tr>
</tbody>
</table>

**No COI.**
| University. No mention of COI. | years, mean age 46.97 ± 8.37 years. |   |   |   |   |   |


Electrical Therapies
PHONOPHORESIS
Phonophoresis involves the use of ultrasound to deliver topically applied drugs and has been used to treat patients with CTS.(839)

Recommendation: Phonophoresis for Acute, Subacute, or Chronic CTS
Phonophoresis is recommended for treatment of acute, subacute, or chronic CTS.

Indications – CTS that is sufficiently symptomatic to warrant treatment. Patients should generally be given splints and/or a glucocorticosteroid injection prior to considering phonophoresis as a splint or injection are believed to be more effective.

Frequency – The regimen in the highest quality study consisted of 5-15 sessions per week for 4-8 weeks with ketoprofen phonophoresis (PH),(840) US pulse mode (1:4) with 2.5% ketoprofen gel at 1 MHz frequency and 1 W/cm2 intensity.(640) Dexamethasone has also been successfully used,(783, 840) with one trial suggesting the steroid is superior to NSAID (diclofenac). (783) Other NSAIDs and glucocorticoids are presumably equally efficacious.(640)

Discontinuation – Resolution, failure to objectively improve or intolerance.

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – Low

Rationale for Recommendation
One high-quality comparative trial found ketoprofen phonophoresis plus splinting superior to ultrasound plus splinting.(640) One moderate quality comparative trial found dexamethasone administered by phonophoresis superior to iontophoresis.(840) One moderate quality comparative trial found phonophoresis with glucocorticoid superior to phonophoresis with diclofenac or splinting.(783) Phonophoresis is not invasive, has low adverse effects, and is moderately costly. However, phonophoresis with either NSAID or dexamethasone is recommended particularly where splinting is insufficiently controlling symptoms and an injection is declined.

Evidence for the Use of Phonophoresis
There is 1 high-(640) and 2 moderate-quality(783, 840) RCTs incorporated into this analysis. There are 2 low-quality RCT in Appendix 2.(786, 839)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Phonophoresis or phonophoresis, carpal tunnel syndrome, CTS, median nerve neuropathy, median neuropathy, median nerve disease entrapment, neuropathy nerve compression, burning, itching, numbness, tingling, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 19 articles in PubMed, 6 in Scopus, 11 in CINAHL, 43 in Cochrane Library and 0 in other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 4 articles considered for inclusion, 4 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yildiz 2011</td>
<td>RCT</td>
<td>8.0</td>
<td>N = 51 (25 median nerves; 43 female/8 male) with signs and symptoms of CTS for more than a month and mild-to-moderate CTS after electrodiagnostic test confirmation. Age range 39-66 years.</td>
<td>Group 1: sham ultrasound or US, ultrasound in off mode 15 minute sessions once a day 5x a week for 2 weeks plus splinting with neutral custom-molded thermoplastic volar wrist splint at night and during day (n = 17) vs. Group 2: US, pulse mode (1:4) with gel without medication at 1 MHz frequency and 1 W/cm² intensity plus splinting (n = 17) vs. Group 3: ketoprofen phonophoresis (PH), US pulse mode (1:4) 2.5% ketoprofen gel at 1 MHz frequency and 1 W/cm² intensity plus splinting (n = 17). Follow-up for 8 weeks.</td>
<td>Mean±SD VAS (baseline/2 week/8 week): Group 1, 5.76 ±2.45/2.72 ± 2.07/3.28 ± 2.74 vs. Group 2, 4.96 ± 2.50/2.41 ± 2.43/2.77 ± 2.74 vs. Group 3, 6.04 ± 2.40/3.03 ± 1.96/0.98 ± 1.65 (p = 0.002. Group 3 &gt; Group 1; p = 0.004, Group 3 &gt; Group 2). Pain score significantly lower in Group 3 at 8th week compared to other treatment groups (Group 1 and Group 2) (p = 0.002, p = 0.004 and p = 0.001).</td>
<td>&quot;Ketoprofen PH as adjuvant therapy on splinting is effective with respect to reduction of pain.&quot;</td>
<td>Ultrasound plus splinting not superior to splinting alone. Ketoprofen plus splinting was associated with a reduction in pain at 8 weeks.</td>
</tr>
<tr>
<td>Bakhtiary 2013</td>
<td>RCT</td>
<td>7.0</td>
<td>N = 34 (gender not specified) with mild to moderate CTS confirmed by electromyography. Mean age for Iontophoresis and Phonophoresis: 48.2 (14.5) and 44.6 (12.8).</td>
<td>Iontophoresis of Dex-P 0.4% (n = 26) vs. Phonophoresis of Dex-P 0.4%, plus applied over wrist chin, and pulsed (20%) ultrasound waves (n = 26). Follow-up for 4 weeks.</td>
<td>Pain at end of treatment and 4 weeks later significantly favored phonophoresis vs. iontophoresis of Dex-P intervention, (p &lt;0.01). Motor latency/motor action potential amplitude/finger pinch strength/ hand grip strength/ and pain relief: [mean difference 0.8 m/s; 95% (CI), 0.5-1.1]/(4.1 mV; 95% CI, 3.0 - 5.2)/(31.6 N; 95% CI, 15.9-47.3)/(27.1 N; 95% CI, 13.5-40.5)/and 2.1 points on 10-point scale; 95% CI, 1.3-2.9.</td>
<td>&quot;Our clinical trials showed that phonophoresis of Dex-P is more effective than iontophoresis of Dex-p treatment in patients with mild to moderate CTS.&quot;</td>
<td>Data suggest phonophoresis superior to iontophoresis</td>
</tr>
<tr>
<td>Soyupek 2012</td>
<td>RCT</td>
<td>4.5</td>
<td>N = 52 with CTS, EDS confirmed. Mean age splinting, PCS, PNSAI: 47.95±6.93 years, 50.50±8.71 years, 53.79±10.40 years.</td>
<td>Phonophoresis with corticosteroid (betamethasone valerate %0.1 cream), CS (PCS) over carpal tunnel for 10 min/session at frequency of 3 MHz and intensity of 1.5 W/cm² 5 times a week for 3 weeks (n = 28) vs. phonophoresis with non-VAS difference baseline to after 3 months, mean±SD (baseline/after 3 months): splinting group 50.69±23.45/37.91±23.94 (NS); PCS 60.35±18.95/30.35±18.15 (p &lt;0.017); PNSAI 69.13±16.21/45.65±23.65 (p &lt;0.017). Boston</td>
<td></td>
<td>&quot;[T]he most effective treatment modality for CTS was P-CS according to ultrasonographic investigations and other findings.&quot;</td>
<td>PCS group better than splinting or PNSAI groups.</td>
</tr>
</tbody>
</table>
steroidal anti-inflammatory drug (diclofenac diethylammonium gel), NSAI (PNSAI) over carpal tunnel for 10 min/session at frequency of 3 MHz and intensity of 1.5 W/cm² 5x a week for 3 weeks (n = 23) vs. wrist splinting in neutral position during the day and at night for the first 15 days and then when CTS was symptomatic (n = 23). Follow-up 3 months after treatment.

| Questionnaire total difference from baseline to after 3 months, mean±SD (baseline/after 3 months): | splinting group 43.34±10.89/39.26±10.03 (NS); PCS 54.21±11.34/39.14±10.33 (p <0.017); PNSAI 53.69±41.86/41.86±10.03 (p <0.017). Tinel’s sign, %, difference from baseline to after 3 months (baseline/after 3 months): splinting group 65.2/60.9 (NS); PCS 82.1/50.0 (p <0.017); PNSAI 82.6/65.2 (NS). Phalen’s sign, %, difference from baseline to after 3 months (baseline/after 3 months): splinting group 60.9/52.2 (NS); 89.3/50.0 (p <0.017); PNSAI 78.3/39.1 (p <0.017). |
IONTOPHORESIS

Iontophoresis, a drug-delivery system that utilizes electrical current to transdermally deliver either glucocorticosteroids or NSAIDs, has been used to treat distal upper extremity MSDs including CTS. (839, 841, 842) It is believed to be more efficacious in situations where the dermis and adipose tissue overlying the target tissue is thin which facilitates penetration of the pharmaceutical to the target tissue and may be somewhat of an obstacle for treatment of CTS.

**Recommendation: Iontophoresis for Acute, Subacute, or Chronic CTS**

There is no recommendation for or against the use of iontophoresis for treatment of acute, subacute, or chronic CTS.

**Strength of Evidence – No Recommendation, Insufficient Evidence (I)**

**Level of Confidence – Low**

**Rationale for Recommendation**

Iontophoresis has been studied for the treatment of CTS. There is one moderate-quality study comparing iontophoresis with dexamethasone versus distilled water which reported no benefit. (841) However, it was small in size (n = 20) and appears underpowered. The other moderate-quality study found injection to be superior. (842) There is no quality study of sufficient size comparing iontophoresis with placebo, precluding an assessment of quality evidence of efficacy. Iontophoresis with glucocorticosteroid may be a reasonable option for treating patients who decline injection; however, oral glucocorticosteroids have quality evidence of efficacy and may be recommended preferentially as iontophoresis is believed to be less effective than glucocorticosteroid injections. (842) Iontophoresis is not invasive, has low adverse effects, and is of moderate cost. However, other treatments have documented efficacy and should be used preferentially.

**Evidence for the Use of Iontophoresis for CTS**

There are 2 moderate-quality RCTs incorporated into this analysis. (841, 842) There are 2 low-quality RCT in Appendix 2. (786, 839)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Iontophoresis or phonophoresis, carpal tunnel syndrome, median nerve neuropathy, median nerve disease entrapment, neuropathy nerve compression, burning, itching, numbness, tingling, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 19 articles in PubMed, 6 in Scopus, 11 in CINAHL, 43 in Cochrane Library and 0 in other sources. We considered for inclusion 2 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 2 articles considered for inclusion, 2 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amirjani 2009</td>
<td>7.5</td>
<td>N = 20 (19 female/1 male) with mild to moderate NCS confirmed (19 females; 1 male). Mean age: 54 ±10 years</td>
<td>Dexamethasone sodium phosphate in distilled water 0.4% (n = 10) vs. distilled water iontophoresis 80mA a minute continuous DC current at 2mA a minute over carpal tunnel, 6 treatments QOD over 2 week (n = 10). Follow-up for 6 months.</td>
<td>Levine Self-Assessment Questionnaire scores median (25th-75th % CI) (baseline/post first treatment/post 6 treatments): Dex [38 (31-40)/33 (30-48), 26 (24-31)] vs. water controls (36 (33-54)/38 (27-44)/34 (22-41)), (p = 0.73, p = 0.91, p = 0.25))</td>
<td>“Although corticosteroid iontophoresis is feasible in clinical settings and is well-tolerated by patients, iontophoresis of 0.4% dexamethasone was not effective in the treatment of mild to moderate CTS.”</td>
<td>Small sample size. Stratified baseline data not provided. Appears underpowered, although magnitude of a potential benefit also not likely high or moderate.</td>
</tr>
</tbody>
</table>

Copyright © 2016 Reed Group, Ltd.
Injection Therapies

Four major types of injections have been utilized to treat patients with CTS. These include: 1) carpal tunnel injections with glucocorticosteroids (discussed previously); 2) carpal tunnel injections with insulin among diabetics; 3) intramuscular glucocorticosteroid injections; and 4) botulinum injections.

Intracarpal Tunnel Glucocorticosteroid Injections (“Steroid Injections”)

Steroid injections of the carpal canal are frequently performed to treat CTS patients,(631, 650, 777, 789, 839, 842-855) including those with acute cases (i.e., those that typically occur with fractures, trauma, or unaccustomed high-force use and present primarily with acute flexor wrist pain).(856-858) While various injection techniques have been utilized (including distal to proximal), the most common technical injection approach utilizes a fine gauge needle (e.g., 27- or 25-gauge) entering the skin near the distal wrist crease to the medial (ulnar) side of palmaris longus, and angled approximately 45 degrees distally. While it has been suggested that these injections are underutilized,(859) steroid injections should be done by those experienced with administering these injections.

1. Recommendation: Carpal Tunnel Injections for Treatment of Subacute or Chronic CTS

Carpal tunnel injections are strongly recommended for the treatment of subacute or chronic CTS.

Indications – CTS unresponsive to nocturnal wrist splinting, generally with symptoms lasting at least 3 weeks. It is not believed to be necessary to perform EDX prior to injections.

Dose – One high-quality study found lower 1-year surgery rates with methylprednisolone 80mg vs. 40 mg of 73% vs. 81%, which were also superior to placebo.(860) Generally, at least 40mg of methylprednisolone or equivalent is recommended as the minimum initial dose. Although optimum dose remains unclear, evidence in total includes evaluations with methylprednisolone acetate (12, 15, 20, 40, 60mg, 80mg), betamethasone (6.0, 6.4mg), triamcinolone hexacetonide (20mg), and hydrocortisone (25, 100mg) in quality studies. Some physicians increase the dose in proportion to perceived symptom severity. However, there is no quality evidence to support this practice. The type of steroid to inject and whether to use a depot preparation, are also unclear as there are no quality studies comparing the various preparations commonly utilized. Most physicians include at least 1mL of an injectable anesthetic (e.g., 1% lidocaine). Lidocaine allows for rapid assessment immediately after the injection. The limitation of using an anesthetic as an adjuvant is that the numbness that ensues afterwards may limit a patient’s activities. Thus, a shorter-duration anesthetic such as lidocaine is recommended.

Frequency/Duration – A single injection and the results carefully evaluated to document improvement, even if short-term as it is believed to have considerable prognostic significance. There is no evidence that a series of injections is efficacious, although it has been argued that two injections are ideal.(861) There is no evidence that there is a limit to the number of injections to treat an episode or in a lifetime. Failure to respond, particularly if the median nerve was successfully
anesthetized by the injection, should result in a careful re-assessment of the accuracy of the
diagnosis of CTS. A second injection, typically utilizing a moderately higher dose, may be indicated if
there has been insufficient but partial relief, or if the first injection was thought to have not entered the
carpal canal.

*Indications for Discontinuation* – No partial response to carpal tunnel injection(s), then no
recommendation for additional injection(s). Patients who fail to even partially respond to injections
are a priori suspected to not have CTS and a thorough search for an alternate diagnosis should
ensue. Patients who respond to carpal tunnel injections, but redevelop symptoms are believed to be
ideal candidates for surgical release.

*Strength of Evidence – Strongly Recommended, Evidence (A)*
*Level of Confidence – High*

2. *Recommendation: Carpal Tunnel Injections for Treatment of Acute CTS without Fracture*

Carpal tunnel injections are recommended for treatment of acute CTS without fractures.
*(Acute CTS with fractures should be referred for potential emergent surgical release.)*

*Indications/Dose/Frequency –* See above Recommendation #1.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*
*Level of Confidence – Moderate*

Rationale for Recommendations
There is strong consistent evidence that carpal tunnel injections are efficacious with superiority to
placebo.(631, 777, 839, 842, 843, 845, 849, 851, 854, 860) There also is evidence that injections are
superior to oral glucocorticosteroids(648) and iontophoresis with glucocorticosteroids.(842) Duration of
improvements after injection is controversial and may differ by CTS severity. Nearly all quality studies
required electrodiagnostic confirmation and many had patients with symptoms lasting years, suggesting
more severely affected patients benefited. In such patients, injections may be somewhat less efficacious
than in patients with more recent or mild symptoms that are seen initially in primary care settings.

Aside from local tenderness among 50% of patients lasting a mean 1.2 days,(862) long-term
complications are rare. Long-term outcomes are somewhat controversial. One study of 30 patients found
11.4% remained asymptomatic over an 80-week observation period with more undergoing surgery if
treatment had been via an oral steroid than via an injection. Other studies reported only 22% of injected
patients were subsequently referred for surgery during 1 year of follow-up.(845)

Steroid injections into the carpal tunnel are minimally invasive, have minimal adverse effects, and are
moderately costly. These injections provide lasting relief of at least intermediate-term durations in a
majority of CTS patients. They are strongly recommended for treatment of subacute or chronic CTS.

Carpal tunnel injections are also recommended by consensus of the Evidence-based Practice Hand,
Wrist, and Forearm Panel for treatment of acute CTS in cases where there are no fractures. There are
no quality studies of these clinical cases; however, these injections are thought to be the best treatment
for acute CTS presentations.

*Evidence for the Use of Glucocorticosteroids (Oral and Injection) for CTS*
There are 8 high-(646, 648, 843-845, 851, 855, 860) and 19 moderate-quality(631, 636, 643, 644, 647,
777, 835, 840, 842, 848, 849, 852-854, 863-868) RCTs (one with two reports) incorporated into this
analysis. There are 5 low-quality RCT and 1 prospective randomized blinded trial in Appendix 2.(786,
789, 839, 869-871)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane
Library without date limits using the following terms: glucocorticoids, glucocorticosteroids, carpal tunnel
syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment,
neuropathy, nerve disease, syndrome, median nerve, median neuropathy, burning, itching, numbness,
tingling, hand, palm, finger, wrist, pain; controlled clinical trial, controlled trials, randomized controlled
trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 109 articles in PubMed, 268 in Scopus, 5 in CINAHL, and 46 in Cochrane Library. We considered for inclusion 30 from PubMed, 0 from Scopus, CINAHL, Cochrane Library and other sources. Of the 30 articles considered for inclusion, 30 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hui 2001</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>8.0</td>
<td>N = 36 (17 female/1 male) newly diagnosed CTS, EDS confirmed. Age mean for steroids/placebo groups: 44.9 (10.0)/42.9 (7.2).</td>
<td>Oral steroids or prednisolone 25mg a day (N = 18) vs. Placebo 10 day course for 10 days (N = 18). Follow-up at 2 and 8 weeks.</td>
<td>Median (IR) change at 2 weeks steroid -12.5 (-15 to -7) vs. placebo group -4.5 (-14 to 0). (p = 0.027). After 8 weeks, median (IR) reduction of GSS -9 (-14 to -6) vs. placebo -2 (-10 to 0). (p = 0.034).</td>
<td>“This study shows a small but statistically significant reduction in GSS in the group prescribed a short course of prednisolone as compared with placebo.”</td>
<td>Brief report. Data suggest superiority of glucocorticosteroids.</td>
</tr>
<tr>
<td>Chang 1998</td>
<td>RCT</td>
<td>Sponsored by NSC to Ming-Hong Chang. No mention of COI.</td>
<td>7.0</td>
<td>N = 91 (53 female/20 male) with clinical symptoms/signs of CTS and EDS confirmed. Mean age placebo/diuretic/NSAID/steroid groups: 44.2±5.4/45.7±4.8/47.4±5.7/45.4±5.2</td>
<td>NSAID-SR with Tenoxicam-SR, 2mg daily (N = 18) vs. Diuretic treatment with Tenoxicam-SR, 20mg daily (N = 16) vs. Steroid or Prednisolone 2 weeks 20mg daily, then 10mg daily (N = 23) vs. Placebo group (N = 16). Follow-up for 4 weeks.</td>
<td>No significant reduction from baseline GSS seen at 2nd and 4th weeks in placebo, NSAID-SR, and diuretic groups. However, mean score at 4 weeks in steroid group decreased significantly from baseline of 27.9±6.9 to 10±7.4.</td>
<td>“For patients with mild to moderate CTS who opt for conservative treatment, corticosteroids are of greater benefit.”</td>
<td>Suggests oral steroids effective, but diuretic and NSAID are not.</td>
</tr>
<tr>
<td>Chang 2002</td>
<td>RCT</td>
<td>Sponsored by grants VGHKS, NSC, and TCVGH. No COI.</td>
<td>6.5</td>
<td>N = 109 (81 female/18 male) with CTS, EDS confirmed. Mean age 4 and 2 week treatment: 46.2 (5.4)/45.9 (5.1).</td>
<td>Group 1, received prednisolone 20mg daily for 2 weeks then 10mg for 2 weeks (n = 53) vs. Group 2 received prednisolone 20mg daily and 2 weeks of placebo (n = 56). Follow-up for 4 weeks.</td>
<td>Overall improvement (%) (4 week/2 week): Month 1: 66.0/48.2 (35/27); Month 3: 64.1/44.6 (34/25); Month 6: 52/37.5 (28/21); Month 12: 49.0/35.7 (26/20), (NS). Global symptom scores (4-week treatment/2-week treatment): Baseline: mean (SD): 17.88 (6.06)/17.24 (4.95); Month 1: 8.1 (6.8)/ 9.54 (6.56); Month 3: 8.08 (7.72)/ 9.1 (7.95); Month 12: 10.25 (10.6)/10.61 (9.15), (NS).</td>
<td>“In patients with CTS, a 2-week course of prednisolone was as effective as a 4-week course in improving symptoms.” “Short term low dose oral steroids are effective treatment for carpal tunnel syndrome. The dose of steroids and the duration treatment are not key determinants of efficacy.”</td>
<td>Suggests minimal difference with 4 week course vs. 2 weeks course.</td>
</tr>
<tr>
<td>Herskovitz 1995</td>
<td>RCT</td>
<td></td>
<td>6.0</td>
<td>N = 15 (12 female/3 male) suspected CTS. Mean for</td>
<td>Prednisone, 20mg daily 1st week followed by 2nd week of 10 mg daily (n = 6) vs. Placebo matched</td>
<td>Prednisone-treated group better at 2 weeks (p &lt;0.05). After treatment discontinued, 4- and 8-week mean GSS not different.</td>
<td>“This approach may provide a treatment alternative in the short-term, conservative management of CTS.”</td>
<td>Small sample size.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>Sponsorship or COI</td>
<td>Participants</td>
<td>Interventions</td>
<td>Outcomes</td>
<td>Comments</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>--------</td>
<td>-------------------</td>
<td>-------------</td>
<td>--------------</td>
<td>---------</td>
<td>---------</td>
<td></td>
</tr>
<tr>
<td>Mishra 2006</td>
<td>4.0</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>N = 66 with symptoms suggestive of CTS at least 1-month duration. Age range from 28 to 60 years.</td>
<td>Splint group carpal tunnel (n = 36 hands) vs. steroid or oral prednisolone 20mg a day for 2 weeks followed by 10mg a day for another 2 weeks (n = 36 hands). Follow-up for 3 months.</td>
<td>Mean±SD for splint vs. steroid: symptom severity score (SSS): 0-1: 0.34±0.42 vs 0.40±0.30 (p = 0.52); 0-3: 0.30±0.54 vs. 0.49±0.44 (p = 0.42). Sensory distal latency (SDL): 0-1: 0.16±0.63 vs. 0.13±0.71 (p = 0.86); 0-3: 0.35±0.76 vs. 0.55±0.66 (p = 0.25).</td>
<td>&quot;There was significant improvement in both groups, clinically as well as electrophysio-logically, at 3 months.&quot;</td>
<td></td>
</tr>
<tr>
<td>No blinding. Suggests splinting is as effective as oral steroid, though function was slightly better with splinting.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Celiker 2002</td>
<td>5.5</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>N = 23 with bilateral or unilateral CTS, EDS confirmed. Mean age for Group A and B; 49.6±15.3 and 46.9±10.0.</td>
<td>Group A: acemetacine 120mg a day with splints at night. Group B: 40mg methylprednisolone acetate (1ml)</td>
<td>VAS pain scores (baseline/2nd week/8th week): NSAID vs. steroid: 7.9±1.4/4.3±0.9/1.7±1.0 vs. 7.0±2.2/3.1±2.5/1.8±1.9 (P&gt;0.05). Symptom severity scale results similar (P&gt;0.05).</td>
<td>&quot;Both splinting combined with the use of a nonsteroidal anti-inflammatory drug and steroid injection into the carpal tunnel resulted in significant improvement in carpal tunnel syndrome.&quot;</td>
<td></td>
</tr>
<tr>
<td>No placebo controlled. Suggests splinting and NSAID may be as effective as injection.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Karadas 2011</td>
<td>4.5</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>N = 99 with clinical and electrophysiologic evidence of CTS, older than 18 years.</td>
<td>Group 1 40mg triamcinolone acetonide (n = 34) vs. Group 2 4ml 1% procaine HCl (n = 32) vs. Group 3 both 40mg triamcinolone acetonide and 4ml 1% procaine HCl (n = 33). Follow-up at baseline, 2 and 6 months after injection.</td>
<td>VAS scores improved significantly in each group at 2 and 6 months after treatment, (p&lt;0.05). No significant differences for electrophysiologic findings at baseline, 2, and 6 months, (p &gt;0.05).</td>
<td>&quot;Local procaine HCl injection and steroid injection effectively reduced the symptoms of CTS and equally improved electrophysiologic findings.&quot;</td>
<td></td>
</tr>
<tr>
<td>Combined triamcinolone acetonide and procaine HCl may be superior to individual medications alone.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Karadas 2012</td>
<td>4.5</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>N = 57 with clinically suspected primary CTS. Age &gt;18 years.</td>
<td>Group 1 injected with 1ml 0.09% saline (n = 19) vs. Group 2 injected with 40mg triamcinolone acetonide (n = 20) vs. Group 3 injected with 4ml 1% procaine HCl (N = 18). Follow-up at 1, 2 and 6 months.</td>
<td>Clinical/electrophysiological evaluations improved significantly in groups 2 and 3 at post-treatment, (p &lt;0.05). No significant changes in group 1, (p &gt;0.05). Groups 2 and 3 better scores vs. group 1 at 2, 6 months, (p &lt;0.05). No difference between groups 2 and 3 in terms of change</td>
<td>&quot;Triamcinolone acetonide and procaine HCl injections are effective regarding short- and long-term outcomes compared with placebo injections, and procaine HCl injection was as effective as steroid injection.&quot;</td>
<td></td>
</tr>
<tr>
<td>Both active interventions superior to saline injection.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Gender</td>
<td>Diagnosis</td>
<td>Age</td>
<td>Treatment</td>
<td>Outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------------</td>
<td>----</td>
<td>--------</td>
<td>-----------</td>
<td>-----</td>
<td>-----------</td>
<td>----------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bakhtiyari 2013</td>
<td>7.0</td>
<td></td>
<td>CTS</td>
<td>48.2 ± 8.2</td>
<td>Iontophoresis of Dex-P 0.4% (n = 26) vs. Phonophoresis of Dex-P 0.4%, plus applied over wrist chin and pulsed (20%) ultrasound waves (n = 26). Follow-up for 4 weeks.</td>
<td>Pain at end of treatment and 4 weeks later significantly favored phonophoresis vs iontophoresis of Dex-P intervention, (p &lt; 0.01). Motor latency/motor action potential amplitude/finger pinch strength/hand grip strength/ pain relief: [mean difference 0.8 m/s; 95% (CI), 0.5-1.1]/ (4.1 mV; 95% CI, 3.0-5.2)/ (31.6 N; 95% CI, 15.9-47.3)/ (27.1 N; 95% CI, 13.5-40.5)/ and 2.1 points on 10-point scale; 95% CI, 1.3 - 2.9.</td>
<td>&quot;Our clinical trials showed that phonophoresis of Dex-P is more effective than iontophoresis of Dex-P treatment in patients with mild to moderate CTS.&quot;</td>
<td></td>
</tr>
<tr>
<td>Gökölğlu 2005</td>
<td>4.0</td>
<td></td>
<td>CTS</td>
<td>48.0 ± 8.2</td>
<td>Group 1: 40mg methylprednisolone acetate injected (n = 15) vs. Group 2: iontophoresis of dexamethasone sodium phosphate (n = 15). Follow-up for 2 and 8 weeks.</td>
<td>Symptoms severity scores (baseline/week 2/week 8): injection 2.7±0.8/1.9±0.7/ 1.6± 0.6 vs. iontophoresis 3.1±0.8/2.5±0.9/ 2.2±1.0 (p &lt;0.05) for Weeks 2 and 8 favoring injection. Functional status scale and VAS scores similarly favored injection.</td>
<td>&quot;Success of both iontophoresis of dexamethasone sodium phosphate and injection of corticosteroids, but symptom relief was greater at 2 and 8 weeks with injection of corticosteroids.&quot;</td>
<td></td>
</tr>
<tr>
<td>Bilgic 2010</td>
<td>5.5</td>
<td></td>
<td>CTS</td>
<td>47.33 (7.44) and 44.15 (9.30).</td>
<td>Group A, ultrasound treatment (n = 16) vs. Group B, local corticosteroid injection plus splinting (n = 18). Follow-up for 8 weeks.</td>
<td>VAS pain/severity of symptoms/functional status / grip strength. (p &lt;0.001) and two point discrimination (p &lt;0.016). Group A, improved for all clinical outcomes, (p &lt;0.001), except grip strength.</td>
<td>&quot;Both ultrasound treatment and corticosteroid injection plus splinting were effective on the clinical symptoms and the electrophysiological findings of CTS.&quot;</td>
<td></td>
</tr>
<tr>
<td>Dammers 2006</td>
<td>9.0</td>
<td></td>
<td>EDS</td>
<td>45/43/44</td>
<td>20mg methylprednisolone injections (n = 45) vs. 40mg methylprednisolone injections (n = 43) vs. 60mg methylprednisolone injections (n = 44). Follow-up for 3 months.</td>
<td>73% of 60mg, 53% of 40mg and 56% of 20mg groups symptom free or requiring no further treatment at 6 months. Only 22% treated with 1-2 injections methylprednisolone during first year referred to surgery (p &lt;0.05).</td>
<td>&quot;One injection of methylprednisolone close to the carpal tunnel reduces the number of patients requiring surgery.&quot; 60mg dose more effective than lower doses, with 2nd injection possibly increasing.</td>
<td>Injection site 4cm proximal to distal wrist crease.</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Age</td>
<td>Injection Method</td>
<td>Outcome</td>
<td>Recurrence</td>
<td>Notes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>----</td>
<td>-----</td>
<td>------------------</td>
<td>---------</td>
<td>------------</td>
<td>-------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Üstün 2013</td>
<td>4.0</td>
<td>N = 46 with idiopathic CTS. Mean age for US-guided/Palpation-guided group: 45.96±10.49/23.71±11.38.</td>
<td>US-guided device of 20mg methylprednisolone (n = 23) vs. Palpation-guided approach or blind injection group of 20mg methylprednisolone using ulnar side approach (n = 23). Follow-up at 6 and 12 weeks.</td>
<td>Scores for symptom severity and functional status improved at 6 and 12 weeks after the treatment, (p &lt; 0.05). Boston Carpal Tunnel Questionnaire (BCTQ) symptoms/function: 6 weeks: 1.33±0.55 and 12 weeks: 1.30±0.45 vs 1.41±0.59 and 1.67±0.73 Palpation group, (p &lt;0.001)/1.33±0.46 and 1.36±0.49 vs 1.52±0.87 and 1.86±1.09, (p &lt;0.001).</td>
<td>Both US-guided and blind steroid injections were effective in reducing the symptoms of CTS and improving the function, an earlier onset/better improvement of symptom relief suggests that US-guided steroid injection may be more effective than are blind injections in CTS.</td>
<td>Data suggest ultrasound guided injection superior to blind for providers with this level of experience.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wong 2001</td>
<td>9.0</td>
<td>N = 62 (53 female/7 male) with newly diagnosed CTS &gt;3 months.</td>
<td>Steroid or prednisolone 25mg PO QD for 10 days (n = 30) vs. Oral steroid or prednisolone acetate 15mg injection (n = 30). Follow-up for 12 weeks.</td>
<td>Global symptom scores (injection/oral): baseline (25.0±6.4/25.7±8.3), 2 weeks (13.6±7.5/17.8±10.0), 8 weeks (13.7±8.3/20.8±8.7), and 12 weeks (14.3±8.4/21.4±9.6). GSS scores borderline significant at 2 weeks (p = 0.07), but significant at 8 and 12 week follow-ups (p = 0.002 and p = 0.004).</td>
<td>Both US-guided and blind steroid injections were effective in reducing the symptoms of CTS and improving the function, an earlier onset/better improvement of symptom relief suggests that US-guided steroid injection may be more effective than are blind injections in CTS.</td>
<td>Data suggest injections superior to oral glucocorticosteroids.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Habib 2006</td>
<td>5.0</td>
<td>N = 42 with symptoms of CTS and EDS confirmed. Age &gt;18 years old.</td>
<td>Local corticosteroid classic injection (n = 21) vs. 2-3cm distal to the wrist crease. Both with 12mg methylprednisolone acetate with 0.15ml lidocaine (n = 21). Follow-up for 12 months.</td>
<td>81% of classical injection and 71% new method injection patients had favorable response rate after 3 weeks (p = 0.468). Procedure time 8.48±1.123 seconds in new method group vs. 26.71±32.83 in classical group (p = 0.021). Mean±SD grade of pain: new method 4.38±1.523 vs. classical method 3.62±1.071 (p = 0.065).</td>
<td>Local corticosteroid injection using the novel approach for the treatment of carpal tunnel syndrome is helpful, and the favorable response rates are comparable to those using the classic approach after 1, 3, 6, and 12 weeks.</td>
<td>Suggests traditional injection technique may be superior.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Özdönüm 1984</td>
<td>6.0</td>
<td>N = 37 with idiopathic CTS. Mean age 45.8±8.7 years.</td>
<td>Steroid injection, 1.5mg betamethasone disodium phosphate and acetate</td>
<td>7 from carpal injection group and 6 from IM injection group returned with symptoms after 1 month and</td>
<td>Suggests injections superior to intramuscular steroids.</td>
<td>Carpal injections appear superior to intramuscular steroids.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Intracarpal Tunnel Injection with Glucocorticosteroids vs. Saline or No Injection

<table>
<thead>
<tr>
<th>Study</th>
<th>Methodology</th>
<th>N</th>
<th>Age</th>
<th>Intervention Details</th>
<th>Outcomes</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>Armstrong 2004</td>
<td>RCT</td>
<td>81 (63 female/18 male)</td>
<td>18-80</td>
<td>Steroid injections or Betamethasone 6mg (n = 43) vs. Placebo group or saline (n = 36). Follow-up for 18 months.</td>
<td>Changes in median sensory latencies -0.19±0.27 vs. -0.04±0.14 (p = 0.01). Changes in symptoms scores also favored corticosteroid injections - 0.78 ±0.80 vs. -0.19 ±0.62 (p &lt;0.01). Satisfaction rates 70% vs. 34% (p = 0.001). In subsequent open label follow-up, additional injections performed per patient requests (up to 7 injections for a few); 18 (39.1%) referred for surgery, 37.0% reported adequate symptom relief.</td>
<td>Unblinded after 2 weeks.</td>
</tr>
<tr>
<td>Peters-Veluthamaningal 2010</td>
<td>RCT</td>
<td>69 (53 female/16 male)</td>
<td>Mean age: NaCl group = 57.6 years, TCA group = 56.5 years.</td>
<td>1ml triamcinolonacetonide (TCA) 10mg/ml (n = 36) vs. 1ml saline (NaCl) 0.9%, placebo 1-2 injections (n = 33). Follow-up 1, 3, 6 and 12 months</td>
<td>Steroid-group showed better direct treatment response (p = 0.013), perceived improvement (p = 0.01) and more improvement than NaCl-group in outcomes SSS BCTQ score (from 2.872 to 1.948 in TCA group vs. from 2.815 to 2.529 in NaCl group) and FSS BCTQ score (2.456 to 1.881 in TCA group vs. 2.353 to 2.366 in NaCl group). Mean difference in change score 0.637 (95% CI: 0.320, 0.960; p &lt;0.001)) for SSS BCTQ and mean difference in change score 0.588 (95% CI: 0.232, 0.944; p = 0.02) for FSS BCTQ. Number Needed to Treat to achieve satisfactory partial treatment response or complete</td>
<td>Multiple injections given if patient result was “not satisfactory” Data suggest steroid injections superior to NaCl for short term outcomes.</td>
</tr>
<tr>
<td>Study</td>
<td>Duration</td>
<td>N = 32 with</td>
<td>Methodology</td>
<td>Results</td>
<td>Comments</td>
<td></td>
</tr>
<tr>
<td>------------------------</td>
<td>----------</td>
<td>-------------</td>
<td>-------------</td>
<td>---------</td>
<td>----------</td>
<td></td>
</tr>
<tr>
<td>O'Gradaigh 2000</td>
<td>3.0</td>
<td>suspected CTS and EDS confirmed</td>
<td>Hydrocortisone 25mg or 100mg (A), hexacetonide 20mg (B), plus phase II; Triamcinolone 20mg or Hydrocortisone 100mg (n = 33) vs. Control no injection (n = 20). Follow-up 6 weeks and 6 months.</td>
<td>Results from Phase 1 (25mg/ 100mg/no injection) 66% vs. 65% vs. 5% better or much better (NS between injected groups’ differences). Symptoms improved in Phase 2 in 72% vs. 67% (NS).</td>
<td>“As low dose steroid is as effective, and potentially less toxic, this should be the recommended dose for injection of carpal tunnel syndrome.”</td>
<td></td>
</tr>
<tr>
<td>Girlanda 1993</td>
<td>4.0</td>
<td>clinical and EDS evidence of CTS. Age 36-60 years.</td>
<td>Methylprednisolone acetate 15mg acetate injection locally (n = 9) vs. saline solution same amount as treatment group (n = 8). Study on long-term effects (n = 8). Follow-up at 2 months and 2 years.</td>
<td>Paresthesias significantly improved from baseline in both groups, but more improved in steroid group (p &lt;0.0001 vs. p &lt;0.01); statistical significance of improvements in saline disappeared at 1 month; persisted through 2 months in steroid. 50% of nerves worse within 6 months; 90% within 18 months; 8% of nerves remained improved at 2-year.</td>
<td>“Only a small percentage (8%) of the nerves remained improved at the 2-years follow-up.”</td>
<td></td>
</tr>
<tr>
<td>Wong 2005</td>
<td>9.0</td>
<td>newly diagnosed CTS and NCS confirmed, mean age for single and double injection groups: 46.4 (5.6)/47.3 (9.6).</td>
<td>Single injection group or methylprednisolone 15 mg injection (n = 20) vs. Double-injection group at 8 weeks of steroid or placebo (n = 20). 40 week follow-up.</td>
<td>Global Symptom Score Single vs. Double injections (pre/8/24/ 40 weeks): Single 26.7±10.1/15.2±9.9/15.9±10.6/12.6±9.1 vs. Double 25.6±11.6/11.4±7.6/13.0±9.7/14.1±11.0 (p&gt;0.19) all times. No differences in grip strengths or in NCS other than right hand which was borderline different at baseline (p = 0.08).</td>
<td>“The results suggest that an additional steroid injection confers no added benefit to a single injection in terms of symptoms relief.”</td>
<td></td>
</tr>
<tr>
<td>Atroshi 2013</td>
<td>8.5</td>
<td>(81 female/30 male) with idiopathic CTS not previously treated with Steroid vs. Placebo</td>
<td>Methylprednisolone (n = 37) vs. Placebo (n = 37).</td>
<td>At baseline CTS symptom severity score at 10 weeks improved those who received methylprednisolone vs. placebo (p = 0.003 for 80mg; p = 0.001 for 40mg</td>
<td>“Methylprednisolone injections for CTS have significant benefits in relieving symptoms at 10 weeks and reducing the rate of surgery 1 year after</td>
<td></td>
</tr>
</tbody>
</table>

**One vs. Two Injections**

**Steroid vs. Placebo**

Two studies in one report with the first finding benefits of injection. Second trial found minimal incremental gain for higher dose.

Methods details sparse, especially for long duration components of study. Patients had symptoms over 4 years.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Criteria</th>
<th>Intervention</th>
<th>Outcome Measures</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dammers 1999</td>
<td>RCT</td>
<td>8.0</td>
<td>N = 60 with carpal tunnel symptoms &gt;3 months and NCS confirmed.</td>
<td>Intervention group or methylprednisolone 40mg plus 10mg lidocaine (n = 30) vs. Control group or lidocaine alone (n = 30). Follow-up for one year.</td>
<td>Percentage not needing 2nd treatment (1/3/6/9/12 month): steroid (77/63/57/53/50%) vs. placebo (20/7/7/7/7%), significant but no p-value reported. In open phase, 24 of 28 crossed over from controls and 50% of those had surgery, no p-value reported.</td>
<td>“A single injection with steroids close to the carpal tunnel may result in long term improvement and should be considered before surgical decompression.” Data suggest injection effective and 50% need no treatment for 1 year.</td>
</tr>
<tr>
<td>Ucan 2006</td>
<td>RCT</td>
<td>5.0</td>
<td>N = 57 with CTS diagnosis, mean age for Group A, B, and C: 44.50±7.24, 44.46±8.52, and 45.2±13.19.</td>
<td>Group A or Splinted (S) hands splinted in neutral position with standard cotton polyester splint (n = 23) vs. Group B or single steroid injection (20mg triamcinolone acetate plus 20mg lidocaine) and splinted (SLSI) (n = 23) vs. Group C: Surgery (OCTR) (N = 11). Follow-up for 3 and 6 months.</td>
<td>Boston Questionnaire scores (baseline/3rd month/6th month): splinting 2.66±0.35/1.39±0.37/1.54±0.31 vs. splint plus steroid 2.79±0.63/1.41±0.32/1.96±0.63 vs. CTR 3.09±0.5/1.86±0.6/1.41±0.31 (p = 0.004). Palm-wrist median sensory nerve velocities: splint 27.26±5.3/29.6±7.16/29.56±4.83 vs. splint plus steroid 26.35±4.12/31.57±4.33/28.74±6.19 vs. CTR 23.98±4.28/32.20±4.17/33.15±4.1 (NS between groups). Those completely/almost satisfied 3rd/6th months splinting 69.6%/34.8% vs. splint plus steroid 100%/82.6% vs. CTR 45.5%/90.9%.</td>
<td>“All treatment methods were found to be effective, but despite the complications and the relatively long period to return to work, OCTR was superior to conservative methods in long term.” Baseline differences present. Appears to have targeted lower enrollment for surgery without stating such.</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Description</td>
<td>Results</td>
<td>Conclusion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-------</td>
<td>----------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Glucocorticosteroid vs. Surgery</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hui 2005</td>
<td>8.0</td>
<td>N = 50 (48 female/2 male) with EDS confirmed idiopathic CTS. Follow-up at 6 and 20 weeks.</td>
<td>Mean improvements in global symptoms scale: 24.2±11.0 vs. 8.7±13.0 (p &lt;0.001). Grip strengths were: surgery 23.4±8.2 to 21.8±7.9 vs. injection 24.2±7.0 to 26.6±7.4 (p = 0.009). Sensory nerve conduction velocities: surgery 34.2±7.9 to 42.2±6.0 m/s vs. injection 37.3±8.0 to 40.5±6.3 (p = 0.003).</td>
<td>“Open carpal tunnel release resulted in better symptomatic and neurophysiologic outcome but not grip strength in patients with idiopathic carpal tunnel syndrome over a 20-week period.”</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ly-Pen Arth Rheum 2005; Ly-Pen Med Clin 2005</td>
<td>6.5</td>
<td>N = 123 (163 wrists) with CTS. Mean age 51.9 years. Betamethasone 6.4mg, 2 injections 2 weeks apart (n = 83 wrists) vs. Open Carpal Tunnel Release (n = 80). One year study. Follow-up at 3, 6, and 12 months.</td>
<td>70% improvements in nocturnal paresthesias present (3/6/12 months): injection 86.7/69.9/61.4% vs. surgery 61.3/68.8/ 73.8% (p = 0.001/p = 1.0/p = 0.098).</td>
<td>“Over the short term, local steroid injection is better than surgical decompression for the symptomatic relief of CTS. At 1 year, local steroid injection is as effective as surgical decompression for the symptomatic relief of CTS.”</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ly-Pen 2012</td>
<td>6.0</td>
<td>N = 101 with clinical diagnosis and neuro-physiological confirmation of CTS. Surgical decompression (n = 83 wrists) vs. Local steroid injection (n = 83 wrists). Follow-up of 2 years.</td>
<td>56 underwent surgery, 24 had CTS in both hands. 84% required 2 injections. At 24-months follow-up, 60.2% of wrists in injection group and 68.8% in surgery group achieved 20% response in nocturnal paraesthesias, (p = 0.256). Surgery more effective than injection for self-perceived functional impairment, with mean VAS score of 6.21 (8.81) in injection group vs. 2.02 (7.23) in surgery group, (p = 0.008).</td>
<td>“Our findings suggest that both local steroid injection and surgical decompression are effective treatments in alleviating symptoms in primary CTS at 2-year follow-up.”</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Copyright© 2016 Reed Group, Ltd.
INTRAMUSCULAR INJECTIONS

Intramuscular injections have been used to treat CTS.(854)

Recommendation: Intramuscular Injections for Acute, Subacute, or Chronic CTS
Intramuscular injections are not recommended for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – Not Recommended, Evidence (C) – Chronic CTS
Not Recommended, Insufficient Evidence (I) – Acute, subacute CTS

Levels of Confidence – Moderate

Rationale for Recommendation
Intramuscular injections for CTS are not recommended as they have been found to be inferior to carpal tunnel injections.(854)

Evidence for the Use of Intramuscular Injections for CTS
There is 1 moderate-quality RCT incorporated into this analysis.(854)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: intramuscular injections, carpal tunnel syndrome, CTS, median nerve neuropathy, median nerve disease, entrapment, neuropathy, nerve compression, burning, itching, numbness, tingling, wrist, hand, palm, finger, pain, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 36 articles in PubMed, 722 in Scopus, 3 in CINAHL, 40 in Cochrane Library and 0 in other sources. We considered for inclusion 8 from PubMed, 0 from Scopus, 1 from CINAHL, 2 from Cochrane Library and 0 from other sources. Of the 11 articles considered for inclusion, 3 randomized trials and 1 systematic study met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Özdoğan 1984</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>6.0</td>
<td>N = 37 females: symptoms: burning, pain, tingling, numbness in thumb, index and long fingers and palm. Mean age 45.8 years</td>
<td>Group A: 1.5mg betamethasone disodium phosphate and acetate suspension into carpal tunnel and same volume of placebo (0.5 ml saline) into the deltoid muscle on same side (n = 18) vs. Group B: 1.5mg betamethasone disodium phosphate and acetate suspension into deltoid muscle and same volume of placebo into carpal tunnel (n = 19). Follow-up at 1 week, 1 month, and 10 months after study completion.</td>
<td>Seven patients from carpal injection group and 6 patients from IM injection group returned with symptoms after 1 month and required 2nd shot. One from 1st group and 2 from 2nd group required 3rd shot after 7.3±3.7 months. Response rate 50% in hand injections compared to 15.8% IM.</td>
<td>“Steroid injected at the site of entrapment is effective and suggest superiority to the intramuscular route in the management of ICTS.”</td>
<td>Data suggest intracarpal tunnel injections much more effective.</td>
</tr>
</tbody>
</table>

INSULIN INJECTIONS

Treatment of CTS with carpal tunnel insulin injections has been attempted.(634, 872)

Recommendation: Insulin Injections for Acute, Subacute, or Chronic CTS
There is no recommendation for or against use of insulin injections for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low
Rationale for Recommendation

There is one quality study which included CTS patients that suggests benefit from 7 weekly injections of insulin. (872) A second moderate quality trial found a lack of benefits compared with physiotherapy. (873) The consensus of the Evidence-based Practice Hand, Wrist, and Forearm Panel is that these results require replication. Applicability of these results, even if confirmed, are suggested to be relatively limited to a narrow subset of diabetic patients with CTS who fail to improve with other therapies and either decline surgery or have significant symptoms of focal intracarpal nerve dysfunction after surgery. These injections are invasive, may have adverse effects that also require ascertainment, and are moderate to high cost. There is no recommendation for or against insulin injections for treatment of diabetic patients with CTS.

Evidence for the Use of Insulin Injections for CTS

There are 2 moderate-quality RCT incorporated into this analysis. (872, 873)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Insulin injections and carpal tunnel syndrome, CTS, median nerve neuropathy, median neuropathy, median nerve disease entrapment, neuropathy nerve compression, burning, itching, numbness, tingling, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 6 articles in PubMed, 836 in Scopus, 1 in CINAHL, 39 in Cochrane Library and 0 in other sources. We considered for inclusion 3 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 3 articles considered for inclusion, 2 randomized trials and 1 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intracarpal Tunnel NPH Insulin vs. Saline Injection</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ozkul 2001</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>6.0</td>
<td>N = 43 (gender not specified) with non-insulin-dependent diabetes mellitus (NIDDM) with mild to moderate CTS. Mean age: insulin 47.1±1.3 years placebo 48.4±0.9 years</td>
<td>NPH insulin (0ml-12 U) (n = 22) vs. placebo injected into carpal tunnel weekly for 7 weeks after initial glucocorticoid injection for all (n = 21). Follow-up 23 weeks.</td>
<td>Mean±SD median nerve motor distal latency (MNMDL): decrease 5 weeks in insulin group 4.52±0.12 vs. placebo 4.80±0.03ms (p &lt;0.05) and continued to 23 weeks (p &lt;0.01). Mean±SD median nerve sensory velocity (MNSV): difference more significant insulin group vs placebo over whole study (p &lt;0.01).</td>
<td>“Local insulin injections more significantly decreased MNMDL [median nerve motor distal latency], increase MNSV [median nerve sensory velocity] and reduces GSS [global symptom score] than the placebo in NIDDM patients with CTS.”</td>
<td>All had glucocorticosteroid injection. Suggestive results that need confirmation.</td>
</tr>
<tr>
<td>Ashraf 2009</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>4.0</td>
<td>N = 50 with non-insulin dependent diabetes mellitus; 20 had bilateral involvement, had symptoms and signs of CTS confirmed by standard electro diagnosis.</td>
<td>Group 1 received injection into carpal tunnel (10IU of NPH insulin) (n = N/A) vs. Physiotherapy (2 periods with 10 sessions) (n = N/A). Follow-up period not mentioned.</td>
<td>In both groups decrement of distal motor latency (DML) of median nerves statistically significant. In both groups the increment of sensory nerve conduction velocity was statistically significant. Also, decrement of pain, paresthesia, numbness, weakness/</td>
<td>“In conclusion, in the present study, local insulin injections significantly reduced symptoms as the physiotherapy in NIDDM patients with CTS. But clinical significant difference in compare with physiotherapy was not seen. In summary two local insulin injections had no significant difference with compare to 20.</td>
<td>No differences between groups.</td>
</tr>
</tbody>
</table>
Mean age 51.3.

clumsiness and nocturnal awaking was statistically significant in both groups. But no significant difference between two groups. sessions physiotherapy. Although these findings are promising, further studies with insulin are needed to verify its effectiveness as a treatment for CTS and other degenerative nerve diseases."

BOTULINUM INJECTIONS
Botulinum injections have been used to treat CTS. (874, 875)

Recommendation: Botulinum Injections for Acute, Subacute, or Chronic CTS
Botulinum injections are not recommended for treatment of acute, subacute, or chronic CTS.

Strength of Evidence – Not Recommended, Evidence (C) – Chronic CTS
Not Recommended, Insufficient Evidence (I) – Acute or subacute CTS

Level of Confidence – Low

Rationale for Recommendation
There is one quality study that included CTS patients that does not show clear benefit from botulinum injections, but did show weakness in two patients lasting a few weeks. (874) There are no other quality studies identified for management of other distal upper extremity disorders, including tendinoses. Botulinum injections are invasive, have adverse effects when the effects of the toxin are beyond the site where they were injected that include fatalities, (752, 876) and are costly. They are not recommended for management of CTS.

Evidence for the Use of Botulinum Injections for CTS
There is 1 moderate-quality RCT incorporated into this analysis. (874)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: botulinum toxin, botox or botulinum Injection, carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, burning, itching, numbness, tingling, hand, palm, finger, wrist, and pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 11 articles in PubMed, 201 in Scopus, 2 in CINAHL, and 1 in Cochrane Library. We considered for inclusion 1 from PubMed, 0 from Scopus, CINAHL, and Cochrane Library. Of the 1 article considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breuer 2006</td>
<td>RCT</td>
<td>Sponsored by Elan</td>
<td>7.5</td>
<td>N = 20</td>
<td>2,500 units of</td>
<td>Response rates</td>
<td>&quot;Botulinum toxin B is not dramatically superior to placebo for the relief of CTS symptoms.&quot;</td>
<td>Small sample size. Few screened (20/388) randomized. Suggests not effective.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pharmaceuticals, San</td>
<td></td>
<td>(gender</td>
<td>botulinum toxin B</td>
<td>for botulinum</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Francisco, California. No</td>
<td></td>
<td>not specified</td>
<td>(n = 11) vs.</td>
<td>toxin B and</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>mention of COI.</td>
<td></td>
<td>Placebo</td>
<td>Placebo or</td>
<td>placebo groups:</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>or normal</td>
<td>normal saline</td>
<td>126/143 (88.1%) vs. 117/117 (100%).</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>solution (n = 9).</td>
<td>Follow-up for 13 weeks.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Surgery
CARPAL TUNNEL SURGICAL RELEASE
Surgical consultation may be indicated for CTS patients who:
• Have red flags of a serious nature;
• Fail to respond to non-surgical management including worksite modifications; or
• Have clear clinical and special study evidence of a lesion that has been shown to benefit, in both the short and long term, from surgical intervention.

Surgical considerations depend on the confirmed diagnosis of the presenting hand or wrist complaint. If surgery is a consideration, counseling regarding likely outcomes, risks, and benefits, and especially expectations is important. If there is no clear indication for surgery, referring the patient to a provider experienced in non-operative treatment of CTS may aid in formulating a treatment plan.

Treatment of CTS with surgical release of the carpal flexor retinaculum has been utilized for many decades with surgical case series suggesting significant benefits.(762, 877-903) In the late 1980s, endoscopic releases were reported, gained prominence, utilized various equipment,(894, 895, 904-913) and were initially reported as superior to open releases.(907, 911, 914-918) However, the endoscopic technique reportedly has a higher incidence of injuries to the nerve, particularly in inexperienced surgical hands, as well as higher rates of incomplete surgical releases.(919) A large endoscopically treated case series of 2,402 cases involving 1,698 patients reported an overall success rate of 95% and recurrence rate of 0.5% in experienced hands.(920) More recently, the open technique has been revised towards a minimal incisional technique(921) and continues to be successfully performed with little apparent difference in outcome versus endoscopic releases.(922-925) Currently, there is a trend towards performing these minor surgical releases in uncomplicated patients in clinics as opposed to in hospitals or ambulatory surgical centers that is facilitating return to work on the same day as surgery.

Many adjunctive procedures and modifications of surgical release have been attempted in order to obtain better clinical results. These include neurolysis, epineurectomy, epineurectomy, tenosynovectomy, excision of the carpal ligament, cutaneous nerve sparing, two small open incisions, use of a Knifelight, hypothenar fat pad and other flaps, and concomitant release of the ulnar nerve in Guyon’s canal.(926-941)

Most, but not all surgical studies required patients to have preoperative confirmation with electrodiagnostic studies (EDS), although the EDS criteria are usually not specified. How results compare among those without EDS confirmation is unclear. Risks of surgical decompression include complications of anesthesia (addressed separately in this document), wound infection, complex regional pain syndrome, and damage to the median nerve.(894, 895, 942, 943) Incomplete decompression or recurrence of symptoms can lead to the need for further surgery. Early return to work is the main cost driver regardless of the type of carpal tunnel surgical approach utilized. Early return to work appears more dependent on the attitude of the employer and patient than on the surgical technique,(944) with self-employed patients incurring less lost work time.(944) The durations of lost time have been shown to vary from days to weeks, further suggesting that surgical approaches are not the primary determinants of return-to-work status.

1. Recommendation: Surgical Release for Treatment of Subacute or Chronic CTS

Surgical release is strongly recommended for patients who fail non-operative treatment for subacute or chronic CTS.(911, 914-918, 922-925, 945, 946) It is also recommended for patients who have emergent or urgent indications (e.g., acute compression due to fracture, arthritides, or compartment syndrome with unrelenting symptoms of nerve impairment).

Indications – Failure of non-operative treatment or severe symptoms such as continuous tingling and numbness. Many surgeons will not operate on a patient without a positive EDS. Most patients should have had at least 1 glucocorticosteroid injection with documentation of at least partial or complete relief followed by a return of symptoms. Patients should have an electrodiagnostic study (EDS) consistent with CTS (see Electrodiagnostic Studies). The decision to undergo surgery is typically driven by nocturnal symptoms.(947) Mild CTS with normal EDS exists, but a clinical impression of moderate or severe CTS with normal EDS is very rare and generally indicates a mistaken diagnosis. Positive EDS in asymptomatic individuals is very common, is not CTS, and suggests the need to carefully select patients for EDS and properly interpret the results.
2. **Recommendation: Open or Endoscopic Release for Subacute or Chronic CTS**

   Either open or endoscopic release is moderately recommended for treatment of subacute or chronic CTS. With either open or endoscopic, the effectiveness results from complete division of the flexor retinaculum. The procedure that the surgeon is most comfortable performing is recommended. (922, 923, 925, 945, 946)

   **Indications** – See indications above for Recommendation #1.

   **Strength of Evidence** – Moderately Recommended, Evidence (B)
   **Level of Confidence** – Moderate

3. **Recommendation: Knifelight for Subacute or Chronic CTS**

   The use of a Knifelight is recommended for treatment of subacute or chronic CTS.

   **Indications** – See indications above for Recommendation #1.

   **Strength of Evidence** – Recommended, Evidence (C)
   **Level of Confidence** – Low

4. **Recommendation: Other Adjunctive Procedures or Techniques for Subacute or Chronic CTS**

   While there may be limited indications for the following procedures or techniques, their routine use is not recommended for treatment of subacute or chronic CTS.
   
   **Epineurotomy is moderately not recommended.**
   
   - **Strength of Evidence** – Moderately Not Recommended, Evidence (B)
   - **Level of Confidence** – Moderate

   **Internal neurolysis is strongly not recommended.**
   
   - **Strength of Evidence** – Strongly Not Recommended, Evidence (A)
   - **Level of Confidence** – High

   **Flexor retinacular lengthening is moderately not recommended.**
   
   - **Strength of Evidence** – Moderately Not Recommended, Evidence (B)
   - **Level of Confidence** – Moderate

   **Ulnar bursal preservation is moderately not recommended.**
   
   - **Strength of Evidence** – Moderately Not Recommended, Evidence (B)
   - **Level of Confidence** – Moderate

   The mini palmar incision using the ring finger as a guide does not require any special changes in the location of the incision. Therefore, altering the location of the incision to “superficial nerve-sparing incision” is not recommended.

   - **Strength of Evidence** – Not Recommended, Evidence (C)
   - **Level of Confidence** – Low

   **Flexor tenosynovectomy is not recommended.**

   - **Strength of Evidence** – Not Recommended, Evidence (C)
   - **Level of Confidence** – Moderate

   As discussed above, an incision that is placed too far ulnarly may result in damage to the ulnar nerve or artery; therefore, an ulnar incisional approach is not recommended.

   - **Strength of Evidence** – Not Recommended, Evidence (C)
   - **Level of Confidence** – Low

5. **Recommendation: Biopsy of Abnormal Tenosynovium for Subacute or Chronic CTS**

   Biopsy of abnormal tenosynovium is recommended for treatment of subacute or chronic CTS.
Indications – Abnormal appearing tenosynovium, including potential amyloidosis, infectious agents, or evidence for inflammatory conditions.

Strength of Evidence – Recommended, Insufficient Evidence (I)

Level of Confidence – Moderate

Rationale for Recommendations
Evidence that surgery is effective is strong. Six quality studies have compared carpal tunnel release with other interventions. Three quality studies document superiority compared with splinting.(763, 777, 778) Two of three studies suggest superiority of surgical release compared with injection(777, 851-853) over longer timeframes mostly of 1 year. One study suggested superiority compared with physical therapy.(641) These appear to indicate a slight superiority of surgery to injection over 1 year and a modestly stronger benefit compared with nocturnal splinting. Longer-term outcomes are believed to further favor surgery.

Seventeen quality trials have compared open versus endoscopic techniques.(907, 911, 914-918, 922-925, 945, 946, 948-952) Six of 11 studies reported since 2000 have failed to demonstrate better outcomes with endoscopic releases,(922, 923, 925, 945, 946, 948-952) which appears to be due to the successful use of minimal incisional techniques that utilize incisions as small as 2cm. These small incisions appear to have removed the primary advantage of endoscopic releases. Quality evidence of superiority of endoscopic versus minimal incisional releases is now lacking and one study has reported no differences at 5-year follow-up, also importantly documenting no differences in reoperation rates.(945) Differences in recovery time between the endoscopic or minimally invasive techniques reported mostly in the 1990s appear to have largely or completely disappeared in the 2000s with 4,(922, 923, 925, 945, 946) of 6 studies(917, 918) showing a lack of superiority of the endoscopic release.(922-925, 945, 946) However, it is the surgeon’s experience and comfort that are determining factors in the selection of the procedure performed.

Overall, the available evidence suggests either the open or endoscopic procedures are successful surgical procedures. Thus, the Evidence-based Practice Hand, Wrist, and Forearm Panel agreed regarding the overall recommendation for surgery. The mini-procedures continue to improve outcomes,(921) while most early studies compared endoscopic to the traditional open procedure. Outcome measures vary with each study making a direct comparison difficult. Studies have shown that with well-motivated individuals and a mini-palm technique, return to modified work the next day is possible.(398)

Recently, a Knifelight has been utilized for carpal tunnel releases.(928, 932) This technique involves use of an instrument through a small palmar incision to perform a blinded division of the flexor retinaculum. While there is sufficient quality evidence to document efficacy and recommend the procedure,(928, 932) further studies are needed comparing the Knifelight with a standardized, minimal incisional technique and using larger sample sizes.

There have been many alterations on standard operative techniques(953) and/or adjunct surgical procedures performed to attempt to derive superior outcomes for patients who have been subjected to quality studies. Without exception, none of the following were found beneficial – epineurotomy,(929, 930, 935, 954, 955) neurolysis,(937, 938) flexor tenosynovectomy,(939) flexor retinacular lengthening,(956) nerve sparing incisions,(940) double-limited incisions,(941) ulnar incisions,(957) and ulnar bursal preservation.(931) Evidence indicates that even in the presence of synovial hypertrophy and histological changes, tenosynovectomy has not been shown to be beneficial in a moderate-quality study.(939) However, biopsy of abnormal tissue is indicated for diagnostic and therapeutic purposes. Examples of potential findings to be sought include amyloidosis, infectious agents, and evidence for inflammatory conditions.

The primary cost driver for CTS claims is lost work time.(778) CTS is not different from other MSDs, with reportedly worse outcomes and greater delays in return to work among patients receiving workers’ compensation.(914, 925) In quality studies, lost time ranged from 12 days for open releases in the Netherlands(778) to 88 days for endoscopically treated patients in Sweden,(922) with most trials
reporting these data between 12 and 40 days. There is no clear pattern by country or procedure, other than a propensity for greater lost time in the 1990s in the open release groups compared with the endoscopic releases, prior to the dissemination of limited incisional techniques.

**Evidence for the Use of Carpal Tunnel Surgical Release**

There are 7 high-quality (763, 851, 931, 937, 938, 955, 956) and 36 moderate-quality (one with two reports) RCTs and crossover trials incorporated into this analysis. There are 13 low-quality RCTs (407, 846, 913, 930, 961-969) in Appendix 2.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: carpal tunnel surgical release, Knifelight, open release, endoscopic, epineurotomy, neurolysis, flexor retinacular, ulnar bursal preservation, mini palmer incision, flexor tenosynovectomy, biopsy of abnormal tenosynovium and carpal tunnel syndrome, median nerve neuropathy, median neuropathy, median nerve disease entrapment, neuropathy nerve compression, burning, itching, numbness, tingling, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective studies, prospective studies, epidemiological studies, epidemiological research, and Nonexperimental Studies. We found and reviewed 179 articles in PubMed, 84 in Scopus, 17 in CINAHL, 45 in Cochrane Library and 0 in other sources. We considered for inclusion 56 articles from PubMed, 2 from Scopus, 2 from CINAHL, 1 from Cochrane Library and 3 from other sources. Of the 64 articles considered for inclusion, 51 randomized trials and 12 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gerritsen 2002 RCT Sponsored by a grant from the Health Care Insurance Council of the Netherlands. No mention of COI.</td>
<td>8.5</td>
<td>N = 176 (143 female/33 male) EDS confirmed. Mean age 49 years.</td>
<td>Open release (n = 87) vs Splinting for 12 months (n = 89). Follow-up at 1, 3, 6, 12 and 18 months.</td>
<td>Overall success rates statistically superior for all 5 measurements other than 1st month (1/3/6/12/18 months): 29 vs. 42% (p = 0.07)/80 vs. 54% (p = 0.001)/94 vs. 68% (p &lt; 0.001)/92 vs. 72% (p = 0.002) /90 vs. 75% (p = 0.02). Nights awakening with symptoms and paresthesias not significantly different at 12 or 18 months. Five (5.7%) in surgery group had wound infection; CRPS/RSD in one. Median-ulnar latency differences borderline favored splinting (baseline: 1.7 vs. 1.8 months; 12 months: 1.1 vs. 0.7 months), as did other measures.</td>
<td>“Treatment with open carpal tunnel release surgery resulted in better outcomes than treatment with wrist splinting for patients with CTS.”</td>
<td>Duration of symptoms was somewhat worse in splinting group (median 52 vs. 40 weeks, NS). Both treatment arms document substantial improvement, which may reflect a good natural history.</td>
</tr>
<tr>
<td>Jarvik 2009 RCT Sponsored by the Intramural Research Program of the NIH Clinical Center. No COI.</td>
<td>7.0</td>
<td>N = 116 (62 female/54 male) considering surgery for diagnosed CTS. Mean age 50.7 years.</td>
<td>Surgery Group: Open surgery or Endoscopic surgery depending on surgeon’s preference. (n = 57) vs. non- Surgical therapy group: 6 visits with physical therapy and prescribed NSAIDS. (n = 59). Follow-up 3, 6, 9 and 12 months.</td>
<td>Primary outcome Carpal Tunnel Syndrome Assessment Questionnaire (CTSAQ). Surgical group significantly lower CTSAQ function score vs. non-surgical group at 6 months: 1.91 vs. 2.44 (p = 0.006) and at 12 months: 1.74 vs. 2.17 (p = 0.0081). Secondary outcome of CTSAQ symptoms also significantly lower in surgery vs. non-surgery at 6 months: 2.02 vs. 2.42 (p = 0.018) and 12 months: 1.74 vs. 2.07 (p = 0.036).</td>
<td>“Overall, these data indicate that, in patients with carpal tunnel syndrome without denervation, surgery modestly improves hand function and symptoms by 3 months compared with a multimodality non- surgical treatment regimen. And this benefit is sustained through 1 year.”</td>
<td>At 12 months, surgical group was statistically significant for improved symptoms and function.</td>
</tr>
<tr>
<td>Korthals-de Bos 2006 RCT Sponsored by grant from Health Care Insurance Council of the Netherlands. No COI.</td>
<td>4.0</td>
<td>N = 13 EDS confirmed. Mean age not provided.</td>
<td>Open release: Incision size not specified. Numerous specialists performed (n = 73) vs. Nocturnal splinting plus daytime “if they wished to.” Follow-up 3, 6, 12 months.</td>
<td>Success rates higher at 12 months for surgery group, 92% vs. 72%, difference is 20% (8-31 95% CI). Night awakening due to complaints not different (3.6±2.9 vs. 2.9±3.0). Severity of main complaint higher in surgery group (6.4±2.7 vs. 5.1±3.1). Mean aggregate costs 2,126€ surgery vs. 2,111€ splint. Absenteeism comparable (50 vs. 52 days).</td>
<td>“In the Netherlands, surgery is more cost-effective compared with splinting, and recommended as the preferred method of treatment for patients with CTS.”</td>
<td>Population-based study with likely relatively suboptimal control over treatments. Small sample size. Applicability of cost data to US is questionable.</td>
</tr>
</tbody>
</table>
## Carpal Tunnel Release vs. Injections

<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Comparison</th>
<th>Median/ Mean Age</th>
<th>Intervention</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hui 2005</td>
<td>8.0</td>
<td>N = 50 (48 female/2 male) EDS confirmed. Mean age 49.5 years.</td>
<td>Injection Group- Methypred-nisolone acetate 15mg (n = 25) vs. Open carpal tunnel release (n = 25). Follow-up at 6 and 20 weeks.</td>
<td>Mean improvements in the global symptoms scale 24.2±8.3 vs. 8.7±3.8 (p &lt; 0.001). Grip strengths: surgery 23.4 ±8.2 to 21.8±7.9 vs. injection 24.1±7.0 vs. 26.6±7.4 (p = 0.009). Sensory nerve conduction velocities: surgery 34.2±7.9 to 42.2±8.0 m/s vs. injection 37.3±8.0 to 40.5±6.3 (p = 0.003).</td>
<td>“Open carpal tunnel release resulted in between symptomatic and neurophysiologic outcome but not grip strength in patients with idiopathic carpal tunnel syndrome over a 20-week period.”</td>
</tr>
</tbody>
</table>

Ly-Pen Arth Rheum 2005; Ly-Pen Med Clin 2005 | 6.5 | N = 123 (163 wrists) with CTS. Mean age 51.9 years. | Bethamethasone 6.4mg, 2 injections 2 weeks apart (n = 83 wrists) vs. Open Carpal Tunnel Release (n = 80). 1 year study. Follow-up 3, 6, and 12 months. | 70% improvements in nocturnal paraesthesias present (3/6/12 months): injection 86.7/69.9/61.4% vs. surgery 61.3/68.8/73.8% (p = 0.001/p = 1.0/p = 0.098). | “Over the short term, local steroid injection is better than surgical decompression for the symptomatic relief of CTS. At 1 year, local steroid injection is as effective as surgical decompression for the symptomatic relief of CTS.” |

Ucan 2006 | 5.0 | N = 57 (57 hands) with mild to moderate idiopathic carpal tunnel syndrome. Mean age 44.6 years. | Group A: splinted for 3 months (n = 23 Hands) vs. Group B: Single steroid injection (20mg triamcinolone acetate with 20mg lidocaine) and splinted for 3 months (n = 23 Hands) vs. Group C: surgery (n = 11 Hands). Follow-up assessments 3 and 6 months. | Boston Questionnaire scores (baseline/ 3rd month/6th month): splinting 2.66±0.35/1.39±0.37/1.54±0.31 vs. splint plus steroid 2.79±0.63/1.41±0.32/1.96±0.63 vs. CTR 3.09±0.5/1.86±0.6/1.41±0.31 (p = 0.004 at 6 months). Palm-wrist median sensory nerve velocities: splint 27.26±5.3/29.6±7.16/29.56±4.83 vs. splint plus steroid 26.35±4.12/31.57±4.33/28.74±6.19 vs. CTR 23.98±4.28/32.20±4.17/33.15±4.1 (NS). Completely satisfied/ almost satisfied (3rd/6th months): splinting 69.6%/34.8% vs. splint plus steroid 100%/82.6% vs. CTR 45.5%/90.9%. | “All treatment methods were found to be effective, but despite the complications and the relatively long period to return to work, OCTR was superior to conservative methods in long term.” |

## Endoscopic vs. Open Release

<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Comparison</th>
<th>Median/ Mean Age</th>
<th>Intervention</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saw 2003</td>
<td>7.5</td>
<td>N = 123 with carpal tunnel syndrome. Mean age 51.9 years.</td>
<td>Open Carpal Tunnel Release Group: Open incision 2cm (n = 76) vs. 1-portal endoscopic release (n = 74). Follow-up at 1, 3, 6 and 12 weeks.</td>
<td>Anterior carpal tenderness not significantly different 22±7 vs. 24±6 (p = 0.18). Grip strength was also not different, but favored endoscopic (p = 0.21). Endoscopic group returned to work average 8 days (95% CI 2-13, (p = 0.005)) sooner than open. Lost time offset</td>
<td>“On the basis of these findings, we recommend that endoscopic carpal tunnel release should be considered in the employed as a cost-effective procedure, but perhaps not in the long term.”</td>
</tr>
</tbody>
</table>
Atroshi 2006  
**RCT**  
Sponsored by research grants from Skane county council’s research and development foundation, Kristianstad University, and Swedish Society of Medicine. No COI.  
| 7.0 | N = 128 (88 female/40 male) with idiopathic CTS. Mean age 44 years | Open Surgery Group-4cm open (n = 65) vs. 2-portal endoscopic release-1cm endoscopic (n = 63). Follow-up at 3 and 6 weeks and 3 and 12 months. | Post-operative pain scores (3 weeks/6 weeks/ 3 months/12 months): open 60.5±23/51.3±23/36.2±20/13.9±22 vs. endoscopic 52.1±23/43.3±23/23.5±26/8.7±21 (p = 0.028, p = 0.03, p <0.001, p = 0.13 respectively). Lost time median 28 days in both groups (range 17-44). | increased costs of endoscopic surgery, resulting in net savings of 438€ ($661.63 USD 2009) per patient. | general population as a whole. |

Atroshi 2009  
**RCT**  
Sponsored by research grants from Skane County Council’s research and development foundation, Kristianstad University, and The Swedish Society of Medicine. No mention of COI.  
| 7.0 | Same as above Atroshi 2006 | Same as above Atroshi 2006 | Symptom severity scores at 5 years were endoscopic 1.45±0.7 vs. open 1.42±0.7 (NS). 52/61 open vs. 53/63 endoscopic had “no pain” (NS). No differences in functional status scores, although both improved from pre-operative status (p <0.001). In 1st year, 1 open and 2 endoscopic required repeat surgery; between Years 1 and 5, 2 open and 1 endoscopic required repeat surgery. | “In carpal tunnel syndrome, endoscopic surgery was associated with less postoperative pain than open surgery, but the small size of the benefit and similarity in other outcomes make its cost effectiveness uncertain.” | Minimal advantage to endoscopic of less pain, but not earlier return to work. |

Brown 1993  
**RCT**  
No sponsorship or COI.  
| 6.5 | N = 145 (169 hands) with CTS. Mean age 56 years. | Open Carpal Tunnel Release: Open incisions 3.5-4.5cm (n = 75, 85 hands) vs. 2-portal endoscopic release-endoscopic incisions 2cm and 1.5cm (n = 76, 84 Hands). Follow-up at 21, 42, 84 days. | Symptoms relieved in 98-99% among each group. Open group more likely to have incisional tenderness (61% vs. 36%). Return-to-work occurred earlier for endoscopic group (p <0.05). | “Preliminary analysis suggests that functional outcomes are achieved more quickly when the endoscopic method is used. However, the greater rate of complications indicates that intraoperative safety must be improved before endoscopic carpal-tunnel release is performed on a widespread basis.” | Suggested endoscopic superior. |

Ferdinand 2002  
**Crossover Trial**  
| 6.5 | N = 25 (50 hands; 20 female/5 male) with | Open carpal tunnel release (n = 25) vs 1-portal endoscopic release (n = 25). | Data presented graphically. Persisting symptoms in 1 (4%) of open vs. 0% endoscopic. Persisting pain in 1 in each group. No “In comparison with open release, single-portal endoscopic carpal tunnel release has a similar | “The improvements in symptoms of CTS and hand-related disability 5 years after open and 2-portal endoscopic carpal tunnel release were equivalent.” | Very high response rate for 5-year study (only missing 2 who died). Suggests no long term differences. Same rates of palmar pain for both groups. No differences in reoperation rates. |

---

**Copyright © 2016 Reed Group, Ltd.**
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N &amp; Description</th>
<th>Incision Sizes</th>
<th>Follow-up Period</th>
<th>Results</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trumble 2002</td>
<td>RCT</td>
<td>N = 147 (192 hands) with idiopathic CTS. Mean age 56 years.</td>
<td>Open incision 3-4 cm (n = 72, 95 hands) vs. 1-portal endoscopic release (n = 75, 97 hands). Follow-up assessments made at 2, 4, 8, 12, 26, and 52 weeks.</td>
<td>Symptom severity scores different for weeks 2; 3.1 vs. 2.3 (p &lt; 0.01), 4; 3.0 vs. 2.0 (p &lt; 0.01), 8; 2.7 vs. 1.9 (p &lt; 0.01), and 12; 2.5 vs. 1.8 (p &lt; 0.01) among open group vs. endoscopic group. Open group also showed significant increase in functional status score vs. endoscopic group at week 2; 3.0 vs. 2.2 (p&lt;0.01), 4; 2.6 vs. 1.9 (p&lt;0.01), 8; 2.5 vs. 1.9 (p&lt;0.01), and 12; 2.4 vs. 1.7 (p &lt;0.01). Median time to return to work 38 vs. 18 days, (p = 0.0086), favoring endoscopic group.</td>
<td>“Good clinical outcomes and patient satisfaction are achieved more quickly when the endoscopic method of carpal tunnel release is used. Single portal endoscopic surgery is a safe and effective method of treatment carpal tunnel syndrome.”</td>
<td></td>
</tr>
<tr>
<td>Wong 2003</td>
<td>Crossover</td>
<td>N = 30 (60 hands) with bilateral idiopathic CTS. Mean age 47 years.</td>
<td>Open Group: using Strickland instrumentation. 1.5cm open incision (n = 15, 30 hands) vs. 2-portal endoscopic release (n = 15, 30 hands). Follow-up 2, 4, 8, 16 weeks, 6 and 12 months.</td>
<td>At 1 year, 17 (57%) of endoscopic vs. 19 (63%) of limited open had complete resolution (p = 0.65). Trend toward increased strength in open group (NS). Pain scores lower in limited open group 2 weeks: 2.5 vs. 3.3 (p = 0.004) and 4 weeks: 1.5 vs. 2.5 (p = 0.008).</td>
<td>“The results showed that the outcome was similar at follow-up of one year using both techniques. However, the LOCTR group had significantly less tenderness of the scar at the second and fourth postoperative week. There was also less thenar and hypothenar (pillar) pain after LOCTR.”</td>
<td></td>
</tr>
<tr>
<td>Erdmann 1994</td>
<td>RCT</td>
<td>N = 105 with CTS. Mean age 53.4 years.</td>
<td>Open carpal tunnel release (n = 52) vs. 2-portal endoscopic release (n = 53). Incision sizes not specified.</td>
<td>Symptoms relieved in 1.1 vs. 1.75 days. Return to work in 14 vs. 39 days (p&lt;0.005) for the endoscopic group vs. the open group. Grip strength returned to preoperative</td>
<td>“This trial illustrates that endoscopic carpal tunnel release has distinct advantages over open surgery, in a select group”</td>
<td></td>
</tr>
</tbody>
</table>

Data suggest the long-term outcomes were identical, although the benefits were short-term for the endoscopic technique.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Gender</th>
<th>Age (Mean)</th>
<th>Inclusion Criteria</th>
<th>Intervention</th>
<th>Follow-up</th>
<th>Outcome Measures</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>MacDermid 2003</td>
<td>RCT</td>
<td>6.0</td>
<td>N = 123</td>
<td>53.6 years</td>
<td>Open carpal tunnel syndrome (n = 32) vs. 2-portal endoscopic release (n = 91). Incision sizes not specified. Follow-up assessments at 1, 6 and 12 weeks.</td>
<td>McGill Pain Questionnaire scores favored endoscopic release, e.g., Week 1: 13 vs. 28 and Week 6: 12 vs. 22, both (p &lt; 0.05). Symptom Severity Scale scores not significantly different. Grip strengths at 1 and 6 weeks favored endoscopic release (e.g., week 1, 11 vs. 15kg, (p &lt; 0.05)).</td>
<td>No substantive difference in benefit was shown for these 2 methods of carpal tunnel release.</td>
<td>The data indicate less pain and better grip strength at 1 to 6 weeks in the endoscopically treated group.</td>
<td></td>
</tr>
<tr>
<td>Sennwald 1995</td>
<td>RCT</td>
<td>5.5</td>
<td>N = 47</td>
<td>52.6 years</td>
<td>Open carpal tunnel release (n = 22) vs. 1-portal endoscopic release- Endoscopic incision 2cm (n = 25). Follow-up at 4, 8 and 12 weeks.</td>
<td>Grip strength recovery significant at 4 weeks (p = 0.005), 8 weeks (p = 0.003) and 12 weeks (p = 0.0002) in favor of endoscopic group compared to open group. Endoscopic group could use operated hand normally after 24 days vs. 42 days after open procedure (p &lt; 0.001).</td>
<td>The study is strongly in favor of endoscopic release. However, this technique does not allow any analysis of the pathology or structure to be treated.</td>
<td>Baseline mean grip strength approximately 26 vs. 32 (p = 0.29). Appears to have contributed to post-operative differences.</td>
<td></td>
</tr>
<tr>
<td>Ejiri 2012</td>
<td>RCT</td>
<td>5.5</td>
<td>N = 79</td>
<td>51.2 years</td>
<td>Endoscopic carpal tunnel release (ECTR group) (n = 40, 51 hands) vs. Open carpal tunnel release (OCTR) (n = 39, 50 hands). Follow-up assessments at week 4 and 12.</td>
<td>At week 12, rate of improved cases higher in OCTR group vs. ECTR group (p = 0.08), however not significant. No significant differences between groups for improvement in ADL impairment. At week 4, mean improvement in grip strength significantly higher in ECTR group vs. OCTR: -4.6 vs. -8.1 (p = 0.04). But not significant at 12 weeks: -1.2 vs. -3.6 (p = 0.27).</td>
<td>These results suggest that while no difference exists between ECTR and small incision methods in terms of improved subjective symptoms, sensation, or electrophysiological findings, recovery of muscle strength is superior with ECTR.</td>
<td>At 4 weeks, ECTR was significantly better than OCTR for muscle strength, but ECTR may increase the risk of transient nerve dysfunction which resolved at 6 months.</td>
<td></td>
</tr>
<tr>
<td>Larsen 2013</td>
<td>RCT</td>
<td>5.5</td>
<td>N = 90</td>
<td>52.6 years</td>
<td>Classic incision group 7cm curved incision (n = 30) vs. short incision group: incision 3cm in mid-palm (n = 30). Endoscopic group using Linvatec system (n = 30). Follow-up at 1, 2, 3, 6, 12, 24 weeks.</td>
<td>No significant difference between groups for post-op pain at any time point (p &gt;0.05). No significant difference for disappearance of paresthesia between treatment groups (p &gt;0.05). Tendency for earlier return of grip strength (significant at weeks 2 and 3 only (p &gt;0.05)), as well as ROM (significant at weeks 1 and 3) in endoscopic groups vs. other two groups.</td>
<td>These results are in accordance with the findings in the literature: faster rehabilitation and earlier return to work after ECTR... (Endoscopic Carpal Tunnel Release), few complications but a risk of nerve branch neuropathy with transient neurological problems.</td>
<td>At 24 weeks, the endoscopic group had quicker return to work and faster rehabilitation.</td>
<td></td>
</tr>
<tr>
<td>Study Reference</td>
<td>Design</td>
<td>Sponsored by</td>
<td>Patient Characteristics</td>
<td>Procedure Details</td>
<td>Outcome Measures</td>
<td>Results</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-----------------</td>
<td>--------</td>
<td>-------------</td>
<td>------------------------</td>
<td>----------------</td>
<td>----------------</td>
<td>---------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dumontier 1995</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>N = 96 with idiopathic CTS; Mean age 52.3 years.</td>
<td>Open carpal tunnel release group: Open incisions 3-4cm (n = 40) vs. 2-portal endoscopic release (n = 56). Follow-up assessments made at 2 weeks, 1, 3, 6 months.</td>
<td>Loss of grip strength conventional group vs. endoscopic group (mean±SD): 2 W-pre-op: -15.02±10.27/-13.84±9.50 (p = 0.67); 1 M-pre-op: -12.80±9.84/-6.25±6.81 (p &lt; 0.01); 3 M-pre-op: -8.26±6.37/-3.66±6.84 (p = 0.02).</td>
<td>“No statistically significant differences were found regarding pain, disappearing of paresthesiae or time to return to work. However, better recovery of grip strength was observed in the endoscopic group at 1 and 3 months.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jacobsen 1996</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>N = 29 EDS confirmed (32 hands) with idiopathic CTS. Mean age 46 years.</td>
<td>Open carpal tunnel release group (n = 16 Hands) vs. 2-portal endoscopic release (n = 16 hands). Incision sizes not specified. Follow-up at 2 and 6 weeks and 6 months.</td>
<td>Sick length average 17 days (0-31) in endoscopic group vs. 19 days (0-42 days) in open group. No significant difference between groups for average sick day length (p &gt; 0.05). At final follow-up, 8 in endoscopic group returned to normal vs. 9 in open group (p &gt; 0.05).</td>
<td>No differences in surgical results were found, but three patients in the endoscopic group suffered transient numbness on the radial side of the ring finger.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kang 2013</td>
<td>RCT</td>
<td>No mention of sponsorship. No COI.</td>
<td>N = 59 with bilateral CTS. Each hand randomly assigned to different surgery. Mean age 55 years.</td>
<td>Endoscopic Group: carpal tunnel release surgery performed with Agee technique (n = 59 hands) vs. Mini-Open Group: release performed with small (1.5cm) incision. (n = 59 hands). Follow-up at 3 months post-op.</td>
<td>Boston Carpal Tunnel Questionnaire symptom (BCTQ-S) and function (BCTQ-F) score main outcome. No significant differences between endoscopic vs. mini-open at 3 months for BCTQ-S: 1.5 vs. 1.4 (p = 0.774) or for BCTQ-F: 1.5 vs. 1.7 (p = 0.832). No significant difference in mean DASH (Disabilities of the Arm, Shoulder and Hand) score (p = 0.978).</td>
<td>“Endoscopic and mini-incision open carpal tunnel releases seem to have comparable early subjective outcomes after carpal tunnel release has been performed in patients who had idiopathic carpal tunnel syndrome.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Agee 1992</td>
<td>RCT</td>
<td>Sponsored in part by the 3M Orthopedic Products Division, St. Paul, Minn. No mention of COI.</td>
<td>N = 122 (147 hands) with CTS. Mean age not provided.</td>
<td>Open carpal tunnel release-Control Group (65 hands) vs. 1-portal endoscopic release-Endoscopic incision 2cm (n = 82 hands). Follow-up at weeks 1, 2, 3, 6, 9, 13, and 26.</td>
<td>Median return to work 25 days vs. 46.5, (p &lt;0.01). Stratified analyses 71 vs. 16.5 days for workers’ comp vs. non-WC treated with endoscopic technique vs. 78 vs. 45.5 days in open group (WC vs. non-WC). Less immediate postoperative scar tenderness.</td>
<td>“Improvement in most of the variables measured translated into earlier return to work and to ADL.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jugovac 2002</td>
<td>RCT</td>
<td>Sponsored by Croatian Ministry of Science and Technology grant No. 0082076 to Dr.</td>
<td>N = 72 with NCS finding of CTS. Mean age 53.4 years.</td>
<td>Open carpal tunnel release group (n = 36) vs. mini-incision group-using an operating microscope (n = 36). 3 month follow-up.</td>
<td>Symptomatic relief open (31/36 complete relief) vs mini (31/36) (NS). Hand function return to daily activities in 5 days with limited incision vs. 10 days open (p = 0.001). RTW 15 vs. 30 days (p = 0.001).</td>
<td>“Limited palmar incision CTR is as effective and safe as traditional CTR technique, but with better postoperative recovery and cosmetic results.”</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Open vs. Mini Incision**

Possibly 2:1 assignment, not noted. Variable follow-ups with 45.3% dropout at 3 months. Higher risks in endoscopic group. Sparse methodology. Comparable outcome efficacy at 3 months, but patient preference towards endoscopic procedure. Suggests endoscopic superior to open. Some baseline differences. Follow-up timing unclear.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Funding</th>
<th>COI</th>
<th>Participant characteristics</th>
<th>Intervention</th>
<th>Outcome measures</th>
<th>Results/Findings</th>
<th>Conclusions</th>
</tr>
</thead>
</table>
| Aslani 2012  
RCT  
No mention of sponsorship or COI. | 4.0 | N = 105 who qualified for carpal tunnel release surgery. Mean age 54.2 years. | Open surgery group (n = 36) vs. Endoscopic surgery group (n = 32) vs. Mini Palmer incision group (n = 28). Follow-up at 2 weeks, 4 weeks and 4 months. | Endoscopic (2 wrists showed weakness at 4 months) and Mini Palmer incision (0 wrists weakness) groups showed significant improvement in weakness vs. open surgery (4 wrists showed weakness) (p <0.05). No other significant differences for other variables (p >0.05). 0 participants expressed pain in the open group at final follow-up and 4 participants expressed pain in both endoscopic and mini-palmer groups. | “Satisfactory results with all three surgery techniques of open, mini-incision or endoscopic and has a low chance of complications. Endoscopic treatment and mid-palmar mini incision have less pain and greater satisfaction among patients in the first weeks, however, overall results are the same and satisfactory in all three groups after 4 months.” | Cross-sectional study shows early patient satisfaction with endoscopic and mini-techniques, but at 4 months comparable satisfaction between all groups. |
| Tarallo 2014  
RCT  
Sponsored by National Institutes of Health (NIH), Wellcome Trust, Howard Hughes Medical Institute (HHMI) and other(s). No COI. | 4.0 | N = 120 with CTS with moderate-to-severe symptoms. Mean age 64 years. | Group A: carpal tunnel release by traditional open carpal tunnel release (TOCTR) (n = 60) vs. Group B: carpal tunnel release by minimal-access carpal tunnel release (MACTR) (n = 60) Follow-up at 7 days, 6 and 12 months. | At final follow-up mean static 2-point discrimination score difference not significant between Group A and B; 4.3 mm vs. 4.7mm (p >0.05). At final follow-up, 2 patients (3.6%) in Group A had evidence of recurrent disease vs. 1 (1.8%) in Group B (p <0.01). In each subsection of STSS questionnaire, Group B showed significantly better results than Group A at both 6 month follow-up 1.4 vs. 2.3 (p <0.001) and 12 month follow-up; 1.1 vs. 1.5 (p<0.001). | “In our opinion, median nerve release is strongly advocated by MACTR as a safe, easily reproducible, low-grade learning curve, low time and a low-cost surgery and it can be performed with standard surgical equipment. In our perspective randomised study, MACTR showed statistically significant improvement compared to TOCTR.” | MACTR group was significantly better than TOCTR group at 6 and 12 months. |
| Zyluk 2006  
RCT  
No mention of sponsorship or COI. | 6.5 | N = 79 (50 female/15 male; 82 hands) EDS confirmed CTS. Mean age 48. | 1 limited incision group- Single (2cm) (n = 39, 44 hands) vs. 2 limited open incisions group 1 and 2cm incisions (n = 40, 40 hands). Follow-up at 1, 3, 6, 12 months. | Functional scores not different. Total grip strength (kg) Method 1/Method 2: Pre-op: 16.6/18.1; at 1 month: 16.1/14.9; at 3 months: 20.3/18.9; at 12 months: 24.2/24.1. No significant differences between groups for grip strength (p >0.05). | “We found that the single incision method offers better results in respect of grip and pinch strengths: less weakness at 1 month after surgery and a faster improvement relative to pre-operative values which is statistically significant.” | Minor advantage to one small incision. |
| Cnkovic 2012  
RCT  
No mention of COI. | 9.0 | N = 50 (33 female/17 male) with Epineurotomy Group: Open field surgical release followed by | At 90 days, mean nerve volume increase somewhat higher in epineurotomy group vs. no | “In conclusion, in line with other reports, the results suggest that even failure to provide superiority for epineurotomy after carpal |

### Epineurotomy

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Funding</th>
<th>COI</th>
<th>Participant characteristics</th>
<th>Intervention</th>
<th>Outcome measures</th>
<th>Results/Findings</th>
<th>Conclusions</th>
</tr>
</thead>
</table>
| Marin F. Stanèiæ.  
No mention of COI. | 6.9 | N = 105 who qualified for carpal tunnel release surgery. Mean age 54.2 years. | Open surgery group (n = 36) vs. Endoscopic surgery group (n = 32) vs. Mini Palmer incision group (n = 28). Follow-up at 2 weeks, 4 weeks and 4 months. | Endoscopic (2 wrists showed weakness at 4 months) and Mini Palmer incision (0 wrists weakness) groups showed significant improvement in weakness vs. open surgery (4 wrists showed weakness) (p <0.05). No other significant differences for other variables (p >0.05). 0 participants expressed pain in the open group at final follow-up and 4 participants expressed pain in both endoscopic and mini-palmer groups. | “Satisfactory results with all three surgery techniques of open, mini-incision or endoscopic and has a low chance of complications. Endoscopic treatment and mid-palmar mini incision have less pain and greater satisfaction among patients in the first weeks, however, overall results are the same and satisfactory in all three groups after 4 months.” | Cross-sectional study shows early patient satisfaction with endoscopic and mini-techniques, but at 4 months comparable satisfaction between all groups. |

### 1 vs. 2 Limited Open Incisions

- **Ongoing Study:** |- | | | | | | | | |
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Sample Size</th>
<th>Eligibility Criteria</th>
<th>Intervention</th>
<th>Follow-up</th>
<th>Outcome Measures</th>
<th>Findings</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leinberry 1997</td>
<td>1997</td>
<td>RCT</td>
<td>7.0</td>
<td>N = 44 (gender not specified) EDS confirmed (50 hands) with CTS. Mean age 64.8 years.</td>
<td>Group 1: Release of transverse carpal ligament. No epineurotomy (n = 22, 25 hands) vs. Group 2: carpal tunnel release and adjuvant epineurotomy of median nerve (n = 22, 25 hands). Follow-up 1 and 6 weeks; 6 and 12 months.</td>
<td>At 12-months, 60% of non-epineurotomy group vs. 56% of epineurotomy group asymptomatic (p &gt;0.05). Two-point discrimination, grip strength and sensory nerve latencies all not significantly different.</td>
<td>This suggests that epineurotomy of the median nerve offers no benefit compared with sectioning of the transverse carpal ligament alone.</td>
<td>Patient blinding unclear, but seems probable.</td>
<td></td>
</tr>
<tr>
<td>Blair 1996</td>
<td>1996</td>
<td>RCT</td>
<td>6.0</td>
<td>N = 86 EDS confirmed (117 hands) with CTS. Mean age 48.7 years.</td>
<td>Open release group- 4cm incision (n = 48) vs. carpal tunnel release with epineurotomy. 4cm incision (n = 27). Follow-up for minimum of 24 months.</td>
<td>Synovial hypertrophy graded as marked or moderate in 18.8% of epineurotomy group vs. 33.3% of non-epineurotomy group. Non-significant trends in favor of epineurotomy present for pain (epineurotomy: 87.5% pre-op pain decreased to 12.5% 2 years post-op vs. no epineurotomy: 92.6% decreased to 29.6%). Nerve conduction velocities increased in both groups and did not differ between (pre/post-op): epineurotomy 31.1/43.8ms vs. 30.0/40.4 (p = 0.32). Patients happy/very happy with results in 73% epineurotomy vs. 70%.</td>
<td>&quot;The study data do not support the use of Epineurotomy as an adjunctive procedure during carpal tunnel release.&quot;</td>
<td>The trial is described as a comparative trial, but appears to involve a randomization procedure based on hospital chart number. Demographic variables were balanced between the two groups; however, the group sizes were not.</td>
<td></td>
</tr>
<tr>
<td>Foulkes 1994</td>
<td>1994</td>
<td>RCT</td>
<td>4.0</td>
<td>N = 33 (36 hands) with CTS who had not had previous surgery on same side.</td>
<td>Epineurotomy Group (n = 23, 26 hands) vs. Non-Epineurotomy Group- Non-treatment group (n = 10, 10 hands) Follow-up 6, 12 months post-op.</td>
<td>Results for sensibility not significant between groups at 6 months (p = 0.64) and 12 months (p = 0.99). No significant difference in grip strength between groups at 6 months (p = 0.79) or 12 months (p = 0.28).</td>
<td>&quot;The addition of an adjunctive epineurotomy, although safe, offers no clinical benefit in the surgical treatment of carpal tunnel syndrome in our series of patients.&quot;</td>
<td>Sparse methodological details. Operating surgeons cannot be blinded. Epineurotomy not superior in carpal tunnel surgery.</td>
<td></td>
</tr>
<tr>
<td>Lowry 1988</td>
<td>1988</td>
<td>RCT</td>
<td>8.0</td>
<td>N = 50 (gender not specified) Standard ligament release Surgery alone group (n = 25) vs. Epineurotomy Group; 10.5 mm3 vs. 7.2 mm3 (p = 0.056); not significant. No significant difference found at 180 day follow-up (p = 0.452). Both groups significantly increased in nerve volume size compared to baseline (p&lt;0.001).</td>
<td>in selected patients longitudinal epineurotomy of the median nerve does not confer any relevant electrophysiological or clinical benefit (nor harm), as compared to a simple dissection of the carpal ligament.&quot;</td>
<td>&quot;The results of this study indicate that standard surgical release of the tunnel release, but some pain relief in the control group compared to study group.</td>
<td>No benefit shown for severe CTS.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
No mention of sponsorship or COI.

<table>
<thead>
<tr>
<th>Study</th>
<th>Characteristics</th>
<th>Procedure</th>
<th>Follow-up</th>
<th>Findings</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mackinnon 1991</td>
<td>N = 79 (gender not specified) with idiopathic CTS. Mean age 58.5 years.</td>
<td>Open carpal tunnel release with internal neurolysis (n = 29, 31 hands) vs. open carpal tunnel release without internal neurolysis (n = 30, 32 hands) Follow-up for 6 months.</td>
<td>Relief of symptoms 88% in release only group vs. 81% of neurolysis group. Among those with abnormal pre-op 2-point discrimination, 62% recovered normal sensation in open release group vs. 55% of neurolysis group. Grip strengths increase from 15-19kg in open release only group vs. from 14 to 17kg in neurolysis group.</td>
<td>“While the technique of internal neurolysis has been proven to be safe and is essential in the surgical evaluation of continuity and in peripheral nerve reconstruction using interfascicular nerve grafting, it would appear from this study that it does not confer improved sensory or motor outcome in patients with primary CTS.”</td>
<td>No benefit</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Flexor Tenosynovectomy</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Shum 2002</td>
<td>N = 87 EDS confirmed (88 wrists) with idiopathic CTS. Mean age 58 years.</td>
<td>Open carpal tunnel release with flexor tenosynovectomy (n = 44 wrists) vs. Open carpal tunnel release without flexor tenosyno-ectomy (n = 44 wrists). Follow-up for 12 months.</td>
<td>Both groups’ symptom severity scores improved after surgery (tenosynovectomy 3.0±0.88 to 1.6±0.68 vs. from 2.9±0.64 to 1.5±0.7, (p ≤0.0002)). No correlations between pre- or post-operative symptoms severity scores and the intraoperative tenosynovial ratings (r = 0.038) or subsequent pathological analyses (r = 0.004 to 0.032).</td>
<td>“We observed neither an added benefit nor an increased rate of morbidity in association with the performance of a flexor tenosynovectomy at the time of carpal tunnel release. We identified no clinical correlations that might predict which individuals would benefit from flexor tenosynovectomy on the basis of either the gross (intraoperative) or histological evaluation of the flexor tenosynovium.”</td>
<td>No benefit. No relationship found with tenosynovial ratings of appearance.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Superficial Nerve Sparing</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Siegmeth 2006</td>
<td>N = 42 (84 hands) with bilateral idiopathic CTS. Age &gt;18 years.</td>
<td>Open carpal tunnel release with superficial nerve sparing (n = 42, 42 hands) vs. open carpal tunnel release</td>
<td>No differences in pain scores at any follow-up interval (graphic presentations of data, 6 weeks; (p = 0.73), 3 months; (p = 0.59), and 6 months; (p = 0.13). No differences found between groups in PEM</td>
<td>“Scar pain scores in this series of open carpal tunnel decompressions were similar, whether or not an attempt was made to identify and preserve...”</td>
<td>Small sample size. Comparable efficacy but the standard carpal tunnel decompression technique took less time to perform.</td>
</tr>
<tr>
<td>Study</td>
<td>Duration</td>
<td>Sample Size</td>
<td>Intervention</td>
<td>Key Findings</td>
<td>Notes</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>----------</td>
<td>-------------</td>
<td>-------------------------------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------</td>
</tr>
<tr>
<td>Macaire 2008</td>
<td>4.0</td>
<td>N = 60</td>
<td>Ultrasound Group-Nerve blocks guided</td>
<td>No differences in grip strength, pillar tenderness or scar sensitivity (p &gt;0.05).</td>
<td>Similar efficacy, but procedure times shorter in polyglactin.</td>
</tr>
<tr>
<td>Citron 1997</td>
<td>4.0</td>
<td>N = 47 with CTS. Mean age 52.1 years.</td>
<td>Standard incision parallel to thenar crease (n = 26) vs. Ulnar L-shaped incision (n = 21). Follow-up at 6 weeks, 3, 6, 9 and 12 months</td>
<td>Mean pain scores at 10 days (nylon, polyglactin and stainless steel): 1.7 (+/-2.2), 3.1 (+/-2.3) and 1.9 (+/-2.3). No significant differences in redness or wound hypertrophy.</td>
<td>“No difference was found in pillar pain between the two incisions, but one had a lower incidence of scar sensitivity.”</td>
</tr>
<tr>
<td>Menovsky 2004</td>
<td>5.0</td>
<td>N = 61 EDS confirmed with CTS. Mean age 50.4 years.</td>
<td>Lengthening of retinaculum (n = 26 hands) performed on one hand vs. simple division of flexor retinaculum standard release (n = 26 hands) performed on other hand. Follow-up at 2, 6, 12, and 25 weeks.</td>
<td>Levine symptom scores (baseline/Weeks 2/6/12/25): open 3.1/ 1.3/1.4/1.2/1.3 vs. lengthen 2.8/1.4/1.3/ 1.2/1.3 (p = 0.63). Function scores were negative (p = 0.66). Grip strengths not different (p = 0.79).</td>
<td>“The study has failed to demonstrate any measurable benefit for this technique. Simple division of the retinaculum is adequate.”</td>
</tr>
<tr>
<td>Dias 2004</td>
<td>8.5</td>
<td>N = 26 EDS confirmed (52 hands) with bilateral CTS. Mean age 57 years.</td>
<td>Preservation of parietal layer of ulnar bursa beneath flexor retinaculum during open release (n = 57) vs. Bursal division (n = 61). Final follow-up at 8-9 weeks.</td>
<td>Grip strengths at follow-up 79% of pre-op values in those with ulnar bursal preservation vs. 82% among other group (p &gt;0.05).</td>
<td>“In this group of patients, preservation of the ulnar bursa around the median nerve during open carpal tunnel release produced no significant difference in grip strength or self-rated symptoms.”</td>
</tr>
<tr>
<td>Forward 2006</td>
<td>8.5</td>
<td>N = 118 (84 female/34 male) with CTS. Mean age 57 years.</td>
<td>Nerve blocks guided at 3, 6, 9 and 12 months after surgery. Follow-up at 6 weeks, 3 and 6 months after surgery.</td>
<td>scores at 6 weeks (p = 0.93), 3 months (p = 0.43), and 6 months (p = 0.38).</td>
<td>superficial nerve branches crossing the wound.”</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Group Description</td>
<td>Procedure Details</td>
<td>Results</td>
<td>Summary</td>
</tr>
<tr>
<td>------------------------------</td>
<td>-------</td>
<td>----------------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>Ambulatory Endoscopic Carpal Tunnel Release</strong>&lt;br&gt;Reed Group, Ltd.</td>
<td>47.5 years</td>
<td>Ambulatory endoscopic carpal tunnel release (n = 30) vs. Nerve Stimulation-Nerve blocks using sensory-motor stimulation (n = 30). Follow-up immediately after surgery.</td>
<td>Significantly less time (s) than nerve stimulation to perform median nerve block: 55 s vs. 100 s (p = 0.002) and time (s) to perform ulnar block: 58 s vs. 80 s (p = 0.02). Mean VAS pain score not significant between groups for venipuncture (p = 0.26) and block puncture (p = 0.72). Ultrasound-guided nerve blocks reduce the performance time while the total time until readiness for surgery remains unaltered compared with nerve stimulation.</td>
<td>Ultrasound-guided nerve blocks reduce the performance time while the total time until readiness for surgery remains unaltered compared with nerve stimulation.</td>
<td></td>
</tr>
<tr>
<td><strong>Open Release vs. Knifelight</strong>&lt;br&gt;Open Release vs. Knifelight</td>
<td>6.5</td>
<td>N = 26 (23 female/9 male) with bilateral CTS. Mean age 48 years. 2.5cm open incision (n = 26, 26 hands) vs. 1-1.5cm Knifelight incision (n = 26, 26 hands). Follow-up at 2 and 6 weeks.</td>
<td>Knifelight vs. Open release (Median): return to work (in weeks): 2.0 vs. 2.0 (p = 0.80); grip strength recovery (%): 89 vs. 84 (p = 0.25); scar tenderness: 1 vs. 10 (p = 0.01)</td>
<td>There was little difference between the two techniques with regard to time taken to return to work, return of grip strength, symptom relief, complications, incidence of pillar pain and patient preference. However, the incidence of scar tenderness was significantly lower with the Knifelight technique.</td>
<td>Faster return to work and less scar tenderness with Knifelight.</td>
</tr>
<tr>
<td><strong>Helm 2003</strong>&lt;br&gt;RCT</td>
<td>6.5</td>
<td>N = 82 (50 female/32 male) with CTS. Mean age 53 years. Open release vs. Knifelight. Incision sizes not specified</td>
<td>Post-op CTS symptoms and grip strengths not different between groups. Mild or moderate scar tenderness Knifelight (89.7%) vs. open (48.8%) (p &lt;0.001). Return to work Knifelight vs. open CTR: 20 vs. 28 days, (p &lt;0.001).</td>
<td>We found no difference in discomfort reported during surgery, in the operative time, in the grip strength measured at 2 and 6 weeks post-operatively or in the proportion of patients cured of their pre-operative symptoms. Knifelight group had a statistically significant improvement in the time to return to work and in scar tenderness at 6 weeks post-operatively.</td>
<td>Some details sparse. No workers' compensation patients.</td>
</tr>
<tr>
<td><strong>Lorgelly 2005</strong>&lt;br&gt;RCT</td>
<td>4.0</td>
<td>N = 185 with CTS. Mean age not reported</td>
<td>Knifelight (2cm incision) (n = 92) vs. Limited open (3-4cm) (n = 89). Mean 30 month follow-up.</td>
<td>First section Boston CTS questionnaire (baseline/19/30 months); Knifelight (3.84/1.46/1.28) vs. open (3.66/2.04/1.39), (NS other than 19 month, p &lt;0.001). RTW 16.6 vs. 25.4 days (p &lt;0.001). Recurrent disease in Knifelight 1% vs. 5% (p &lt;0.01).</td>
<td>Minimally invasive carpal tunnel decompression appears to be more effective but more costly.</td>
</tr>
<tr>
<td>Chandra 2013 RCT</td>
<td>5.0</td>
<td>N = 100 affected by CTS. Mean age 45.59 years.</td>
<td>Early surgery group (&lt;1 week after diagnosis) (n = 51) vs. delayed surgery group (&gt;6 months after diagnosis) (n = 49). Delayed determined by wait-listing. Follow-up after at least 6 months (range, 6-13.2 months; mean, 7.2 months).</td>
<td>Both groups improved in pre-op clinical score (p &lt;0.0001). Mean post-op clinical score lower in early surgery group vs. late surgery group at final follow-up; 8.11 vs. 18.19 (p &lt;0.001). Early group had 100% return to normal activity compared to the late group with 89% (43) with partial return of activity and 11% (6) with normal return to activity (p&lt;0.001).</td>
<td>&quot;On the basis of this study, we propose early surgical (1 week) intervention in patients with moderately severe (grade 3–4) CTS.&quot;</td>
</tr>
</tbody>
</table>
PERIOPERATIVE ANTIBIOTICS
Perioperative antibiotics have been administered to patients undergoing carpal tunnel release, most commonly as pre-incisional antibiotics rather than post-operative antibiotic courses. Some surgeons use antibiotics in all patients. Also, some institutions have implemented policies mandating use in all cases.

1. **Recommendation: Perioperative Antibiotics for Patients Undergoing Carpal Tunnel Release**

   Pre-incisional antibiotics are recommended for consideration for patients with risk factors undergoing carpal tunnel release. Thresholds for use in other patients should be generally low.

   *Indications* – Patients with risk factors (e.g., diabetes mellitus, susceptibility to infections) who are undergoing carpal tunnel release surgery. Institutions may also mandate use through policies.

   *Strength of Evidence* – **Recommended, Insufficient Evidence (I)**
   *Level of Confidence* – **Low**

2. **Recommendation: Routine Use of Antibiotics for Patients Undergoing Carpal Tunnel Release**

   Routine use of antibiotics for all patients undergoing carpal tunnel release is not recommended.

   *Strength of Evidence* – **Not Recommended, Insufficient Evidence (I)**
   *Level of Confidence* – **Low**

**Rationale for Recommendations**

There are no quality studies regarding the administering of peri-operative antibiotics to patients undergoing carpal tunnel release. Infections among these patients are quite uncommon. Antibiotics are invasive when administered intravenously, have low adverse effects, and are moderate to high cost depending on frequency and route of administration. Risk factors among patients, such as diabetics or those who are susceptibility to infections, should be considered. As noted, some institutions mandate the use of these antibiotics, and there is no quality evidence to overturn those policies. However, routine use is not generally recommended.

**Evidence for the Use of Perioperative Antibiotics**
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: perioperative antibiotics or antibiotic prophylaxis, carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, burning, itching, numbness, tingling, hand, palm, finger, wrist, pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 3 articles in PubMed, 177 in Scopus, 0 in CINAHL, and 41 in Cochrane Library. We considered for inclusion 0 from PubMed, Scopus, CINAHL, and Cochrane Library and 0 from other sources. Zero articles met the inclusion criteria.

**ANESTHETIC ISSUES FOR CARPAL TUNNEL SURGICAL RELEASES**

Anesthetic techniques for carpal tunnel release and other hand surgery have ranged from general anesthesia to axillary/regional blocks to local infiltration.(888, 970) Tourniquets have also been used.(971)

**Recommendation: Anesthesia during Carpal Tunnel Release**

Anesthesia, either local or regional, is recommended during carpal tunnel release.

*Strength of Evidence* – **Recommended, Insufficient Evidence (I)**
*Level of Confidence* – **High**

**Rationale for Recommendation**
There are no head-to-head comparative trials for most of these anesthetic techniques, thus evidence-based recommendations are not supportable. Ketorolac has been found useful as an adjunct to bier blocks for hand surgery. (972)

Evidence for Use of Anesthesia during Carpal Tunnel Release

There is 1 high- (973) and 8 moderate-quality RCTs (974-981) incorporated into this analysis. There are 7 low-quality RCTs in Appendix 2. (982-988)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: anesthesia, local, carpal tunnel syndrome, median neuropathy, CTS, carpal tunnel, median nerve, compression, entrapment, neuropathy, nerve disease, syndrome, median nerve, median neuropathy, burning, itching, numbness, tingling, hand, palm, finger, wrist, pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, retrospective, and prospective studies; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 15 articles in PubMed, 3165 in Scopus, 11 in CINAHL, and 44 in Cochrane Library. We considered for inclusion 15 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 15 articles considered for inclusion, 15 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peng 2002 RCT No mention of sponsorship or COI</td>
<td>9.5</td>
<td>N = 51 (24 female/16 male) undergoing hand surgery. Mean age for Lidocaine and Ropivacaine group: 43±19 and 42±13.</td>
<td>Ropivacaine 0.375% injected over a period of 1 minute (n = 20) vs Lidocaine 0.5% forearm regional anesthesia (n = 20). Follow-up for 15 minutes.</td>
<td>Onset of anesthesia 6.5±2.9 minutes for lidocaine vs. 8.0±4.1 minutes for ropivacaine. Pain ratings lower among ropivacaine group throughout first 90 minutes.</td>
<td>&quot;0.375% ropivacaine provides effective anesthesia and superior postoperative analgesia compared with 0.5% lidocaine when forearm IVRA is used.&quot;</td>
<td>Study demonstrates ropivacaine provides superior anesthetic effect to lidocaine in IV regional anesthesia for hand surgery.</td>
</tr>
<tr>
<td>Bigat 2006 RCT Sponsored in part by Akdeniz University Scientific Research Project Unit, Antalya / Turkey. No mention of COI.</td>
<td>7.5</td>
<td>N = 50 (28 female/22 male) undergoing elective carpal tunnel release surgery. Mean age for group L/LD and LDC: 42.3±15.8/40.6 ±14.0/45.1±12.4.</td>
<td>Group L received 3mg/kg lidocaine (n = 25) vs Group LD received 3mg/kg lidocaine plus 8mg dexamethasone (n = 25) vs. Group LDC received 3mg/kg lidocaine for IVRA and 8mg dexamethasone IV (n = 25). Follow-up at 5, 10, 15, 30, 60, and 120 minutes.</td>
<td>Duration of motor blockade 13 minutes LD group vs. 8 IVRA and 6 LDC, p = 0.04. LD requested less analgesics post-operatively (36% vs. 72% and 60%), p = 0.033. Mean analgesics consumed: IVRA 520±390 vs. LD 200±285 vs. LDC 420±445mg (p = 0.016 between LD and IVRA).</td>
<td>&quot;The addition of 8mg dexamethasone to lidocaine for IVRA in patients undergoing hand surgery improves postoperative analgesia during the first postoperative day.&quot;</td>
<td>Baseline differences; blinding details sparse.</td>
</tr>
<tr>
<td>Alayurt 2004 RCT Sponsored by the Akdeniz University Scientific Research Project Unit, Antalya / Turkey. No mention of COI.</td>
<td>7.0</td>
<td>N = 60 (gender not specified) scheduled for surgery of hand or forearm. Mean age Group L/LS/LT and LC: 32±13/31±13/31±11/and 33±14.</td>
<td>Group L: 35ml 0.5% lignocaine with 5ml saline (n = 15) vs. Group LS: sufentanil 25µg (n = 15) vs. Group LT: tramadol 100mg (n = 15) vs. Group LC: clonidine 1µg.kg-1 (n = 15). Follow-up for 24 hours.</td>
<td>No difference between groups in intra-operative hemodynamic data, time to recovery of sensory block, onset and recovery of motor block, sedation scores or postoperative pain. Group with saline had a longer delay of sensory block (p&lt;0.001).</td>
<td>&quot;Addition of sufentanil, tramadol, or clonidine to lignocaine shortened the onset of the sensory block, delayed the onset time of the tourniquet pain and reduced the intraoperative consumption of opioid, but did not affect postoperative pain.&quot;</td>
<td>Blinding details sparse.</td>
</tr>
<tr>
<td>Bigat 2005 RCT Sponsored by the Akdeniz University Scientific Research Project Unit, Antalya / Turkey. No mention of COI.</td>
<td>7.0</td>
<td>N = 50 (gender not specified) undergoing elective hand surgery for CTS. Mean age for Group L and R: 41.95 ± 16.44 and 49.40 ± 7.60.</td>
<td>Group R received 1% ropivacaine (n = 25) vs. Group L received 2% lidocaine intravenous regional anesthesia (N = 25). Follow-up for 24 hours after the surgery.</td>
<td>Pain scores elevated from 30-120 minutes lidocaine vs. ropivacaine group (graphic data, p &lt;0.05). Time to first analgesics lidocaine 226.4±237.1 for ropivacaine vs. 91.7±214.2 minutes (p &lt;0.05). (Data appear reversed between groups for that outcome). Mean paracetamol consumption 550±390 vs. 175±335mg, p &lt;0.05.</td>
<td>&quot;[R]opivacaine 1 mg/kg provided effective anaesthesia and long-lasting postoperative analgesia compared with lidocaine.&quot;</td>
<td>Randomization, allocation details sparse. No assessor blinding.</td>
</tr>
<tr>
<td>Antalya/Turkey</td>
<td>No COI.</td>
<td>7.0</td>
<td>N = 56 (gender not specified) with CTS undergoing a release procedure. Mean age in saline group/30µg clonidine/90µg clonidine/and 51 in 300µg clonidine group: 48/49/49/and 51.</td>
<td>30µg clonidine in 400mg lidocaine group (n = 14) vs. 90µg clonidine in 400mg lidocaine group (n = 14) vs. saline control group in 400mg lidocaine (n = 14). Follow-up at baseline, 20, 40, 60, 80, 140, 200 and 260 minutes post release.</td>
<td>Sensory blockage significantly more prominent at all assessments vs. saline group (p &lt;0.01). At 20 and 30 minutes, all clonidine-dose groups significantly higher sedation rates vs. saline control group, (p &lt;0.01). Those in 30µg and 300µg clonidine groups exhibited significantly higher sedation rates at 20, 40, 140 minute assessments vs. those who received saline: 20 (p &lt;0.05), 40 (p &lt;0.01), 140 (p &lt;0.05). At 40 minute assessment, 300µg group had higher sedation rate vs. 90µg group (p &lt;0.05).</td>
<td>Most lidocaine patients (60%) used analgesics vs. 20% ropivacaine.</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Lawrence 2002</td>
<td>Sponsored by the Wishbone Trust. No mention of COI.</td>
<td>7.0</td>
<td>N = 56 (34 female/22 male) undergoing carpal tunnel decompression. Mean age EMLA 55 years, placebo 52 years.</td>
<td>Eutectic mixture off local anesthetics (EMLA) 5ml (n = 29) vs. placebo 5ml (n = 27) at least 1 hour before surgery. All then received 8ml 0.5% bupivacaine infiltrated over 60 second period. Follow-up post-op.</td>
<td>Lower pain scores with EMLA group, 23±10, vs placebo, 35±16 for both needle insertion (p = 0.0012) and anesthetic injection, EMLA 29±14 vs. placebo 46±19 (p = 0.0005).</td>
<td></td>
</tr>
<tr>
<td>Reuben 1996</td>
<td>RCT</td>
<td>7.0</td>
<td>N = 60 (gender not specified) undergoing either elective carpal tunnel release or tenolysis performed by the same surgeon. Mean age for group 1 / 2 and 3: 47.3 ± 13.6 /52.8 ± 18.3 / and 51.9 ± 15.8.</td>
<td>All 40mL 0.5% lidocaine IV regional anesthesia and 1% lidocaine infiltration. Group 1 or control: no adjuvant (n = 20) vs. Group 2: 60mg ketorolac with IVRA (n = 20) vs. Group 3: 60mg ketorolac infiltration to surgical site (n = 20). Follow-up 24 hours post-op.</td>
<td>VAS scores lower in 2 groups who received ketorolac (p &lt;0.05). Mean time from tourniquet release to first medication 109+/−73 minutes for Group 1, 467+/−431 for Group 2, and 393+/−312 for Group 3 (p &lt;0.05). Numbers of tablets taken: 4.1+/−1.3 Group 1; 1.8+/−1.2 Group 2; and 2.0+/−1.3 Group 3 (p &lt;0.05).</td>
<td></td>
</tr>
<tr>
<td>Patil 2006</td>
<td>RCT</td>
<td>5.5</td>
<td>N = 29 with bilateral carpal tunnel syndrome. Mean age 54 (35 – 81). (Modified Gale) 6mL 2% lignocaine site infiltration (n = 14) vs. (modified Altissimi and Mancini technique (p = 0.02).</td>
<td>Six patients experienced intra-operative pain with the Gale technique, versus none with the Altissimi and Mancini technique (p = 0.02).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>N</td>
<td>Criteria</td>
<td>Interventions</td>
<td>Outcomes</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>--------</td>
<td>---</td>
<td>----------</td>
<td>---------------</td>
<td>----------</td>
</tr>
<tr>
<td>Nabhan 2011 RCT</td>
<td>Nabhan 2011</td>
<td>RCT</td>
<td>5.0</td>
<td>N = 44 with CTS confirmed by nerve conduction testing and physical exam lasting &gt;3 months with no prior surgery; Mean (±SD) age 55.0 (±14.0) for all participants.</td>
<td>Local anesthesia group receiving 20ml of pilocaine via 22 gauge needle (n = 22) vs. Intravenous regional anesthesia group receiving 30ml of 1% prilocaine via 20 gauge cannula (n = 22). Assessments at baseline, 2 weeks and 6 months post-op.</td>
<td>Both groups showed significant improvement at 2 weeks and 6 months after procedure for hand function, ADLs, work performance, pain, and patient satisfaction values when compared to baseline. Mean tourniquet inflation time significantly higher in IVRA group compared to LA group: 27.5 (±2.3) vs. 13.0 (±2.8) minutes, (p = 0.01). Mean operating room time also higher in IVRA group vs. LA group: 45 (±3.9) vs. 28 (±3.5) minutes, (p = 0.01).</td>
</tr>
</tbody>
</table>

Mancini) 3.5mL 2% lignocaine infiltrated in incision line and 2.5mL 2% lignocaine infiltrated into carpal tunnel (n = 15). Follow-up 24 hours after surgery. | Altissimi and Mancini technique required significantly lower numbers of analgesic tablets. | Surgical procedures of different hands at different times. |
Follow-up Visits

Carpal tunnel surgical patients usually have a good recovery, although it can be variable and determined by many factors, including severity of the condition, surgical results, complications, coexisting medical conditions, motivation, pain tolerance, compliance with post-operative instructions, speed of returning to activities of daily living, and speed of returning to work. Carpal tunnel release patients have undergone numerous formal rehabilitation programs. However, as the surgical procedure has become less invasive, the overall trend is towards less formal rehabilitation or courses with fewer appointments. In an increasing number of cases this now includes home exercises and graded increased use. Rehabilitation has included range-of-motion exercises, strengthening exercises, splinting, and a virtual reality system. (989) Home exercise programs appear to be the most effective for regaining function. (990)

Most patients require one or two follow-up clinical appointments for wound care and instructions. Patients with less optimal outcomes may require additional appointments to monitor and facilitate recovery. Patients with physically demanding jobs whose initial restrictions are not accommodated may require a greater number of appointments to monitor their recovery and help facilitate their return to work at appropriate intervals.

While most recovery occurs within the first 3 months after surgery, a full functional recovery from carpal tunnel release including attaining a maximum grip strength is estimated to minimally occur at 6 months and for some patients as long as 1 year. For more information regarding post-operative rehabilitation, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

Triangular Fibrocartilage Complex (TFCC) Tears

Diagnostic Criteria

The history should include ulnar wrist joint pain and a catching, snapping or popping sensation in the wrist with movement. The physical examination should reproduce these symptoms. Imaging studies should be consistent with a triangular fibrocartilage complex (TFCC) tear of sufficient magnitude to explain the symptoms. Other TFCC tears do not have all these features, yet are found in the course of imaging for wrist abnormalities. (14, 220, 221) These other tears generally represent asymptomatic prevalent tears discovered through imaging. Treatment of those tears is usually not indicated. TFCC tears are most commonly classified by the Palmer classification system (223) which has been utilized to develop treatment recommendations (see Table 5). However, overlap may be present between the types in particular due to concomitant degenerative and traumatic issues.

Table 5. Palmer Classification of TFCC Tears and Treatment Recommendations. Type I are acute, traumatic injuries and Type II are degenerative.*

<table>
<thead>
<tr>
<th>Type</th>
<th>Anatomy</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>IA</td>
<td>Avascular articular disc tear</td>
<td>Immobilization. Arthroscopic debridement if immobilization unsuccessful.</td>
</tr>
<tr>
<td>IB</td>
<td>Base of the styloid tear</td>
<td>Immobilization. Arthroscopic or open surgery if immobilization fails.**</td>
</tr>
<tr>
<td>IC</td>
<td>Carpal detachment</td>
<td>Immobilization. Arthroscopic or open surgery if immobilization fails.</td>
</tr>
<tr>
<td>ID</td>
<td>Detachment off the radius</td>
<td>Immobilization. Arthroscopic or open surgery if immobilization fails.</td>
</tr>
<tr>
<td>IIA</td>
<td>Thinning of the articular disc without tear</td>
<td>Address degenerative joint disease risks.*** Surgery rarely indicated. Possible ulna shortening in select cases.</td>
</tr>
<tr>
<td>IIB</td>
<td>Thinning of articular disc accompanied by chondromalacia of the lunate or ulna</td>
<td>Address degenerative joint disease risks. Surgery rarely indicated. Possible ulna shortening in select cases.</td>
</tr>
<tr>
<td>IIC</td>
<td>Central disc tear with chondromalacia</td>
<td>Address degenerative joint disease risks. Surgery for residual symptoms, including ulna shortening and wafer procedure.</td>
</tr>
<tr>
<td>IID</td>
<td>Central tear, chondromalacia and lunotriquetral ligament disruption</td>
<td>Address degenerative joint disease risks. Surgery for residual symptoms including ulna shortening and wafer procedures. Possible arthrodesis.</td>
</tr>
</tbody>
</table>
Central tear, chondromalacia and lunotriquetral ligament disruption and ulnocarpal arthritis

Address degenerative joint disease risks. Surgery for residual symptoms

**Surgery of these is felt to be rarely necessary due to vascular supply.
***Degenerative joint disease risks include body mass index, gout, rheumatoid arthritis, other inflammatory arthropathies, and repeated forceful wrist use.

### Special Studies and Diagnostic and Treatment Considerations

#### X-RAYS

**Recommendation: X-rays to Diagnose Triangular Fibrocartilage Complex (TFCC) Tears**

*X-rays are recommended to diagnose triangular fibrocartilage complex (TFCC) tears.*

**Indications** – Suspected TFCC tear and/or to rule out other sources of wrist pain.

**Frequency/Duration** – Obtaining x-rays once is generally sufficient.

**Strength of Evidence** – **Recommended, Insufficient Evidence (I)**

**Level of Confidence** – **Moderate**

**Rationale for Recommendation**

There are no quality studies evaluating x-rays for the diagnosis of triangular fibrocartilage complex (TFCC) tears. Some patients do not require initial x-rays and can be managed clinically. However, x-rays may assist particularly in ruling out other potential sources of wrist pain. They are also indicated for those who fail to improve or have other symptoms suggesting consideration of other potential diagnoses. X-rays also assist with analysis for evidence of other conditions such as osteoarthrosis. Positive ulnar variance (an ulna that extends more distally than the radius) is thought to increase risk.

#### MR ARTHROGRAPHY AND MRI

**Recommendation: MR Arthrography or MRI to Diagnose Triangular Fibrocartilage Complex (TFCC) Tears**

*MR arthrography or MRI is recommended to diagnose triangular fibrocartilage complex (TFCC) tears.*

**Strength of Evidence** – **Recommended, Insufficient Evidence (I)**

**Level of Confidence** – **Moderate**

**Rationale for Recommendations**

There are no quality studies evaluating MR arthrography or MRI for the diagnosis of triangular fibrocartilage complex (TFCC) tears. MR arthrography is thought to be superior. Traditional arthrography without MRI has mostly been replaced by MR arthrography and MRI. (991-996) Virtual MR arthroscopy is in development, but its utility is not yet demonstrated. (997)

#### ARTHROSCOPY

Diagnostic arthroscopy is often combined with surgical repair (see Surgery section).

### Initial Care

Splinting has been used for treatment of TFCC tears (998) as have ice, heat and rest.

1. **Recommendation: Relative Rest for Acute, Subacute, or Chronic Triangular Fibrocartilage Complex (TFCC) Tears**

   *Relative rest is recommended for treatment of acute, subacute, or chronic triangular fibrocartilage complex (TFCC) tears.*

   **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**

   **Level of Confidence** – **Low**
1. **Recommendation: Splinting for Moderate or Severe Acute or Subacute Triangular Fibrocartilage Complex (TFCC) Tears**

   Splinting is recommended for treatment of moderate or severe acute or subacute triangular fibrocartilage complex (TFCC) tears, particularly to reduce forearm rotation.

   **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   **Level of Confidence** – Low

2. **Recommendation: Self-application of Ice for Acute, Subacute, or Chronic Triangular Fibrocartilage Complex (TFCC) Tears**

   Self-application of ice is recommended for treatment of acute, subacute, or chronic triangular fibrocartilage complex (TFCC) tears.

   **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   **Level of Confidence** – Low

3. **Recommendation: Self-application of Heat for Acute, Subacute, or Chronic Triangular Fibrocartilage Complex (TFCC) Tears**

   Self-application of heat is recommended for treatment of acute, subacute, or chronic triangular fibrocartilage complex (TFCC) tears.

   **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   **Level of Confidence** – Low

---

**Rationale for Recommendations**

There is no evidence of the efficacy of wrist splints to treat acute, subacute, or chronic TFCC tears (knee menisci which are structurally similar, appear to heal with use). Yet, there may be cases where a wrist splint seems helpful and others have recommended immobilization (see Table 5). Splints may help with avoiding aggravating activities or actions that provoke symptoms and therefore, may be more appropriate for acute or moderate to severe injuries. There are also no quality studies evaluating relative rest, rest, ice, or heat for TFCC tears. However, limitations are often needed for more symptomatic cases. Though not invasive, limitations can be moderate to high cost over time; however, relative rest may preclude the need for surgical intervention. Ice and heat may help particularly with more acute symptoms. These treatments may help with symptomatic relief, are not invasive, have no adverse effects, and are not costly and are recommended.

**Evidence for the Use of Initial Care**

There are no quality studies incorporated into this analysis.

**Splinting**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Splinting or Immobilization; Triangular fibrocartilage complex (TFCC) tears; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 6 articles in PubMed, 16 in Scopus, 1 in CINAHL, and 52 in Cochrane Library. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 1 from Cochrane Library and 2 from other sources. Of the 4 articles considered for inclusion, 0 randomized trials and 2 systematic studies met the inclusion criteria.
Ice
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Ice; Triangular fibrocartilage complex (TFCC) tears; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library and 0 in other sources. Zero articles met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Heat, Self-application of heat; Triangular fibrocartilage complex (TFCC) tears controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion zero articles in PubMed, zero in Scopus, zero in CINAHL, zero in Cochrane Library and zero in other sources. Zero articles met the inclusion criteria.

Follow-up Visits
Patients generally require from 1 to 6 appointments, depending on severity and need for workplace limitations. Greater numbers of appointments may be required for evaluating and treatment pain and monitoring function and work status over time. Severe TFCC tears, especially those that either are immobilized for many weeks or undergo surgery may require occupational or physical therapy typically for teaching mobilization exercises and strengthening exercises.

Medications
Over-the-counter medications are generally helpful. Prescription medications may be needed in moderate to severe cases.

NSAIDS/ACETAMINOPHEN
Recommendation: NSAIDs or Acetaminophen for Acute, Subacute, or Chronic Triangular Fibrocartilage Complex (TFCC) Tears
NSAIDs or acetaminophen are recommended to control pain associated with acute, subacute, or chronic TFCC tears particularly for patients with significant pain.

Indications – Pain due to acute, subacute, or chronic TFCC tears.

Frequency/Duration – Scheduled dosage rather than as needed is generally preferable.

Indications for Discontinuation – Resolution of pain, lack of efficacy, development of adverse effects particularly gastrointestinal.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
There are no quality studies evaluating NSAIDs or acetaminophen for controlling pain associated with TFCC tears. However, NSAIDs may help particularly with more acute symptoms. These medications are not invasive, have low adverse effects for short-term use in employed populations, and are not costly. Thus, NSAIDs or acetaminophen are recommended for treatment of pain from acute, subacute, or chronic TFCC tears.

Evidence for the Use of NSAIDs/Acetaminophen
There are no quality studies incorporated into this analysis.
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Acetaminophen, anti-inflammatory agents, non-steroidal, NSAIDS, triangular fibrocartilage, TFCC, triangular fibrocartilage complex, tears, injuries, lesions, triangular fibrocartilage injuries, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion zero articles in PubMed, zero in Scopus, zero in CINAHL, zero in Cochrane Library and zero in other sources. Zero articles met the inclusion criteria.

**Physical Methods/Rehabilitation**

**EXERCISE**

Exercise is generally not indicated acutely; however, exercise may be needed in the recovery or post-operative phases.

**Evidence for the Use of Exercise**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Exercise; triangular fibrocartilage, TFCC, triangular fibrocartilage complex, tears, injuries, lesions, triangular fibrocartilage injuries, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed zero articles in PubMed, zero in Scopus, zero in CINAHL, and 1 in Cochrane Library. We considered for inclusion zero from PubMed, zero from Scopus, zero from CINAHL, zero from Cochrane Library and zero from other sources. Of the zero articles considered for inclusion, zero randomized trials and zero systematic studies met the inclusion criteria.

**Surgery**

1. **Recommendation: Arthroscopic or Open Surgical Repair for Subacute or Chronic Triangular Fibrocartilage Complex (TFCC) Tears**

   Surgical repair (arthroscopic or open) is recommended for patients with instability, concomitant fractures, or symptoms that persist without trending towards resolution despite non-operative treatment and the passage of approximately 3 to 6 weeks.
   
   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – Moderate

   **Rationale for Recommendation**

   There are no quality studies evaluating surgical repair for TFCC tears. Location of the TFCC tear is thought to be related to prognosis with peripheral tears having a better probability of success with non-surgical treatment due to vascular supply; however, central tears also may become asymptomatic.(999) Arthroscopic repair is most typically used with excellent or good results reported in 74% of a case series of 35 patients(1000) with other estimates of success up to 93%,(220, 1001-1004) although open repairs may be performed.(1005)

2. **Recommendation: Ulna Shortening and Wafer Procedures for Chronic Triangular Fibrocartilage Complex (TFCC) Tears**

   Ulna shortening(1006) and wafer procedures are recommended for select cases of chronic Types IIC and IID (Table 5) TFCC tears for which non-surgical treatment is unsuccessful and there is a demonstrable ulna positive variance.
   
   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – Low

   **Rationale for Recommendation**
There are no quality studies evaluating ulna shortening and wafer procedures for TFCC tears. However, in select cases with ulna positive variance and without resolution of considerable or incapacitating symptoms or lacking trending towards resolution, this procedure is recommended. This procedure is invasive, has adverse effects, may not be effective, but also may provide either cure or relief of symptoms and thus is recommended for select cases.

Evidence for the Use of Surgery
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Open surgical repair, triangular fibrocartilage, TFCC, triangular fibrocartilage complex, tears, injuries, lesions, tear, injury, triangular fibrocartilage injuries, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 7 articles in PubMed, 29 in Scopus, 0 in CINAHL, and 0 in Cochrane Library. We considered for inclusion 3 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Zero articles met the inclusion criteria.

Crush Injuries and Compartment Syndrome

Special Studies and Diagnostic and Treatment Considerations

X-RAYS
Recommendation: X-rays for Evaluating Crush Injuries or Compartment Syndrome
X-rays are recommended for evaluating patients with crush injuries or compartment syndrome.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – High

Rationale for Recommendation
There are no quality studies evaluating the use of x-rays for crush injuries or compartment syndrome. However, x-rays are essential for evaluating the extent of injuries and identification of fractures.

Evidence for the Use of X-rays
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: x-ray, crush injury, upper extremity; diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 7 articles in PubMed, 2 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 1580 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

MRI/CT
Recommendation: MRI or CT for Crush Injuries or Compartment Syndrome
MRI or CT is recommended for follow-up of select patients with crush injuries or compartment syndrome.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
Initial evaluation of crush injuries or compartment syndrome generally does not require MRI or CT. However, some patients require MRI or CT for evaluation of symptoms and extent of injury and are recommended in select cases.

Evidence for the Use of MRI/CT
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: magnetic resonance imaging or MRI, CT, crush injury, upper extremity; diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, predictive value of tests, efficacy, efficiency. We found and reviewed 5 articles in PubMed, 18 in Scopus, 6 in CINAHL, 1 in Cochrane Library, and 1490 from Google Scholar. Zero articles met the inclusion criteria.

**Initial Care**

1. **Recommendation: Elevation and Relative Rest for Acute Crush Injuries or Compartment Syndrome**
   Elevation and relative rest are recommended for treatment of acute crush injuries or compartment syndrome.
   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Moderate

2. **Recommendation: Splinting for Subacute Crush Injuries or Compartment Syndrome**
   Splinting is recommended after initial treatment for moderate or severe acute and subacute crush injuries or compartment syndrome.
   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Low

3. **Recommendation: Self-application of Ice for Acute Crush Injuries or Compartment Syndrome**
   Self-application of ice is recommended for treatment of acute crush injuries or compartment syndrome. Other cryotherapies may be required in hospital settings for more severe cases.
   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Low

4. **Recommendation: Self-application of Heat for Acute Crush Injuries or Compartment Syndrome**
   Self-application of heat is not recommended for treatment of acute crush injuries or compartment syndrome.
   - **Strength of Evidence** – Not Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Low

**Rationale for Recommendations**
There are no quality studies evaluating rest/elevation, splinting, or self-application of ice or heat to treat crush injuries or compartment syndrome. However, elevation, rest, and ice are believed to be helpful for treatment of these conditions and in milder cases may be the principal treatments administered. These interventions are not invasive, have no adverse effects, and are not costly (other than repeated administrations of cryotherapies in hospital settings where monitoring is required); thus, they are recommended. Splints may assist in symptomatic relief, are not invasive, have few adverse effects, and are low to moderate cost. The type of splint required depends on the type of injury and subsequent debility. Splints are frequently custom made for patients with these injuries. Splints are recommended particularly for patients with moderate to severe injuries.

Evidence for the Use of Initial Care
There are no quality studies incorporated into this analysis.
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: rest, bed rest, initial elevation, initial care, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion zero articles in PubMed, zero in Scopus, zero in CINAHL, 197 in Cochrane Library, 266 in Google Scholar and zero in other sources. Zero articles met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: splint, splints, nocturnal splint, splinting, upper extremity, wrist, wrist injury, crush injury, compartment syndrome, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 22 articles in PubMed, 11 in Scopus, 0 in CINAHL, 52 in Cochrane Library, and 1,929 in Google Scholar and zero in other sources. Zero articles met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: ice, self-application of ice, crush injuries, wrist injury, compartment syndrome, upper extremity, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 43 articles in PubMed, zero in Scopus, 2 in CINAHL, 4 in Cochrane Library and 5,690 in Google Scholar. We considered for inclusion 1 from PubMed, zero from Scopus, zero from CINAHL, zero from Cochrane Library, zero from Google Scholar and zero from other sources. Of the 5,739 articles considered for inclusion, zero randomized trials and 1 systematic studies met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: heat, self-application of heat, crush injuries, wrist injury, compartment syndrome, upper extremity, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 4 articles in PubMed, 1 in Scopus, zero in CINAHL, 85 in Cochrane Library, 8252 in Google Scholar, and zero other sources. Zero articles met the inclusion criteria.

**Follow-up Visits**

Patients generally require multiple follow-up appointments with the number dependent on the severity of the injury. The mildest cases of crush injuries may require 1 to 3 follow-up appointments. Severe cases of compartment syndrome or crush injuries that have major medical complications and activity limitations may require dozens of appointments to evaluate, treat, advance activity limitations and otherwise monitor and actively facilitate clinical progress. Moderate and severe crush injuries and compartment syndrome usually require occupational or physical therapy for teaching mobilization and strengthening exercises. Therapy needs can be extensive (see below).

**Medications**

Over-the-counter medications may be helpful, but most patients require prescription medications for pain, particularly for moderate to severe injuries. Mannitol has been reported as a treatment. (1007)
NSAIDs/ACETAMINOPHEN
Recommendation: NSAIDs or Acetaminophen for Acute or Subacute Crush Injuries or Compartment Syndrome

NSAIDs or acetaminophen are recommended to control pain associated with acute or subacute crush injuries or compartment syndrome.

**Indications** – Pain due to acute or subacute crush injuries or compartment syndrome.

**Frequency/Duration** – Scheduled dosage rather than as needed is generally preferable.

**Indications for Discontinuation** – Resolution of pain, lack of efficacy, development of adverse effects, particularly gastrointestinal.

**Strength of Evidence** – Recommended, Insufficient Evidence (I)

**Level of Confidence** – Moderate

*Rationale for Recommendation*

There are no quality studies evaluating NSAIDs or acetaminophen for patients with crush injuries or compartment syndrome. There is one trial with non-specific limb injury suggesting efficacy of diclofenac (Woo 05). These medications are helpful for numerous other musculoskeletal disorders and are believed helpful for these injuries. As NSAIDs and acetaminophen are non-invasive, have low adverse effects, and are low cost, they are recommended for treatment of pain associated with acute or subacute crush injuries or compartment syndrome.

*Evidence for the Use of NSAIDs/Acetaminophen*

There is 1 moderate-quality RCT incorporated into this analysis.(1008)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDs; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 3 articles in PubMed, 0 in Scopus, 0 in CINAHL, 110 in Cochrane Library, 510 in Google Scholar, and 1 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 1 from Cochrane Library, 0 Google Scholar, and 1 from other sources. Of the 2 articles considered for inclusion, 1 randomized trials and 1 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Woo 2005 RCT Double-blind No mention of sponsorship or COI.</td>
<td>5.5</td>
<td>N = 300 (No mention of Gender) w/ painful isolated limb injuries. Mean Age: Paracetamol group 35.6±12.2; Diclofenac group 38.2±13.1; Indomethacin group 34.2±11.0; Diclofenac and Paracetamol group 38.3±12.7</td>
<td>Paracetamol and placebo group monitored every 30 minutes for 2 hours, same dosage for 3 days. (N=66) Vs Diclofenac and placebo group monitored every 30 minutes for 2 hours, same dosage for 3 days. (N=69). Vs Indomethacin and placebo group monitored every 30 minutes for 2 hours, same dosage for 3 days.</td>
<td>In stage 1 in the emergency department, analog pain scores and rest and with activity was &gt;13 mm in all groups for the first hour. The diclofenac-paracetamol group achieved &lt;13mm range at 90 minutes after ingestion as well as greatest pain reduction score in 2 hours. After 90 minutes all groups pain score was &lt;13mm. No statistical difference between groups at any time. In stage 2, the diclofenac-paracetamol combination was found to be more effective than the other groups.</td>
<td>&quot;Analgesic benefit of oral paracetamol—nonsteroidal anti-inflammatory drug combinations over single nonsteroidal anti-inflammatory drugs or paracetamol treatment is small and of doubtful clinical significance.&quot;</td>
<td>Baseline comparability questionable as diagnoses and distribution of group. No placebo group.</td>
</tr>
</tbody>
</table>
(N=71) Vs Diclofenac and paracetamol group monitored every 30 minutes for 2 hours, same dosage for 3 days. (N=94);
Follow-up at baseline and at 5-8 days after initial presentation. paracetamol group was only group to achieve <13mm average pain reduction score within the first day. It also saw more abdominal pain than any other group. Median patient satisfaction scores (out of 10) with the oral analgesic treatment were 3.0 (3.0 to 4.0; P=.39) and with the study in general were 3.0 (3.0 to 4.0; P=.25).

**OPIOIDS**
See Opioids recommendations in Carpal Tunnel Syndrome section.

**Physical Methods/Rehabilitation**

**EXERCISE**
Exercise is generally not indicated acutely; however, exercise may be needed in the recovery or post-operative phases.

*Evidence for the Use of Exercise*
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Exercise; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 3 articles in PubMed, 43 in Scopus, 5 in CINAHL, 3 in Cochrane Library, 150 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**HYPERBARIC OXYGEN**

*Recommendation: Hyperbaric Oxygen for Acute or Subacute Crush Injuries or Compartment Syndrome*

Hyperbaric oxygen is recommended for treatment of acute or subacute crush injuries or compartment syndrome depending on the nature of the injury. This frequently includes emergency fasciotomy for release of tension from compartment syndromes as well as other surgical procedures to address fractures and other remediable defects.

*Strength of Evidence – Recommended, Evidence (C)*
*Level of Confidence – Low*

*Rationale for Recommendation*
There is one quality study of hyperbaric oxygen (HBO) for treatment of crush injuries with considerable benefits demonstrated including improved healing and reduced need for additional surgeries.(1009) HBO is non-invasive and generally safe,(1010-1014) (Eskes 13, Eskes 11, Dougherty 13, Garcia-Covarrubias 05,
Greensmith 04) although it is high cost. However, HBO is recommended for treatment of patients with moderate to severe crush injuries or compartment syndrome as risks are outweighed by benefits. 

**Evidence for the Use of Hyperbaric Oxygen for Crush Injuries or Compartment Syndrome**

There is 1 moderate-quality RCT incorporated into this analysis. (1009)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Hyperbaric oxygen therapy, hyperbaric oxygenation, HBOT, crush syndrome, crush injury, compartment syndrome, compartment syndromes, upper extremity, hand, arm, forearm; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 15 articles in PubMed, 11 in Scopus, 15 in CINAHL, 5 in Cochrane Library, 1050 in Google Scholar, and 0 from other sources. We considered for inclusion 6 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 2 Google Scholar, and 0 from other sources. Of the 8 articles considered for inclusion, 1 randomized trial and 5 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Compariso n Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bouachour 1996</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>6.5</td>
<td>N = 36 (gender not specified) with Class II or III soft tissue injuries.</td>
<td>HBO therapy 100% O2 at 2.5 atmospheres vs. placebo</td>
<td>Complete wound healing without tissue necrosis requiring surgical excision in 17 HBO patients vs. 10 placebo (p &lt;0.01). Tissue necrosis 1/18 HBO vs. 8/18 placebo. New surgical procedure = 2 (1 patient) vs. 8 (6 patients), p = 0.03 (p = 0.04).</td>
<td>“[T]his study shows the effectiveness of HBO in improving wound healing and reducing repetitive surgery.”</td>
<td>Results suggest HBO beneficial for these more severe injuries with better healing and less repeat surgery required.</td>
</tr>
</tbody>
</table>

**Surgery**

**Recommendation: Surgery for Acute or Subacute Crush Injuries or Compartment Syndrome**

Surgery is recommended for treatment of acute or subacute crush injuries or compartment syndrome depending on the nature of the injury. This frequently includes emergency fasciotomy for release of tension from compartment syndromes as well as other surgical procedures to address fractures and other remediable defects.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*

*Level of Confidence – High*

**Rationale for Recommendation**

There are no quality studies evaluating surgeries for crush injuries or compartment syndrome of the hand or forearm and the clinical variability between patients is large. However, fasciotomies are particularly essential for treatment of significant neurovascular compromise from compartment syndrome and is a surgical emergency. (19, 227, 1009-1014) Other procedures may be required based on remediable defects such as fractures, ligament tears, or other injuries.

**Evidence for the Use of Surgery**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Surgery, surgical procedures, operative, general surgery, crush, wrist injuries, wrist injury, compartment syndrome, compartment syndromes, upper extremity, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 212
articles in PubMed, 250 in Scopus, 17 in CINAHL, and 0 in Cochrane Library. We considered for inclusion 5 from PubMed, 0 from Scopus, 2 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 7 articles considered for inclusion, 0 randomized trials and 1 systematic study met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Emergency fasciotomy, crush injuries, crush, injury, injuries, compartment syndrome, upper extremities, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 44 in Scopus, 0 in CINAHL, and 1 in Cochrane Library. We considered for inclusion 0 from PubMed, 3 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 3 articles considered for inclusion, 0 randomized trials and 2 systematic studies met the inclusion criteria.

**Kienböck Disease**

**Diagnostic Criteria**
Patient has non-radiating wrist compartment pain, limited range of motion, and developed x-ray evidence of radiological collapse of the lunate.

**Special Studies and Diagnostic and Treatment Considerations**

**X-RAYS**

*Rationale for Recommendation*
There are no quality studies evaluating the use of x-rays to diagnose Kienböck disease. However, x-rays are used to confirm the diagnosis and are moderately costly, thus they are recommended. X-rays generally should be taken of both hands.

*Strength of Evidence* – **Recommended, Insufficient Evidence (I)**
*Level of Confidence* – **High**

**Evidence for the Use of X-rays**
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Kienböck's disease, X-ray, radiography, radiograph; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 3 articles in PubMed, 347 in Scopus, 2 in CINAHL, 12 in Cochrane Library, 140 in Google Scholar and zero in other sources. Zero articles met the inclusion criteria.

**CT**

*CT is recommended to diagnose Kienböck disease when xrays are negative or unclear.*

*Strength of Evidence* – **Recommended, Insufficient Evidence (I)**
*Level of Confidence* – **Moderate**
**Rationale for Recommendation**

There is one quality study evaluating the use of CT scans that included patients with Kienböck disease suggesting 3-D CT may provide more information than x-ray or plain CT. (1025) (Nakamura 89) CTs are used to assist with diagnosis and management, thus they are recommended where x-rays are negative or unclear.

**Evidence for the Use of CT**

There is 1 moderate-quality study incorporated into this analysis. (1025) (Nakamura 89)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: computed tomography or CT, Kienböck’s disease; diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 33 articles in PubMed, 3 in Scopus, 0 in CINAHL, 0 in Cochrane Library, and 295 from Google Scholar. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 1 article considered for inclusion 1 diagnostic study met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score</th>
<th>Number</th>
<th>Area of Spine</th>
<th>Diagnoses</th>
<th>Type of CT</th>
<th>X-ray used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Surgery Performed</th>
<th>Long term follow-up</th>
<th>Clinical outcomes</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nakamura 1989</td>
<td>Diagnostic Study</td>
<td>4.5</td>
<td>N = 20 (3 female and 17 male) admitted for wrist problems; 3 with Kienböck’s disease, 14 with fractures or dislocation of the carpal bones</td>
<td>Wrist</td>
<td>Wrist problems due to altered bony or joint structure(s).</td>
<td>High resolution CT scanner (Somatom DRH) and accompanying software (3D Display; Version B or C)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>16/17 cases of fracture a three-dimensional CT image was believe to be useful to detect the fracture line. 3 had a flattened lunate due to Kienbock disease. 13 had deformity of the hamate body seen on plain radiography and CT, but the three-dimensional CT image. Presence and location of small fragments not detected by plain radiographs and CT, but “Three-dimensional CT imaging provides a great deal of information that cannot be obtained by conventional radiographs or CT images even at their present stage of technical development.”</td>
<td>Small sample (N=20). Data suggest 3-D CT provides more diagnostic information than either plain radiography or conventional CT.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
MRI

**Recommendation: MRI to Diagnose Kienböck Disease**

MRI is recommended to diagnose Kienböck disease.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*

*Level of Confidence – Low*

**Rationale for Recommendation**

There are 2 moderate-quality articles evaluating the use of MRIs to diagnose Kienböck disease. However, MRI was not shown to have superior performance for diagnostic purposes. MRIs are used to assist with diagnosis and management, thus they are recommended.

**Evidence for the Use of MRI**

There are 2 moderate-quality studies incorporated into this analysis. (1015, 1016)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Magnetic Resonance Imaging, MRI, Kienböck’s disease or Kienbock disease, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 82 articles in PubMed, 68 in Scopus, 1 in CINAHL, 0 in Cochrane Library, and 523 from Google Scholar. We considered for inclusion 2 from PubMed, 0 from Scopus, CINAHL, Cochrane Library, Google Scholar, and from other sources. Of the 2 articles considered for inclusion 2 diagnostic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Year</th>
<th>Study Type</th>
<th>Author</th>
<th>Score</th>
<th>Number</th>
<th>Area</th>
<th>Diagnoses</th>
<th>Type of MRI used</th>
<th>Type of CT used</th>
<th>T1 weighted</th>
<th>T2 weighted</th>
<th>X-ray</th>
<th>Myelography</th>
<th>Clinical</th>
<th>Surgery</th>
<th>Clinical</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>1996</td>
<td>Diagnostic</td>
<td>Hashizume</td>
<td>4.0</td>
<td>10 (2 female/8 male)</td>
<td>Wrist</td>
<td>Kienbock’s Disease</td>
<td>1.5 Tesla signal, both T1 and T2 weighted images.</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>Mean follow-up 29.</td>
<td>Areas of collapse easily identified in x-ray, tomography, CT and microradiographic images. MRI showed complete loss of signal intensity in T1 images of lesion of lunate.</td>
<td>“MRI is at present unable to distinguish bone necrosis, the histological reactive interface or surrounding hyperaemia in detail.”</td>
<td>Small sample size. Data suggest MRI unable to distinguish bone necrosis in detail.</td>
<td></td>
</tr>
<tr>
<td>1992</td>
<td>Diagnostic</td>
<td>Imaeda</td>
<td>4.0</td>
<td>26 (7 female and 19 male)</td>
<td>Wrist</td>
<td>Kienbock’s Disease</td>
<td>1.5 tesla signal with 3-inch surface coil. Both T1 and T2 weighted images.</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td></td>
<td>For normal wrists, bone marrow showed high signal intensity on T1 and iso intensity on T2. For wrists with Kienbock’s disease, T1 weighted images had decrease in signal intensity in all cases. After osteotomy of radius, signal intensity of lunate returned to normal in both T1 &amp; T2.</td>
<td>“After osteotomy of the radius, the signal intensity of the lunate returned to normal and Lichtman’s stage IL cases had better results than those in stage III. M.R. imaging is ideal for evaluating the lunate in Kienbock’s disease.”</td>
<td>Small sample. Data suggest a low signal intensity of lunate on T-1 weighted images is diagnostic of Kienbock’s disease and signal intensity (if high) correlate to disease severity.</td>
<td></td>
</tr>
</tbody>
</table>
**SCREENING FOR SYSTEMIC DISORDERS**

**Recommendation: Screening for Systemic Disorders for Kienböck Disease**

Screening for systemic disorders is recommended for patients with Kienböck disease.

- **Strength of Evidence** – Recommended, Insufficient Evidence (I)
- **Level of Confidence** – Moderate

**Rationale for Recommendation**

There are multiple disorders that are thought to predispose to Kienböck disease. These disorders may be otherwise asymptomatic, there may be potential to develop other manifestations of these diseases including in the other hand, and it may be possible to slow the rate of progression of this condition through active clinical management. Thus, the threshold for evaluations of systemic metabolic issues (e.g., diabetes, glucose intolerance), alcoholism, and rheumatological studies should be low, particularly as potentially modifiable risks may theoretically slow the rate of progression or prevent the disease in the other hand.

**Evidence for the Use of Screening**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Screening for Systemic Disorders, steroid, trauma, Kienböck’s disease or Kienbock disease, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, predictive value of tests, efficacy, efficiency, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 13 articles in PubMed, 0 in Scopus, 0 in CINAHL, Cochrane Library, and 127 from Google Scholar. We considered for inclusion 0 from PubMed, Scopus, CINAHL, Cochrane Library, Google Scholar, and from other sources. Zero articles met the inclusion criteria.

**Initial Care**

Initial care of patients with Kienböck disease involves identification and elimination or control of potential systemic contributing factors. Patient’s limitations should be based on their capabilities.

1. **Recommendation: Self-application of Ice for Acute, Subacute, or Chronic Kienböck Disease**

   Self-application of ice is recommended for treatment of acute, subacute, or chronic Kienböck disease.

   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Low

2. **Recommendation: Self-application of Heat for Acute, Subacute, or Chronic Kienböck Disease**

   Self-application of heat is recommended for treatment of acute, subacute, or chronic Kienböck disease.

   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Low

3. **Recommendation: Splints for Acute, Subacute, or Chronic Kienböck Disease**

   Splints are recommended for treatment of select patients with acute, subacute, or chronic Kienböck disease.

   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Low

**Rationale for Recommendations**

There are no quality studies evaluating the use of ice or heat for treatment of Kienböck disease. However, these treatments may help with symptomatic relief. These interventions are not invasive, have no adverse effects, and are not costly, thus they are recommended. There are no quality studies...
evaluating splinting for Kienböck disease. A trial may be helpful to assess whether splinting provides symptomatic relief. Splints are not invasive and have few adverse effects over the short term although over the long term there are concerns regarding the potential for accelerated debility disuse and weakness of the wrist. Splints are also low cost. Thus, they are recommended for select patients.

**Evidence for the Use of Initial Care**

There are no quality studies incorporated into this analysis.

**Ice:**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Kienböck’s disease or Kienbock disease; Ice; Self Application; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 0 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Heat:**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Kienböck’s disease or Kienbock disease; HEAT/ Self-Application of Heat; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library and 0 in other sources. Zero articles met the inclusion criteria.

**Splints:**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Screening for Systemic Disorders, steroid, trauma, Kienböck’s disease or Kienbock disease, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, predictive value of tests, efficacy, efficiency, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 13 articles in PubMed, 0 in Scopus, 0 in CINAHL, Cochrane Library, and 127 from Google Scholar. We considered for inclusion 0 from PubMed, Scopus, CINAHL, Cochrane Library, Google Scholar, and from other sources. Zero articles met the inclusion criteria.

**Follow-up Visits**

Patients with Kienböck disease generally require periodic appointments to follow the clinical course. Frequencies of appointments may be greater where workplace limitations are required. Post-operative rehabilitation can be considerable, with a requirement for occupational or physical therapy on a prolonged basis in order for the patient to recover as much function as possible.

**Medications**

Over-the-counter medications are generally helpful for pain associated with Kienböck disease. Prescription medications may be needed for moderate to severe cases. Patients with Kienböck disease often develop chronic pain (see Chronic Pain Guideline for a comprehensive approach to managing chronic pain). An abbreviated approach is noted below.

**NSAIDS/ACETAMINOPHEN**

*Recommendation: NSAIDs and Acetaminophen for Acute, Subacute, or Chronic Kienböck Disease*
NSAIDs and acetaminophen are recommended to control pain associated with acute, subacute, or chronic Kienböck disease.

**Indications** – Pain due to acute, subacute, or chronic Kienböck disease.

**Frequency/Duration** – Scheduled dosage rather than as needed is generally preferable.

**Indications for Discontinuation** – Resolution of pain, lack of efficacy, development of adverse effects particularly gastrointestinal.

**Strength of Evidence** – **Recommended, Insufficient Evidence (I)**

**Level of Confidence** – **Low**

**Rationale for Recommendation**
There are no quality studies evaluating NSAIDs and acetaminophen for Kienböck disease. However, these medications may relieve pain and increase function. They are not invasive, have few adverse effects in employed populations, and are low cost, thus they are recommended.

**Evidence for the Use of NSAIDs/Acetaminophen**
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDS, Acetaminophen, Kienböck’s disease; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 11 articles in PubMed, 2 in Scopus, zero in CINAHL, 3 in Cochrane Library, 132 in Google Scholar, and zero in other sources. Zero articles met the inclusion criteria.

**TOPICAL MEDICATIONS**

**Recommendation:** Topical Medications for Treatment of Acute, Subacute, or Chronic Kienböck Disease
Topical medications including topical creams, ointments, and lidocaine patches are recommended for treatment of pain associated with acute, subacute, or chronic Kienböck disease.

**Strength of Evidence** – **Recommended, Insufficient Evidence (I)**

**Level of Confidence** – **Low**

**Rationale for Recommendation**
There are no quality studies evaluating the use of topical medications for treatment of Kienböck disease. However, these treatments may provide symptom relief. They are not invasive, have few adverse effects in employed populations, and are low cost, thus they are recommended. Caution is warranted if there is use of anesthetic agents over large areas of the body, as adverse effects from systemic absorption have been reported.

**Evidence for the Use of Topical Medications**
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Topical Cream, Topical Ointment, lidocaine patch, topical medication, Kienböck’s disease, Kienbock disease; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 1 article in PubMed, 3 in Scopus, zero in CINAHL, 72 in Cochrane Library, 14 in Google Scholar and zero in other sources. Zero articles met the inclusion criteria.

**OPIOIDS**
See Opioids recommendations in Carpal Tunnel Syndrome section.
Physical Methods/Rehabilitation

**EXERCISE**

Exercise is generally not utilized during acute presentations of Kienböck disease. However, exercise is nearly always necessary for post-operative patients and is frequently used for patients in the subacute and chronic phases.

**Evidence for the Use of Exercise**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: exercise, Kienböck’s disease, Kienbock disease upper extremity, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 35 articles in PubMed, 5 in Scopus, zero in CINAHL, zero in Cochrane Library, 492 in Google Scholar, and zero other sources. Zero articles met the inclusion criteria.

**Surgery**

**Recommendation: Surgical Repair for Chronic Kienböck Disease**

Surgical treatment is recommended as an option for patients with moderate to marked impairment if not improved 8 weeks post-injury or after 6 weeks of non-operative treatment due to Kienböck disease.

**Strength of Evidence** – **Recommended, Insufficient Evidence (I)**

**Level of Confidence** – Low

**Rationale for Recommendation**

There are no quality studies evaluating surgical repair for Kienböck disease. There are many different surgical procedures and no quality comparative studies that have been reported. Surgical procedures utilized have included: lunate excision with silicone implants (1028-1030) (no longer recommended), excision with autogenous soft tissue implants including coiled palmaris longus tendon, (1028, 1031-1036) external fixation, (1035, 1037) arthrodesis, (1038, 1039) radial shortening, (1040, 1041) scaphoid-trapezium-trapezoid fusion, (1036, 1042, 1043) in advanced cases, proximal row carpectomy, (1044-1046), lunate core decompression, (1047, 1048) (Mehrpour 11, Rodrigues-Pinto 12) and vascularized bone transfers, (1049) A comparative clinical trial found superior clinical results and better preservation of carpal height ratio using palmaris longus tendon ball with a bone core compared with no bone core. (1034) In the absence of quality studies, the main determinant of surgical technique is the experience and comfort of the surgeon with specific treatment approaches.

**Evidence for the Use of Surgery**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: surgery, surgical fixation, surgical repair, kienbock’s disease, Kienböck’s disease, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 127 articles in PubMed, 17 in Scopus, 9 in CINAHL, 809 in Google Scholar and 1,348 in Cochrane Library. We considered for inclusion 4 from PubMed, zero from Scopus, zero from CINAHL, zero from Cochrane Library, 4 in Google Scholar and zero from other sources. Of the 8 articles considered for inclusion, zero randomized trials and 8 systematic studies met the inclusion criteria.

**Wrist Sprains**


**Diagnostic Criteria**
A history of an acute traumatic event with forceful loading of the wrist, combined with a negative examination other than ligamentous tenderness and negative x-rays.

**Special Studies and Diagnostic and Treatment Considerations**

**X-RAYS**
X-rays are traditionally the first diagnostic imaging study to evaluate wrist sprains and potential fractures.(1035)

*Recommendation: X-rays for Wrist Sprains*
X-rays are recommended to determine whether a fracture is present, particularly for patients with scaphoid pain or scaphoid tubercle tenderness.(1036)

*Strength of Evidence – Recommended, Insufficient Evidence (I)*
*Level of Confidence – High*

*Evidence for the Use of X-rays*
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: X-Ray, Wrist Sprain, Wrist Sprains, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 15 articles in PubMed, 0 in Scopus, 2 in CINAHL, Cochrane Library, and 55 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 57 articles considered for inclusion 0 diagnostic studies met the inclusion criteria.

**CT SCANS**

*Recommendation: CT scans for Wrist Sprains*
CT scans are recommended to determine whether a fracture is present, particularly for patients with scaphoid pain or scaphoid tubercle tenderness with negative x-rays.(1036)

*Strength of Evidence – Recommended, Insufficient Evidence (I)*
*Level of Confidence – Moderate*

*Evidence for the Use of CT Scans*
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Wrist Sprain, Wrist Sprain, Computed Tomography (CT), diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 13 articles in PubMed, 0 in Scopus, 0 in CINAHL, Cochrane Library, and 432 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 445 articles considered for inclusion 0 diagnostic studies met the inclusion criteria. Zero articles met the inclusion criteria.

**MR ARTHROGRAPHY**

*Recommendation: MR Arthrography for Wrist Sprains*
MR arthrography is recommended for patients without improvement in wrist sprains after approximately 6 weeks of treatment.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*
*Level of Confidence – Low*
Rationale for Recommendations
There are no quality studies evaluating x-rays for wrist sprains. Mild wrist sprain may not necessitate x-rays. There is no evidence other studies are helpful in the acute setting (see discussion of scaphoid fractures for other studies in the presence of ongoing, non-resolving pain.) However, x-rays may assist in diagnosing and treating the condition and thus are recommended. There also are no quality studies evaluating MR arthrography. However, MR arthrograms are helpful to particularly identify ligamentous issues such as scapholunate, lunotriquetral, and TFCC tears that may be diagnosed as simple sprains. Thus, MR arthrography is recommended after approximately 6 weeks of clinical management.

Evidence for the Use of MR Arthrography
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: MR Arthrography, Wrist Sprain, Wrist Sprain, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 4 articles in PubMed, 0 in Scopus, 0 in CINAHL, Cochrane Library, and 244 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 248 articles considered for inclusion 0 diagnostic studies met the inclusion criteria.

Initial Care
1. Recommendation: Relative Rest for Acute Wrist Sprains
   Relative rest is recommended for treatment of acute wrist sprains.
   
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Moderate

2. Recommendation: Splinting for Moderate or Severe Acute or Subacute Wrist Sprains
   Splinting is recommended for treatment of moderate or severe acute or subacute wrist sprains.
   
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Moderate

3. Recommendation: Self-application of Ice for Acute Wrist Sprain
   Self-application of ice is recommended for treatment of acute wrist sprain.
   
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Low

   Self-application of heat is recommended for treatment of acute wrist sprain.
   
   Strength of Evidence – Recommended, Evidence (C)
   Level of Confidence – Low

Rationale for Recommendations
There is one moderate-quality RCT that shows heat is effective in reducing pain from wrist sprains. (1037) There are no quality studies evaluating relative rest, splints, or ice for wrist sprains. However, these treatments may help with symptomatic relief. Splints are recommended particularly for patients with moderate to severe sprains. (Physicians should be aware that as early mobilization of ankle sprains results in improved clinical outcomes and those results may be applicable to the wrist.) These interventions are not invasive, have no adverse effects, and are low cost, thus they are recommended.

Rest:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Rest, wrist sprains; controlled clinical
trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed zero articles in PubMed, 477 in Scopus, zero in CINAHL, zero in Cochrane Library, 1224 in Google Scholar, and zero from other sources. We considered for inclusion zero from PubMed, zero from Scopus, zero from CINAHL, and zero from Cochrane Library, zero Google Scholar, and zero from other sources. Zero articles met the inclusion criteria.

**Splints:**
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: splint, splinting, Wrist Sprain, Wrist Sprain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed zero articles in PubMed, 15 in Scopus, zero in CINAHL, zero in Cochrane Library, zero in Google Scholar, and zero from other sources. Zero articles met the inclusion criteria.

**Ice:**
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ice, hypothermia, cryotherapy, ice packs, wrist sprains, wrist sprain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed zero articles in PubMed, 15 in Scopus, zero in CINAHL, zero in Cochrane Library, 3243 in Google Scholar, and zero from other sources. We considered for inclusion 2 from PubMed, zero from Scopus, zero from CINAHL, and zero from Cochrane Library, zero Google Scholar, and zero from other sources. Of the 2 articles considered for inclusion, zero randomized trials and 2 systematic studies met the inclusion criteria.

**Heat:**
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: Wrist sprains, heat, hot temperatures, therapeutics; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 1290 articles in PubMed, 9 in Scopus, 1 in CINAHL, zero in Cochrane Library, and 2610 in Google Scholar. We considered for inclusion one from PubMed, zero from Scopus, zero from CINAHL, zero from Cochrane Library, zero from google scholar, and zero from other sources. Of the one article considered for inclusion, 1 randomized trial and zero systematic studies met the inclusion criteria.

**Evidence for Heat for Wrist Sprain**
There is 1 moderate-quality RCT incorporated into this analysis.(1037)

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Michlovitz 2004</td>
<td>RCT</td>
<td>Sponsored by Procter &amp; Gamble Health Sciences Institute. COI Erasala.</td>
<td>N = 69 (14 males, 15 females) with acute wrist pain, mostly from sprains, tendinosis, strains, osteoarthritis</td>
<td>Self-applied heat wrap group at 104° F (40°C) for 8 hours daily (N= 29) vs. Oral placebo (N= 30) vs.</td>
<td>Mean pain relief greater in heat wrap than oral placebo (mean pain relief 1.68±0.23 vs. 1.15±0.21 (p = 0.045). Grip strength improved more in heat wrap group 6.44± 1.34kg</td>
<td>Continuous low-level heat therapy is a novel strategy in the treatment of musculoskeletal disorders. In this study, increased pain relief, functional gains, and grip strength along with Short (3 days) treatment. Results for acetaminophen and unheated wrap not reported.</td>
<td></td>
</tr>
</tbody>
</table>
Follow-up Visits
Patients generally require 1 to 3 appointments, depending on severity of the sprain and the need for workplace limitations. Severe wrist sprains may require occupational or physical therapy mostly for teaching mobilization exercises. Wrist sprains that do not resolve or trends towards resolution by 6 weeks should have either further diagnostic evaluation or referral for consideration of other diagnostic testing and treatment options.

Medications
Over-the-counter medications are generally helpful for pain associated with wrist sprain. Prescription medications may be needed for moderate to severe cases.

**NSAIDs/ACETAMINOPHEN**
*Recommendation: NSAIDs and Acetaminophen for Acute or Subacute Wrist Sprain*
NSAIDs and acetaminophen are recommended to control pain associated with acute or subacute wrist sprain.

*Indications – Pain due to acute or subacute wrist sprain.*

*Frequency/Duration – Scheduled dosage rather than as needed is generally preferable.*

*Indications for Discontinuation – Resolution of pain, lack of efficacy, development of adverse effects particularly gastrointestinal.*

- **Strength of Evidence – Recommended, Evidence (C) – NSAIDs**
- **Strength of Evidence – Recommended, Insufficient Evidence (I) – Acetaminophen**
- **Level of Confidence – Moderate**

*Rationale for Recommendations*
There are no quality studies directly evaluating the use of NSAIDs and acetaminophen for pain associated with wrist sprain; however, there are moderate-quality studies of lower extremity sprains (1038, 1039) and these injuries are believed to be analogous (see Ankle and Foot Disorders Guideline). These medications may relieve pain and increase function. They are not invasive, have few adverse effects in employed populations, and are low cost, thus they are recommended for pain associated with acute or subacute wrist sprain.
Evidence for the Use of NSAIDs/Acetaminophen
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDs, non-steroidal anti-inflammatory drugs, Wrist Sprains; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 7 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 50 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

OPIOIDS
See Opioids recommendations in Carpal Tunnel Syndrome section.

Physical Methods/Rehabilitation
EXERCISE
Exercise is generally not indicated acutely. Patients with deficits may require a home exercise program during recovery phases. Some patients require a formal exercise program.

Evidence for the Use of Exercise
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: wrist, sprain, sprains, strain, strains, exercise, exercise therapy; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 22 articles in PubMed, 406 in Scopus, 3 in CINAHL, 5 in Cochrane Library, 330 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Surgery
Recommendation: Surgery for Treatment of Acute or Subacute Wrist Sprain
Surgery is not recommended for treatment of acute or subacute wrist sprain in the absence of a remediable defect.

   Strength of Evidence – Not Recommended, Insufficient Evidence (I)
   Level of Confidence – High

Rationale for Recommendation
There are no quality studies evaluating the use of surgery for wrist sprain. Other than among patients with other trauma necessitating surgery, wrist sprains are not believed to respond to surgery. Ongoing symptoms that do not resolve should be evaluated for other diagnoses.

Evidence for the Use of Surgery
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: surgery, surgeries, general surgery, general surgeries; wrist, sprain, sprains, strain, strains; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 213 articles in PubMed, 335 in Scopus, 2 in CINAHL, 0 in Cochrane Library, 2474 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.
Mallet Finger

Diagnostic Criteria
Mallet finger is a clinical diagnosis with a characteristic presentation of inability to extend the distal segment when the extensor tendon is damaged.

Special Studies and Diagnostic and Treatment Considerations

X-RAYS
Recommendation: X-rays for Mallet Finger
X-rays are recommended in most cases of mallet finger to determine if a fracture is present and to what extent.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
There are no quality studies evaluating the use of x-rays for mallet finger. X-rays may assist in identifying fractures and the magnitude of the involvement of the joint surface, which if large enough, alters management to surgery. It is reasonable to omit x-rays if there is no swelling or tenderness.

Evidence for the Use of X-rays
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: x-ray, computed tomography, radiograph, mallet finger, baseball finger; diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 22 articles in PubMed, 10 in Scopus, 2 in CINAHL, 0 in Cochrane Library, and 243 from Google Scholar. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 2 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

ULTRASOUND
Recommendation: Ultrasound to Diagnose Mallet Finger
Ultrasound is not recommended to diagnose mallet finger.

Strength of Evidence – Not Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There are no quality studies evaluating the use of ultrasound to diagnose mallet finger. While ultrasound has been used for imaging,(1040) there is no evidence it alters treatment or prognosis and x-ray studies appear sufficient for diagnostic purposes. Thus, ultrasound is not recommended to diagnose mallet finger.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: ultrasonography, ultrasound, ultrasound scanning, sonography, mallet finger, baseball, hammer; diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 0 articles in PubMed, 10 in Scopus, 0 in CINAHL, 0 in Cochrane Library, and 178 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 3 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.
Initial Care
Care usually involves a splint and follow-up visits. Large fracture fragments are rare(264, 1041-1044) and necessitate surgery.

SPLINTS

1. **Recommendation: Splints for Acute or Subacute Mallet Finger**
   Extension splinting with the joint in a neutral or hyperextended position is moderately recommended for treatment of acute or subacute mallet finger.(264, 1044)
   
   **Indications** – Acute or subacute mallet finger.
   **Frequency/Duration** – Splinting for 6 to 8 weeks, possible nocturnal use for an additional 2 to 4 weeks.
   **Indications for Discontinuation** – Skin complications, non-compliance.

   **Strength of Evidence** – Moderately Recommended, Evidence (B)
   **Level of Confidence** – High

   **Rationale for Recommendation**
   There are 5 moderate-quality RCTs incorporated in this analysis. Splints must hold the finger in continuous, full extension for a minimum duration of 6 weeks.(1045) Some protocols involve 8 weeks, while some involve nocturnal use for an additional 2 to 4 weeks.(264, 1043-1047) There are many different types of (264, 1043, 1044, 1046, 1047) and no quality evidence of the unequivocal superiority of one versus another.(1048-1050) A padded aluminum splint was reportedly superior compared to a Stack (pre-fabricated plastic) splint due to easier fit and fewer skin complications.(264) Another trial found the Stack splint superior to the Abouna splint.(1044) Extension must be maintained even if the splint is removed for skin hygiene, which is often one of the times non-compliance occurs and is believed to cause many of the treatment failures.(1048) One quality study suggested better outcomes with fixation for patients presenting with delayed treatment.(1041)

   A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: splint, splints, splinting, finger, mallet, baseball, drop, hammer; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 12 articles in PubMed, 68 in Scopus, 3 in CINAHL, 17 in Cochrane Library, 4,110 in Google Scholar, and 0 from other sources. We considered for inclusion 8 from PubMed, 3 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 0 from other sources. Of the 12 articles considered for inclusion, 1 randomized trials and 3 systematic studies met the inclusion criteria.

2. **Recommendation: Instructions for Splint Wear**
   It is recommended that careful instructions on splint wear be provided to patients.

   **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   **Level of Confidence** – High

   **Rationale for Recommendation**
   There are no quality studies evaluating the use of instructions for splint wear for mallet finger. However, instructions appear critical for preventing treatment failures and are thus recommended.

   A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: failed splints, splint failure, surgery, finger, mallet, baseball, drop, hammer; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 7 articles in PubMed, 2 in Scopus, 1 in CINAHL, 3 in Cochrane Library, 407 in Google Scholar, and 0 from other
sources. We considered for inclusion 2 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 3 from Google Scholar, and 0 from other sources. Of the 6 articles considered for inclusion, 0 randomized trials and 2 systematic studies met the inclusion criteria.

**Follow-up Visits**

Patients require a few appointments to reinforce importance of splinting and of not removing the splint unsupported. Multiple appointments are generally not required.

**Medications**

Nonprescription medications are usually not required as mallet finger is generally not painful. Prescription medications are rarely required as mallet finger is generally not painful.

*Evidence for the Use of Medications*

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: mallet finger, baseball, hammer, NSAIDs, NSAID, acetaminophen, non-steroidal anti-inflammatory; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 1 in Scopus, 0 in CINAHL, 13 in Cochrane Library, 75 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Physical Methods/Rehabilitation**

**EXERCISE**

Exercise is not indicated acutely and most patients with mallet finger do not require participation in an exercise program. However, patients usually require careful education about splinting (see Education above). For patients with residual deficits, particularly post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

*Evidence for the Use of Exercise*

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: exercise, physical activity, mallet finger, baseball, drop, hammer; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 3 articles in PubMed, 5 in Scopus, 3 in CINAHL, 1 in Cochrane Library, 187 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 2 articles considered for inclusion, 01 randomized trials and 0 systematic studies met the inclusion criteria.

**Surgery**

1. **Recommendation: Surgical Interventions for Mallet Finger with Displaced Fractures**

   Surgical treatment with a fixation wire is recommended for patients with displaced fractures involving more than one third to one half of the articular surface of the DIP joint.

   *Strength of Evidence – Recommended, Insufficient Evidence (I)*
Level of Confidence – Moderate

2. **Recommendation: Surgical Interventions for Failed Splinting Cases of Mallet Finger**

   Surgery is recommended for those cases that fail splinting yet have sufficient symptoms or concerns that an attempt at fixation is desired.

   **Strength of Evidence – Recommended, Insufficient Evidence (I)**

   Level of Confidence – Moderate

Rationale for Recommendation

Quality studies to determine which patients with mallet finger would be optimal for surgical interventions are not currently available. (1048) One study reported a non-statistically significant trend suggesting preference for fixation among those presenting late for treatment; (1041) however, the dropout rate was high. A low-quality study also suggested no difference in splinting outcomes among those presenting late. (1042) Surgery is invasive, has relatively few adverse effects for this disorder, and is high cost; however, surgery is recommended for these select patients.

Evidence for the Use of Splints and Surgery for Mallet Finger

There are 5 moderate-quality RCTs incorporated into this analysis. (264, 1041, 1044, 1049, 1050) There are 3 low-quality RCTs in Appendix 2. (1042, 1043, 1051)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: surgical procedure, surgical intervention, surgery, displaced fracture, finger, mallet, baseball, drop, hammer; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 7 articles in PubMed, 75 in Scopus, 0 in CINAHL, 29 in Cochrane Library, 332 in Google Scholar, and 0 from other sources. We considered for inclusion 5 from PubMed, 4 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 9 articles considered for inclusion, 8 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>O'Brien 2011 RCT, Single blind Sponsored by the Alfred Allied Health Research Grant (A10602). No COI.</td>
<td>6.5</td>
<td>N = 64 (42 males, 22 females) with acute type 1a or 1b mallet finger; Mean (±SE) age 37.6 (±1.9).</td>
<td>Dorsal aluminum (13-mm wide padded aluminum) splint group (N= 21) Vs. Thermoplastic splint (1.6mm Orfit classic soft micro-perforated) group (N=22) Vs. Stack splint control group (N=21)</td>
<td>No significant differences reported between groups for extension lag at 8, 10, 12 or 20 weeks. The dorsal splints and stack control group had significantly higher treatment failure rate compared to thermoplastic group: Dorsal split - 23.8% vs. Control - 23.8% vs. Thermoplastic - 0%, (p=0.04).</td>
<td>“Our findings demonstrate that the majority of mallet finger injuries treated with 8 weeks of immobilization and graded exercise thereafter achieve excellent or good results; adding weight to the argument that these injuries can be managed independently in hand therapist-led clinics. To enable patients to comply with this protocol, the splint provided must be robust enough for daily living requirements and must not cause complications which are intolerable to the patient. In this study, there was no significant difference in the outcomes achieved in the 3 trial splints; however, the custom-made thermoplastic splint was significantly less likely to result in complications that lead to treatment failure thus supporting its use in the treatment of mallet finger.”</td>
<td>High dropout rate. Data suggests mallet finger splints should be designed to accommodate daily living and thus enhance compliance as these splints must be worn continuously for 6-8 weeks. Data found no extensor lag difference or pain or satisfaction between the three splints. Stack and dorsal splints had high treatment failure rates.</td>
</tr>
<tr>
<td>Tocco 2013 RCT No sponsorship or COI.</td>
<td>6.0</td>
<td>N = 57 (35 males, 22 females) with closed mallet fingers (60 fingers total); Mean age 45.</td>
<td>Low temperature thermoplastic lever-type orthosis group (LTTP) (N=30; 30 fingers) Vs. Quickcast orthosis group (QC) (N=27; 30 fingers)</td>
<td>At 12 weeks follow up (follow up 5), the LTTP group had significantly higher extensor lag than the QC group, (p=0.05). The QC group had significantly higher average active extensions of 5 degrees or more compared with the LTTP group, (p=0.05). Success rates were higher in the QC group compared with LTTP group and approached significance; 60% vs. 81%, (p=0.08).</td>
<td>“The findings of this study demonstrate that full-time immobilization with QC of Type 1 mallet fingers was more effective than the traditional approach of fabricating an LTTP orthosis and instructing the patient to remove it daily for skin care.”</td>
<td>Relatively small sample size. Compliance difficult to assess. Group instructions were different. Data suggest LTTP group had significantly greater extensor lag than QC subjects at 12 weeks and age and amount of edema negatively impacted D/P extensor lag.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>N &amp; Gender</td>
<td>Mean Age</td>
<td>Treatment Details</td>
<td>Follow-up</td>
<td>Results</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>------------</td>
<td>----------</td>
<td>-------------------</td>
<td>-----------</td>
<td>---------</td>
</tr>
<tr>
<td>Pike 2010</td>
<td>Randomized clinical trial</td>
<td>Sponsored by the Canadian Orthopedic Association. No COI.</td>
<td>5.5</td>
<td>N = 77 (51 males, 26 females) with acute mallet finger; Mean age 43.</td>
<td>Dorsal aluminum (with padding) splint group (N=26) Vs. Volar aluminum splint (without padding) group (N=27) Vs. Custom thermoplastic with circumferential coverage splint group (N=24)</td>
<td>All groups received 6 weeks of treatment. No overnight splinting required.</td>
</tr>
<tr>
<td>Warren 1988</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>5.0</td>
<td>N = 114 (73 males, 41 females) with mallet finger; Mean age 46.1.</td>
<td>Stack splint group (N=58) vs. Abouna splint group (N=49) Splints worn continuously for 6 weeks, then nightly for 2 weeks</td>
<td>Successes: Stack vs. Abouna splint: 19/58 (33%) vs 19/49 (39%) (NS); 20/70 (28.6%) without vs 17/33 (51.5%) with bony injury; Ages 10-39: 23/38 (60.5%); ages 40-79: 15/69 (21.7%)</td>
</tr>
<tr>
<td>Maitra 1993</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>4.0</td>
<td>N = 60 (37 males, 23 females) with mallet finger; Mean age 44.5 ± 16.6.</td>
<td>Aluminum splint group (N=30) vs. Stack splint group (N=30) All splints worn continuously for 6 weeks, then nightly for 3 weeks.</td>
<td>Success rates 37% vs. 33% (NS); skin complications aluminum vs. stack splint: number of fingers with skin complications: 6.6% vs. 33%; dorsal ulcer: 3% vs. 10%; skin maceration: 3% vs. 20%</td>
</tr>
</tbody>
</table>

Data suggests comparable efficacy as no lag differences were observed between the three splint types. Data suggests increased lag occurs after the splint is discontinued.
<table>
<thead>
<tr>
<th>Source</th>
<th>Level</th>
<th>Study Details</th>
<th>Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Auchincloss 1982</td>
<td>4.0</td>
<td>N = 41 (29 males, 22 females) with mallet finger injuries; Mean age 41.</td>
<td>Kirschner wire percutaneous fixation (6 weeks) group (N=19) vs. Pryor and Howard splint (6 weeks) (N=22) Follow up fourteen to eighteen months after injury.</td>
<td>“Trial showed no particular advantage for either method, but suggested that patients presenting after some delay may achieve better results after internal fixation.”</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>K-wire group vs P&amp;H splint group: Normal function: 19/19 (100%) vs. 20/22 (90.9%). Good objective results: 11 (57.9%) vs. 11 (50%). Unchanged objective results: 1 (5.3%) vs. 4 (18.2%)</td>
<td>High dropout rates preclude strong conclusions.</td>
</tr>
<tr>
<td>Toker 2015</td>
<td>4.0</td>
<td>N = 22 (17 males, 5 females) with mallet fractures; Mean age 32.</td>
<td>Extension block pinning group (N = 16) vs Open reduction and hook plate fixation group (N = 6) Mean follow up 13 months.</td>
<td>“Extension block pinning and open reduction hook plate fixation comparable in efficacy. The cost of open reduction and plate fixation was higher than that of extension block pinning. This difference would be even higher if plate removal is required.”</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>No significant differences reported between groups at follow up for VAS, mean extensor lag or mean flexion. Extension block pinning found to be more cost-effective than hook plate fixation.</td>
<td>Small sample (N=22). Data suggest similar efficacy between extensor block pinning versus open reduction for mallet fractures and pinning more cost effective than open reduction.</td>
</tr>
</tbody>
</table>
Flexor Tendon Entrapment (Tenosynovitis and Trigger Digit)

Diagnostic Criteria
The diagnosis of flexor tendon entrapment is clinical. Patients without triggering will typically have only focal A1 pulley tenderness with or without a tendon nodule. Patients with triggering can usually demonstrate the triggering for the examiner. (36, 40)

Special Studies and Diagnostic and Treatment Considerations
There are no special tests that are typically performed. (36, 40) X-rays are usually not helpful. The threshold for testing for confounding conditions such as diabetes mellitus, hypothyroidism and connective tissue disorders should be low particularly to prevent other morbidity. (36, 40)

Evidence for the Use of Diagnostic Studies
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Flexor Tendon Entrapment, Tenosynovitis, Trigger Finger Disorder, X-Rays, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 5 articles in PubMed, 24 in Scopus, 0 in CINAHL, 0 Cochrane Library, and 195 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Initial Care
The initial care for this condition is not well defined. As quality evidence for efficacy of other interventions is lacking and success of injections is strong, arguably, the initial management should be glucocorticosteroid injection (39) in contrast with most other potentially occupational MSDs where non-invasive treatments are initially indicated.

SPLINTS

Recommendation: Splints for Select Cases of Acute, Subacute, or Chronic Flexor Tendon Entrapment
Splints are recommended for treatment of select cases (i.e., patients who decline injection) of acute, subacute, or chronic flexor tendon entrapment.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There is one moderate-quality RCT evaluating the use of two different splints for flexor tendon entrapment with no apparent differences in outcomes between the types of splints. (1052) Historically splints were widely used for treatment of trigger digits; (27, 36, 40, 1053) however, prior to the advent of glucocorticosteroid injection, the lack of successful treatments often resulted in surgery. Splints have been used to treat trigger digits (27, 36) and they may be reasonable intervention for patients who decline injection, although it is recommended that patients be educated that the use of splints appears substantially less successful than injections (or surgery).

Evidence for the Use of Splints
There is 1 moderate-quality RCT incorporated into this analysis. (1052)
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Splints, Flexor Tendon Entrapment, Tenosynovitis, Trigger Finger Disorder, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 20 articles in PubMed, 21 in Scopus, 5 in CINAHL, 1 in Cochrane Library, and 2130 from Google Scholar. We considered for inclusion 1 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 0 from other sources. Of the 3 articles considered for inclusion, 1 randomized trials and 1 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type Conflict of Interest (COI)</th>
<th>Scoring (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tarbhai 2012 RCT Supported by University Health Network Allied Health research fund.</td>
<td>4.0</td>
<td>N = 30 (17 females, 13 males) with trigger digit. Mean age 63.4 years.</td>
<td>MCP Group: metacarpophalangeal joint blocking splint (n = 15, 15 digits) vs. DIP Group: distal interphalangeal joint blocking splint (n = 15, 17 digits). Follow-up 3 and 6 weeks.</td>
<td>At 6 weeks, MCP group 77% success rate vs. 47% in DIP group, and slight decrease in grip strength; 4/13 MCP vs. 3/15 DIP (p &gt;0.05). No identified functional limitations. No significant difference in pain intensity, severity of triggering, frequency of triggering, functional limitations (p &gt;0.05).</td>
<td>“Initiating conservative treatment with the MCP joint blocking splint has value for patients with trigger finger and positive outcomes in 77% of subjects, whereas use of the DIP joint splint was effective in about half of subjects.”</td>
<td>Small sample. Trends towards different severity at baseline in outcome measures. Data suggest increase comfort with MCP joint blocking splint but both groups showed significant improvement at 6 weeks maintained for 1 year. Data do not show substantive differences between types of splints.</td>
</tr>
</tbody>
</table>

**Follow-up Visits**

Follow-up visits are generally limited unless complications arise. Success of injections is usually high, thus 1 or 2 follow-up appointments are typical. Post-surgical outcomes for minimally invasive approaches are similarly excellent and necessitate few, if any follow-up appointments beyond 1 or 2. Additional appointments are required for complicated courses.

**Medications**

There are no quality studies on use of medications for this condition, although some have recommended NSAIDs.(36) Medications are generally not required as the condition is generally not substantially painful. NSAIDs may be a reasonable option to control pain; however, injections appear to be superior interventions. NSAIDs may be a more appropriate intervention for those who decline initial injection.

**Evidence for the Use of Medications**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Flexor Tendon Entrapment, Tenosynovitis, Trigger Finger Disorder, Anti-Inflammatory Agents, Non-Steroidal, non-steroidal anti-inflammatory, NSAIDS; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 12 articles in PubMed, 2 in Scopus, zero in CINAHL, one in Cochrane Library, 5730 in Google Scholar, and zero from other sources. We considered for inclusion 1 from PubMed, zero from Scopus, zero from CINAHL, zero from Cochrane Library, one from Google Scholar, and zero from other sources. Of the articles considered for inclusion, 1 randomized trial and 1 systematic studies met the inclusion criteria.
Physical Methods/Rehabilitation

EXERCISE
Exercise is not generally indicated acutely and most patients with flexor tendon entrapment do not require an exercise program. For patients with residual deficits, particularly post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

Evidence for the Use of Exercise
There are no quality studies incorporated into this analysis. A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: exercise, exercising; flexor tendon entrapment, trigger finger disorder, trigger thumb, trigger digit, thumb, thumbs, digit, digits; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 0 in Scopus, 0 in CINAHL, 1 in Cochrane Library, 12,060 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Injections

GLUCOCORTICOSTEROID INJECTIONS
Recommendation: Glucocorticosteroid Injections for Acute, Subacute, or Chronic Flexor Tendon Entrapment

Glucocorticosteroid injections are strongly recommended for treatment of acute, subacute, or chronic flexor tendon entrapment.

Indications – Triggering digit or symptoms of pain over the A-1 pulley thought to be consistent with stenosing tenosynovitis. Injection at the first appointment may be the most appropriate initial intervention. (39)

Dose – Optimal dose is unknown. Quality studies have included betamethasone 6mg,(38, 1054) depot preparation of methylprednisolone 20mg;(1055) and triamcinolone 1mL(1056) most of which were generally combined with an anesthetic. However, there are no quality comparisons of doses and the need for topical anesthetic is untested in quality studies. Subcutaneous injection over the A-1 pulley appears as efficacious as attempted intrasheath injection.(1057)

Frequency/Duration – A single injection and results evaluated to document improvement. Ultrasound-guidance is not shown to be helpful.(1079) (Cecen 15)

Indications for Discontinuation – If a partial response, consideration should be given to repeating the injection, typically at a modestly higher dose.

Strength of Evidence – Strongly Recommended, Evidence (A)
Level of Confidence – High

Rationale for Recommendation
There are 2 high-quality and 2 moderate-quality studies incorporated into this analysis.(38, 1075-1077) Glucocorticosteroid injection(s) are the most commonly used intervention for trigger digits.(39, 40) Quality studies have reviewed attempts to inject along the tendon, although a moderate-quality study failed to find superior results among the group with an attempt to inject within the sheath(1078) and a low-quality study performed subcutaneous injections that were efficacious.(1080) Further, ultrasound guidance has not been found to improve the results. (Cecen 15) It has been suggested that many injections are performed along, rather than within the sheaths(1081) and thus it may not matter how precisely these injections approximate the target tissue. Multiple studies have consistently demonstrated efficacy of these injections compared with placebo with estimates of success typically exceeding 80%.(31, 38, 1075-1077, 1082-1090) Two studies compared injection with surgery, but the recurrence rates while lower with surgery still showed strong efficacy of injection (0% vs. 11% recurrence(1090) and 0% vs. 14%.(1089) They are less effective in diabetics, although still are effective(38) and there is weak
evidence that patients failing other medical treatments may respond at lower rates of approximately 60%.(280) These injections are minimally invasive, have low adverse effects, and are moderate cost. Some caution is warranted regarding repeated administrations with adverse effects including atrophy that generally recovers over time. Nevertheless, quality studies have documented their efficacy and thus, they are strongly recommended.

**Evidence for the Use of Glucocorticosteroid Injections for Flexor Tendon Entrapment**

There are 2 high-(38, 1075) and 12 moderate-quality RCTs incorporated into this analysis.(1076, 1077, 1079, 1086, 1089-1096) (Jianmongkol 07; Cecen 15)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Glucocorticosteroid injection/ flexor tendon entrapment, trigger finger disorder, trigger thumb, trigger digit, tenosynovitis; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 31 articles in PubMed, 36 in Scopus, 0 in CINAHL, and 0 in Cochrane Library. We considered for inclusion 18 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 18 articles considered for inclusion, 13 randomized trials and 3 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baumgarten 2007</td>
<td>RCT</td>
<td>Sponsored by Orthopaedic Research and Education Foundation (OREF). One or more authors received funding and grants.</td>
<td>9.0</td>
<td>N = 59 (37 female/22 male) with diabetes mellitus. Age nondiabetic/diabetic and diabetic placebo groups: 63.1±11.7/62.9±9.00/61.4±9.12.</td>
<td>1ml (6mg) of betamethasone sodium phosphate/acetate and 0.5ml (5mg) of 1% lidocaine vs 0.5ml (5mg) of 1% lidocaine and 1ml sterile saline</td>
<td>Non-diabetics: 22/29 (75.9%) responded to 1 injection; 6 required 2nd injection; 86% responded to 1 or 2 injections. Diabetics: 11/19 (57.9%) responded to 1 injection; 63.2% to 1 or 2 injections. Results after 2nd injection significant.</td>
<td>“Corticosteroid injections were significantly more effective in the digits of nondiabetic patients than in those of diabetic patients. In patients with diabetes, corticosteroid injections did not decrease the surgery rate or improve symptom relief compared with the placebo.”</td>
<td>Glucocorticosteroids also effective in diabetics, though less effective.</td>
</tr>
<tr>
<td>Murphy J Hand Surg 1995</td>
<td>RCT</td>
<td>No mention of sponsorship or CTO.</td>
<td>8.0</td>
<td>N = 24 (15 female/9 male) with primary TF; average age 56 years.</td>
<td>1mL of celestone (6mg) plus 3mL 1% lidocaine vs. 4mL 1% lidocaine only in the placebo group</td>
<td>At 3-week follow-up: steroid group 10/14 (71.4%) vs. 2/10 (20%) asymptomatic. 4-month follow-up, 9/14 (64.3%) vs. 2/10 (20%) asymptomatic (p &lt;0.05).</td>
<td>“Since the treatment was well tolerated by patients and without complications, it is reasonable to offer steroid and lidocaine injection as the initial treatment for primary TF.”</td>
<td>Modest sample size and intermediate-term follow-up.</td>
</tr>
<tr>
<td>Lambert 1992</td>
<td>RCT</td>
<td></td>
<td>6.0</td>
<td>N = 41(16 males, 25 females)</td>
<td>20mg methylprednisolone acetate plus lignocaine vs. 0.05ml 1% lignocaine injection.</td>
<td>Steroid group success rate 12/20 (60%) vs. 3/16 (18.8%) for placebo (p &lt;0.02).</td>
<td>“Our prospective, controlled, double-blinded trial shows that steroid injection is a satisfactory treatment for trigger finger in 60% of patients. There were no complications and success rate may be even better if repeat injections are used.”</td>
<td>Depot preparation may have unblinded the treating physician.</td>
</tr>
<tr>
<td>Peters-Veluthamaningal 2008</td>
<td>RCT</td>
<td></td>
<td>6.0</td>
<td>N = 50 (22 males, 28 females)</td>
<td>1ml triamcinolonacetonide (TCA) injection vs. 0.9% NaCl.</td>
<td>Immediate reductions in triggering were 13/24 (54.2%) vs. 6/22 (27.3%), p = 0.053. Pain scores significantly improved with TCA (p &lt;0.0005); 80% TCA group improved at 12 months.</td>
<td>“Local injection with triamcinolone-acetonide is effective and safe for treating trigger fingers as compared to placebo injection. The effects of steroid injections last up to 12 months.”</td>
<td>Mean symptoms duration differ at baseline and favored placebo (7 vs. 24 weeks, p = 0.023).</td>
</tr>
<tr>
<td>Study</td>
<td>RCT</td>
<td>No mention of sponsorship. No COI.</td>
<td>Number of patients</td>
<td>Methodology</td>
<td>Treatment</td>
<td>Follow-up Duration</td>
<td>Outcome Measures</td>
<td>Observations</td>
</tr>
<tr>
<td>---------------</td>
<td>------</td>
<td>------------------------------------</td>
<td>--------------------</td>
<td>-------------</td>
<td>--------------------------------------------------</td>
<td>--------------------</td>
<td>----------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Goldfarb 2007</td>
<td>RCT</td>
<td>No mention of study sponsorship. No COI.</td>
<td>7.5</td>
<td>N = 125 (88 trigger finger and 37 de Quervains tenosynovitis). Mean age 59.</td>
<td>Group 1: Injection of steroid, lidocaine, bupivacaine alone (standard injection, acidic pH) (n = 57) vs. Group 2: Injection of steroid, lidocaine, bupivacaine, bicarbonate (balanced injection, neutral pH) (n = 68). Follow-up for 6 weeks.</td>
<td>Both injections provided significantly immediate pain relief reflected in VAS scores (p &lt; 0.001). No significant difference between groups for pre-injection VAS (p = 0.89). Group 2 lower VAS scores than group 1 on each of first 7 days. But, differences in VAS scores between groups only significant at days 5, 6, and 7 (p = 0.4 on each day).</td>
<td>“Patients respond to extra-articular steroid injections with gradual improvement over the course of the first week…A pH-balanced injection did not significantly decrease the risk of a flare reaction.”</td>
<td>Data suggest an extra-articular steroid injection gradually benefits patients over first week with about 1/3 of patients reporting a flare response in the days following the injection. A pH-balanced injection did not significantly decrease risk of flare response.</td>
</tr>
<tr>
<td>Zyluk 2011</td>
<td>RCT</td>
<td>No mention of sponsorship. No COI.</td>
<td>5.5</td>
<td>N= 105 trigger digits in 95 patients (28 males, 67 females) with trigger finger. Mean age was 56 years.</td>
<td>Surgery Group: A1 pulley release (n = 43 patients, 46 digits) vs. Injection Group- Steroid injection of 1ml 2% plain lidocaine (n = 52 patients, 59 digits). Follow-up at 1 and 6 months.</td>
<td>At 1 month, surgery group significantly lower active ROM of fingers vs. injection group: 264 vs. 270 (p &lt;0.05). Also significantly weaker in surgery group: 85% vs. 99% (p &lt;0.05). No significant differences with regards to other parameters. At 6 months 11% recurrence rate in injection group vs. 0% in surgery group (p = 0.005). At 6 months, surgery group showed significantly lower VAS score; 0.4 vs. 1.3 (p &lt;0.05) and significantly worse ROM: 265 vs. 270 (p &lt;0.05) vs. injection group.</td>
<td>“We conclude that percutaneous A1 pulley release is more effective medium-term therapy for trigger digit than steroid injection, because of lower risk of recurrence.”</td>
<td>Data suggest percutaneous A1 pulley release is better than steroid injection for trigger finger due to lower risk of recurrence (11% vs. 0%). Pain (VAS) was 0.4 in pulley release group vs 1.3 in steroid group at 6 months, and ROM varied only 5 degrees.</td>
</tr>
<tr>
<td>Sato 2012</td>
<td>RCT</td>
<td>No mention of sponsorship. No COI.</td>
<td>5.0</td>
<td>N = 137 patients (18 males, 132 females) with 150 trigger fingers. Mean age 54.4 years.</td>
<td>Open: Conventional open surgery of A1 pulley (n = 56) vs. injection: 2ml of methylprednisolone acetate 40mg at site corresponding to A1 pulley (n = 49) vs. Percutaneous: percutaneous release of A1 pulley (n = 45). Follow-up after 1, 2 weeks and 1, 2, 4, and 6 months.</td>
<td>Cure of trigger finger (N); open 56 vs. 1 injection 28 vs. 2 injections 42 vs. percutaneous 45 (p = 0.004). Topical pain (N) open vs. injection vs percutaneous: 1 week 38 vs. 9 vs. 30 (p = 0.000); 2 weeks 36 vs. 9 vs. 30 (p = 0.000); 1 month 22 vs. 5 vs. 15 (p = 0.008); 2, 4, and 6 months (p = NS). Joint pain (N) open vs. injection vs percutaneous: 1 week 17 vs. 3 vs. 13 (p = 0.014); 2 weeks 18 vs. 3 vs. 12 (p = 0.023); 1 month 15 vs. 3 vs. 13 (p = 0.029); 2, 4, and 6 months (p = NS). Total active motion (TAM) average open vs. injection vs. percutaneous: 1 month 176.41 vs. 207.18 vs. 201.76 (p = 0.012); 2 months 184.89 vs. 209.53 vs. 207.78 (p = 0.048); 4 months and 6 months (p = NS).</td>
<td>“The levels of effectiveness of open surgical and percutaneous methods were superior to the conservative method of using CSs based on the cure and reappearance rates of the trigger.”</td>
<td>Data suggests comparable efficacy between percutaneous and open surgery and both invasive techniques were superior injection to treat trigger finger. Yet recurrence rates were 0% (open/percutaneous) vs. 86% 1-2 injections.</td>
</tr>
<tr>
<td>Ring 2008</td>
<td>RCT</td>
<td>No mention of sponsorship. No COI.</td>
<td>4.0</td>
<td>N = 84 (44 males, 40)</td>
<td>Dexamethasone 4mg/ml (n = 40) vs. NS between groups at 6 weeks for average DASH score (p = 0.43) and 3</td>
<td>“Although there were no differences 3 months</td>
<td>Data suggest at three months there were no differences.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Sponsorship</td>
<td>RCT</td>
<td>COI</td>
<td>Patients</td>
<td>Intervention</td>
<td>Outcome</td>
<td>Results</td>
<td>Conclusions</td>
</tr>
<tr>
<td>-------</td>
<td>-------------</td>
<td>-----</td>
<td>-----</td>
<td>----------</td>
<td>-------------</td>
<td>---------</td>
<td>---------</td>
<td>-------------</td>
</tr>
<tr>
<td><strong>Shakeel 2012</strong></td>
<td>No sponsorship. No COI.</td>
<td>RCT</td>
<td>4.0</td>
<td>N = 100 (30 males, 70 females)</td>
<td>N = 50 for Group A, N = 50 for Group B</td>
<td>Triamcinolone 10mg/ml and 12.5mg dexamethasone sodium injection</td>
<td>Follow-up at 3 weeks and 3 months after initial injection.</td>
<td>Mean improvement in Quinzel grading cortisoloid vs. NSAID: 3 weeks 1.8 vs. 0.9 (p = 0.002); 3 months to 0.3 vs. 0.8 (p = 0.002).</td>
</tr>
<tr>
<td><strong>Callegari 2011</strong></td>
<td>Supported by IBSA Institut Biochimique SA, Pambio-Noranco, Switzerland. No COI.</td>
<td>RCT</td>
<td>4.0</td>
<td>N = 30 with ultrasound-confirmed diagnosis of trigger finger; mean age 52.5 years</td>
<td>Group A: ultrasound-guided injection of methylprednisolone acetate (40mg/mL) with 0.8mL lidocaine with 1mL hyaluronic acid 0.8% 10 days later (n = 15) vs. Group B: open surgical release 1st annular pulley (n = 15). Follow-up for 12 months.</td>
<td>At 6 months complete symptom resolution seen in 14/15 (93.3%) in Group A. All 15 in Group B achieved complete resolution of impairment by 3 weeks after surgery, but 10 needed PT to reach complete resolutions of symptoms approximately 30-40 days after surgery. No significant differences between groups for VAS, DASH&lt; and SVAS scores, (p &gt;0.05).</td>
<td>&quot;...the results of this explorative study suggest that ultrasound-guided injection of a corticosteroid and hyaluronic acid could be a safe and feasible approach for the treatment of trigger finger.&quot;</td>
<td></td>
</tr>
<tr>
<td><strong>Pataradool 2011</strong></td>
<td>No mention of sponsorship. No COI.</td>
<td>RCT</td>
<td>4.0</td>
<td>N = 40 with primary trigger fingers; mean age 57.5 years</td>
<td>CI Group: conventional injection technique 0.1% triamcinolone acetone 1ml and 1% lidocaine hydrochloride without epinephrine 1ml (n = 20) vs. P11 Group: Proximal phalanx injection</td>
<td>At final follow-up, mean VAS score 7.3 in CI group vs. 3.2 in P11 group. Difference significantly lower in P11 group (p &lt;0.001). Rise of recurrent symptoms occurred in both groups at 3 month follow-up, 3/20 (15%) in CI group and 5/20 (25%) in P11 group.</td>
<td>&quot;We concluded that the P11 technique is less painful than the CI technique without any significant difference in recurrence rate between the two groups at three months follow-up.&quot;</td>
<td></td>
</tr>
</tbody>
</table>

Data suggest steroids act faster, but at 3 months, NSAIDs equally effective.
<table>
<thead>
<tr>
<th>Study</th>
<th>Phase</th>
<th>No. of patients</th>
<th>Gender Distribution</th>
<th>Intervention</th>
<th>Comparator</th>
<th>Follow-up</th>
<th>Results</th>
<th>Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Taras 1998 RCT</td>
<td>6.0</td>
<td>N = 95 patients (37 males, 58 females) with 107 trigger digits</td>
<td>Intrasheath glucocorticosteroid injection vs. subcutaneous injection along sheath (betamethasone acetate suspension 6mg with 0.5mL 1% lidocaine with Omnipaque)</td>
<td>Intrasheath complete in 19/55 (37%), 24/55 (46%) partial, 9/55 (17%) no evidence of intrasheath injection. Intrasheath group overall: 27/52 (52%) good, 10/52 (19%) fair, 15/52 (29%) poor results. Complete intrasheath injection 47% good, 16% fair, 37% poor. Partial injection 50% good, 17% fair, 33% poor. Subcutaneous injection 70% good, 11% fair, 19% poor. Subcutaneous group: 39/55 (71%) good, 4/55 (7%) fair, 12/55 (22%) poor results.</td>
<td>The results of this study suggest that true intrasheath injection offers no apparent advantage over subcutaneous injection in the treatment of trigger digits.</td>
<td>Evidence suggests subcutaneous injections usually not completely successful. Excluded those &gt;6 months duration, and diabetes.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jianmongkol 2007 RCT</td>
<td>4.5</td>
<td>N=103 trigger fingers (14 males, 87 females) CI Technique mean age was 53.68, MAI Technique mean age was 52.19.</td>
<td>Conventional technique of injection (CI technique) N=53, 23 thumb, 5 index, 13 long, 12 ring Vs. Mid-Axial injection technique (MAI technique) N=48, 21 thumb, 4 index, 15 long, 8 ring Follow-up at 1, 3, and 6 weeks.</td>
<td>After insertion of injections the mean pain score for the MAI technique group was 40.19 and the mean pain score for the CI technique group was 48.39. No statistical significance in mean paracetamal count during follow up periods for both groups. Chi-squared test revealed a score of 5.7 with statistical significance of p&lt;0.05 for both groups. In CI technique group, two fingers had recurrent symptoms and no recurrent symptoms in the MAI technique group.</td>
<td>“In conclusion, the injection technique by Carlson and Curtis’ approach can provide the good results of treatment and there were no complications from the injection. The technique can be easily used and safe for injection in the primary trigger finger.”</td>
<td>Sparse methodological details. Baseline comparability is questionable. Data suggest MAI technique had less reported post injection pain associated with procedure compared to CI technique. But at 6 weeks there were no differences in reported VAS pain scores.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cecen 2015</td>
<td>4.5</td>
<td>N=74 patients (15 males, 55 females) with persistent or increasing symptoms of a single trigger digit. Blinded group average age 54.31±10.96. Ultrasound group average age was 55.60±11.63.</td>
<td>Ultrasound-guided group (USG) N=37 Vs. Blinded group (BIG) N=37 Follow up at 6 weeks and 6 months.</td>
<td>Of the 35 patients treated in BIG, 29 responded to a single corticosteroid injection and 6 required a second injection. Thirty-two of the 35 USG group responded to a single injection and 3 required a second injection. There was no significant difference between BIG and USG for requirement of second injection (p&gt;0.05). Both groups showed significant improvement in Quinell grades, but no significant improvement between the pre-steroid, post-steroid 6 week, and post-steroid 6 month scores. Both groups showed significant</td>
<td>“In conclusion, corticosteroid injection can be recommended as a sound, low-risk primary treatment option for trigger finger, which can be given in an office setting, as a low-cost procedure. The use of ultrasound-guided injection of corticosteroid may be associated with extra time and higher rate of diabetics in BIG group and more females in USG group. Data suggest non superiority of US guided injections (USG) versus blinded injections (BIG).</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Intrasheath Glucocorticosteroid Injection vs. Subcutaneous Injection**

**Ultrasound-Guided injection**
| improvement in VAS scores. BIG VAS | scores decreased from 4.80 to 1.5 at 6 weeks and 0.5 after 6 months. USG VAS scores decreased from 4.7 to 1.6 at 6 weeks and to 0.5 after 6 months. VAS in each group showed significant reduction. (p<0.01). | effort, and seems to have no superior clinical benefits compared to the blinded technique. |
Surgery

Recommendation: Surgery for Persistent or Chronic Flexor Tendon Entrapment

Open release for persistent or chronic flexor tendon entrapment is moderately recommended. Percutaneous release is also a reasonable option.(1074)

Indications – Triggering digit or symptoms of stenosing tenosynovitis that has been unresponsive to at least 1 glucocorticosteroid injection, or with an inadequate response. Those without any response should be evaluated carefully for possible alternate conditions. Adjunctive surgical treatment with glucocorticosteroid injection could be considered, although that evidence relies on a single moderate-quality study.(1075)

Strength of Evidence – Moderately Recommended, Evidence (B)
Level of Confidence – High

Rationale for Recommendation
Both open (with a scalpel) and percutaneous (with a needle through the skin) releases are performed with evidence both are effective.(1099) Evidence is strong that percutaneous release is as effective, if not more effective than open release,(271, 1089, 1090, 1097, 1100-1105) (Gilberts 01) is faster to perform, requires fewer resources,(1097, 1104) involves less pain, and results in faster recovery.(1097) Failures are believed to be due to incomplete release of the A-1 pulley.(1106) There are concerns, however, that particularly in inexperienced hands, lacerations of digital nerves, arteries, and other structures and other complications have been reported with the percutaneous technique. The thumb appears more prone to these complications. A moderate-quality study attempted to identify which third of the pulley was responsible for triggering; however, failures occurred in all surgical groups regardless of which third of the pulley was released.(1106) A low-quality case series suggested repeat percutaneous release was reasonable for treatment of incomplete releases,(1107) although open release has been favored for percutaneous failures. One moderate-quality study compared injections with percutaneous release combined with glucocorticosteroid injection and reported surgical release was superior,(1098) although the success rates were both lower than other reports. Surgical release is invasive (though less invasive with percutaneous release),(1108) has low adverse effects, but is costly. For those patients failing glucocorticosteroid injection(s), surgery is recommended.

Evidence for Surgery for Flexor Tendon Entrapment
There are 10 moderate-quality RCTs incorporated into this analysis.(1090, 1091, 1097, 1098, 1102, 1103, 1106, 1108-1110) (Pegoli 08)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: flexor tendon entrapment, trigger finger disorder, trigger thumb, trigger digit, tenosynovitis Surgery, Open release surgery, percutaneous release surgery; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 147 articles in PubMed, 13 in Scopus, 8 in CINAHL, 23 in Cochrane Library, 570 in Google Scholar, and 3 from other sources. We considered for inclusion 5 from PubMed, 0 from Scopus, 0 from CINAHL, 1 from Cochrane Library, 1 Google Scholar, and 3 from other sources. Of the 10 articles considered for inclusion, 10 randomized trial and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maneerit 2003</td>
<td>5.5</td>
<td>N = 115 patients with idiopathic trigger thumbs</td>
<td>Percutaneous release with steroid injection vs. steroid injection alone.</td>
<td>Surgical results satisfactory in 59/65 (90.8%) treated surgically vs. 28/60 (46.7%) treated with injection, p = 0.001. No significant differences in pain ratings or paracetamol tablets required post-procedure. After 2nd injection, success rate 56.7% for injections.</td>
<td>&quot;We conclude that percutaneous trigger thumb release combined with steroid injection has a higher success rate than that of steroid injection alone.&quot;</td>
<td>Success rates, especially in injection arm, low compared with other quality evidence raising questions about subject selection/other issues. No mention of gender.</td>
</tr>
<tr>
<td>Gilberts 2001</td>
<td>5.5</td>
<td>N = 96 patients (56 males, 44 females)</td>
<td>Open surgery vs. percutaneous surgery.</td>
<td>Open vs. percutaneous release – Operative time 11 vs. 7 minutes, p &lt;0.0001. Mean post-op pain 5.7 vs. 3.1 days, p = 0.039. Motor recovery 18 vs. 7 days, p &lt;0.002. Return to work 7.5 vs. 3.9 days, p &lt;0.001. Complications 3 vs. 2. Success rate 98 vs. 100%, NS</td>
<td>&quot;We conclude that percutaneous correction of trigger digits is a quicker procedure, is less painful, and shows significantly better results in rehabilitation than open surgery.&quot;</td>
<td>All measures favored percutaneous release. Discrepancy with patient number and gender.</td>
</tr>
<tr>
<td>Bamroongshawg asame 2010 RCT</td>
<td>4.5</td>
<td>N= 142 patients (58 males, 84 females) with 160 trigger fingers. Mean age was 47.4 years.</td>
<td>Open Group: Open release surgery (n = 70/ 80 digits) vs. Percutaneous group (n = 72/80 digits). Follow-up 3 and 6 weeks.</td>
<td>Mean time of open surgery 2.2 minutes; percutaneous 1.8 minutes (p= 0.05). Post-op patient satisfaction scores similar at weeks 3 and weeks 6 (p&gt; 0.05). Percutaneous surgery group had lower mean pain score vs. open group at weeks 1, 2, 3 and 4.</td>
<td>&quot;Percutaneous trigger digit surgery using the full handle knife 45° is effective and safe, and results functional outcomes equal to those with open trigger digit surgery.&quot;</td>
<td>Data suggest comparable efficacy between open and percutaneous release in trigger digits</td>
</tr>
<tr>
<td>Dierks 2008 RCT</td>
<td>4.5</td>
<td>N=36 patients (16 males, 20 females) with trigger finger. Mean age was 62.9 years</td>
<td>Open group- Open surgical release of the A-1 pulley (n = 16) vs. Percutaneous group- percutaneous release of A-1 pulley (n = 20). Follow-up at 1 and 12 weeks.</td>
<td>Both groups showed decrease in pain level, but no significant difference between groups (p &gt;0.05). Mean surgery time 26 s in percutaneous group; 4 minutes 17 s open group (p &lt;0.05). Active ROM of PIP joint significantly lower open group at 1 week; 81 vs. 95 (p &lt;0.05). No significant difference for ROM at 12 weeks.</td>
<td>&quot;Because of lower costs and quicker procedure with equal functional outcome when compared with open surgery, we recommend the percutaneous technique using a L15 blade for trigger finger release.&quot;</td>
<td>Sparse methodological details. Data suggest percutaneous release of A1-pulley for stenosing tendovaginitis as it is quicker, less costly and has comparable efficacy to surgery.</td>
</tr>
</tbody>
</table>
| Pegoli 2008 RCT                               | 4.0          | N=200 patients (60 males, 140 females) | Group A-open surgical release of the A-1 pulley (N=100) Vs. | Three patients in Group A reported dyesthesia for 10 days that resolved and 8 patients from Group B reported dyesthesia for 6 days that resolved. The sum of excellent and good results | "The main complaint of the patients after an open trigger finger release is a discomfort at the incision site. In this prospective study. | Sparse methodological details. Data suggest the endoscopic procedure showed faster recovery at all times of evaluation (7
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>N</th>
<th>Gender</th>
<th>Procedure Description</th>
<th>Results</th>
<th>Conclusion</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>with a trigger finger. Mean age was 58.5 years.</td>
<td>Mean age was 58.5 years. A higher difference in results was observed at 30 days post-operation. Group B showed faster recovery. Aesthetic appearance of incision site had significant statistical analysis (p&lt;0.001) with a variable percentage of 30% between the groups and pain under load (p&lt;0.017)</td>
<td>we compared the two consecutive groups of patients with trigger fingers. One was treated by an open approach and the other by the endoscopic release of the A1 pulley. Pre- and post-operative evaluation at seven, 30 and 90 days showed a faster recovery from the discomfort with a faster return to daily and working activities, after the endoscopic procedure.</td>
<td>days, 30 days &amp; 90 days compared to open procedure although surgical times for both procedures are similar.</td>
</tr>
<tr>
<td>Topper 1997</td>
<td>4.5</td>
<td></td>
<td></td>
<td></td>
<td>Group B-endoscopic surgical release of the A-1 pulley (N=100) Follow-up pre-operatively and at 7, 30, and 90 days post-operatively. (questionnaire) at 90 days post-operation was similar for both groups with a prevalence of excellent results in Group B. A higher difference in results was observed at 30 days post-operation. Group B showed faster recovery. Aesthetic appearance of incision site had significant statistical analysis (p&lt;0.001) with a variable percentage of 30% between the groups and pain under load (p&lt;0.017)</td>
<td>&quot;In all 19 patients, a partial resection of the first annular pulley resulted in continued clinical triggering with active digital flexion. At this point, a standard complete first annular pulley release was performed, with resolution of clinical triggering of the involved digit in all patients.&quot;</td>
<td>&quot;We conclude that there is no &quot;critical third&quot; of the first annular pulley responsible for clinical digital triggering.&quot;</td>
<td>Suggests release of the entire pulley is preferred treatment. No mention of gender.</td>
</tr>
<tr>
<td>Sectioning Different Thirds of the A1 Pulley</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Topper 1997</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yiannakopoulos 2006</td>
<td>5.5</td>
<td>RCT</td>
<td>50</td>
<td>20 males, 28 females</td>
<td>N = 50 patients (20 males, 28 females) with trigger finger syndrome EMLA anaesthesia vs 3ml lidocaine 1% Visual analogue pain scale EMLA vs Lidocaine: VAPS: 0 vs. 5.96±2.41 (p&lt;0.05); Patient Satisfaction: 4.6±0.2 vs. 4.4±0.3 (NS)</td>
<td>&quot;Percutaneous trigger finger release can be performed as an office procedure with the use of EMLA avoiding the use of injectable local infiltration anaesthesia.&quot;</td>
<td>EMLA requires 2-3 hours for effectiveness potentially resulting in NS satisfaction scores despite marked differences in pain scores. *The number of males and females compared to the groups does not add up.</td>
<td></td>
</tr>
<tr>
<td>Topical Anesthesia vs. Lidocaine Injection</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zyluk 2011</td>
<td>5.5</td>
<td>RCT</td>
<td>105</td>
<td></td>
<td>N = 105 triggers in 95 patients (28 males, 67 females) with trigger finger. Mean age was 56 years. Surgery Group- A1 pulley release (n = 43, 46 digits) vs. Injection Group- Steroid injection of 1ml 2% plain lidocaine (n = 52 patients, 59 digits). Follow-up at 1 and 6 months. At 1 month, surgery group significantly lower active range of motion of fingers vs. injection group: 264 vs. 270 (p&lt;0.05). Also significantly weaker group in surgery group: 85% vs. 99% (p&lt;0.05). No significant differences with regards to other parameters. At 6 months, 11% recurrence rate in injection group vs. 0% in surgery group (p = 0.005). At 6 months surgery group showed significantly lower VAS score: 0.4 vs. 1.3 (p&lt;0.05) and significantly worse range of motion; 265 vs. 270 (p&lt;0.05) vs. injection group.</td>
<td>&quot;We conclude that percutaneous A1 pulley release is more effective medium-term therapy for trigger digit than steroid injection, because of lower risk of recurrence.&quot;</td>
<td>Data suggest percutaneous A1 pulley release is better than steroid injection for trigger finger due to study suggesting a lower risk of recurrence. Pain (VAS) 0.4 in pulley release group vs 1.3 in steroid group at 6 months, and ROM varied only 5 degrees.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>Participants</td>
<td>Intervention</td>
<td>Outcomes</td>
<td>Comments</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>--------</td>
<td>--------------</td>
<td>-------------</td>
<td>----------</td>
<td>----------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chao 2009</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>4.5</td>
<td>N= 83 patients (26 males, 57 females,) with 93 trigger thumbs. Mean age was 48.5 years.</td>
<td>Group A: miniscalpel-needle percutaneous release (n = 41, 46 thumbs) vs. Group B: Steroid injection 1ml triamcinolone acetonide (10 mg/ml) injected (n = 42, 47 thumbs). Follow-up at 1 and 12 months</td>
<td>Group A achieved successful release in 93% at 1 month and 86% at 12 months. 45% of thumbs in group B satisfactory at 1 month and 26% were satisfactory at 12 months. The mean percent decrease in pain intensity was significantly higher in group A vs. group B at 1 month; 65.7% vs. 38.4% (p &lt;0.001) and 12 months; 89.4% vs. 6.8% (p &lt;0.001).</td>
<td>“Percutaneous release with a miniscalpel-needle had a higher success rate than steroid injection.”</td>
<td></td>
</tr>
<tr>
<td>Callegari 2011</td>
<td>RCT</td>
<td>Supported by IBSA Institut Biochimique SA, Pambio-Noranco, Switzerland. No COI.</td>
<td>4.0</td>
<td>N = 30 patients (20 females, 10 males) with ultrasound-confirmed diagnosis of trigger finger. Mean age was 52.5 years.</td>
<td>Group A: ultrasound-guided injection of methylprednisolone acetate (40mg/mL) with 0.8mL lidocaine with 1mL hyaluronic acid 0.8% 10 days later (n = 15) vs. Group B: Open surgical release of first annular pulley (n = 15). Follow-up for 12 months.</td>
<td>At 6 months complete symptom resolution observed in 14/15 (93.3%) in group A. All 15 in group B achieved complete resolution of impairment by 3 weeks after surgery, but 10 needed physical therapy to reach complete resolutions of symptoms approximately 30-40 days after surgery. No significant differences between groups for VAS, DASH&lt; and SVAS scores (p &gt;0.05).</td>
<td>“…the results of this explorative study suggest that ultrasound-guided injection of a corticosteroid and hyaluronic acid could be a safe and feasible approach for the treatment of trigger finger.”</td>
<td></td>
</tr>
</tbody>
</table>

Data suggest percutaneous release via miniscalpel-needle had better efficacy than steroid injection.
Extensor Compartment Tenosynovitis (Including de Quervain’s Stenosing Tenosynovitis and Intersection Syndrome)

Diagnostic Criteria
These diagnoses are clinical. Patients without triggering will typically have tenderness that is focal over the affected tendon(s) or compartment. Finkelstein’s maneuver should be positive.

Special Studies and Diagnostic and Treatment Considerations
There are no special tests that are typically performed for extensor compartment tenosynovitis. X-rays are usually not helpful. The threshold for testing for confounding conditions such as diabetes mellitus and hypothyroidism should be low, particularly to prevent other morbidity. There are reports of MRI findings;(1086-1088) however, the utility of MRI has not been demonstrated in quality studies. Distinguish from de Quervain’s.

There is 1 moderate-quality study incorporated into this analysis.(1111) (Chien 01)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: X-Rays, Tomography Scanners, X-Ray Computed, Extensor Compartment Tenosynovitis, De Quervain's Stenosing Tenosynovitis; diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 13 articles in PubMed, 7 in Scopus, 1 in CINAHL, 0 in Cochrane Library, and 393 from Google Scholar. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, from Google Scholar, 0 from Cochrane Library and 0 from other sources. Of the 1 articles considered for inclusion, 1 diagnostic study met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score</th>
<th>Number</th>
<th>Area of Spine</th>
<th>Diagnoses</th>
<th>Type of X-rays</th>
<th>CT Used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Surgery Performed</th>
<th>Clinical outcomes assessed</th>
<th>Long term follow-up</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
</table>

Copyright© 2016 Reed Group, Ltd.
6.5

N = 45, (11 Men (24%), 34 Women (76%)) with de Quervain tenosynovitis. Mean age, 43 years.

Wrist de Quervain tenosynovitis confirmed

Not given

- - ++ - + - -

The association between focal radial styloid abnormality and de Quervain tenosynovitis, for both observers, (p < 0.05). The areas under the receiver operating characteristic curves for both observer: 0.71 (95% CI, 0.62–0.79%) and 0.76 (95% CI, 0.67–0.84%). The Kappa values for inter observer variability = 0.44 (moderate agreement), and intra observer variability = 0.62 (substantial).

"Focal radial styloid abnormality is an indicator of de Quervain stenosing tenosynovitis of the wrist."

A retrospective review of radiography showed that focal radial steroid abnormalities to be an indicator of de Quervain stenosing tenosynovitis.

**Recommendation: MRI to Diagnose Extensor Compartment Tenosynovitis**

MRI is recommended to diagnose extensor compartment tenosynovitis.

*Strength of Evidence – Recommended*

*Level of Confidence – Low*

**Rationale for Recommendation**

There are 2 moderate-quality articles evaluating the use of MRIs to diagnose extensor compartment tenosynovitis.

**Evidence for the Use of MRI**

There are 2 moderate-quality studies incorporated into this analysis. (1115, 1116) There is 1 low-quality study in the Appendix 2. (1117) (Hadidy 09)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: MRI OR Magnetic Resonance Imaging Extensor Compartment Tenosynovitis, De Quervain's Stenosing Tenosynovitis, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 12 articles in PubMed, 60 in Scopus, 0 in CINAHL, and 0 in Cochrane Library, and 1020 from Google Scholar. We considered for inclusion 2 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 3 articles considered for inclusion, 3 diagnostic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Area</th>
<th>Diagnoses</th>
<th>Type of MRI used</th>
<th>Type of CT used</th>
<th>More than one rater</th>
<th>Surgery performed</th>
<th>Long term follow-up</th>
<th>Clinical outcomes</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nieuwenhuis 2015</td>
<td>Diagnostics</td>
<td>Wrist</td>
<td>RA</td>
<td>1.5T</td>
<td>N/A</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>65% had MRI-detected tenosynovitis. RA patients had tenosynovitis vs. non-RA patients, ( p = 0.023 ). Flexor tendons at MCP5/ extensor tendons at MCP2 and MCP4 in extensor compartment I of wrist affected in RA vs non-RA; 2.8, 95% CI: 1.9-42.8/14.2, 95% CI: 1.7 – 115.9 and 4.0, 95% CI: 1.4 – 11.1.</td>
</tr>
<tr>
<td></td>
<td>Sponsored by EU Seventh Framework and DAF. Drs. Nieuwenhuis, Krabben, and van der Helm-van Mil's supported by DAF and Vidi grant, and Drs. Stomp and Reijnierse's sponsored by TRACER project grant. Dr. Stomp received speaking fees from GE healthcare.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parellada 2007</td>
<td>Diagnostics</td>
<td>Wrist</td>
<td>Tenosynovitis</td>
<td>1.5T scanner</td>
<td>N/A</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>5 signs of tenosynovitis/4 had tendons of 2nd and 3rd extensor compartment affected/5 had signs of tenosynovitis of ELP tendon/3 showed tenosynovitis proximal and distal to point of intersection; 2 of 3 had discrete point of intersection.</td>
</tr>
</tbody>
</table>
Initial Care

Initial care usually involves limitation of the physical factors thought to be contributing. (42) Thumb spica splints for de Quervain’s and wrist braces for the other compartment tendinoses are generally believed to be helpful. (42) Thumb spica splints have been widely used for treatment of wrist compartment tendinoses while non-spica wrist splints have been used for treatment of other compartment tendinoses. (42, 43, 45, 349, 1091) NSAIDs are often prescribed for initial treatment. (42) Perhaps the largest question in the management of these conditions is how soon to inject, including whether patients might not be most effectively treated by injection at initial clinical presentation, however, there are no quality studies to address that question.

SPLINTS

**Recommendation: Thumb Spica and Wrist Splints for Acute and Subacute Thumb Extensor Compartment Tenosynovitis**

**Thumb spica splints for treatment of acute and subacute thumb extensor compartment tendinoses, and non-spica wrist splints for treatment of other extensor compartment tendinoses** (42) are recommended.

**Indications** – Patients with extensor compartment tendinoses.

**Frequency/Duration** – Generally recommended to be worn while awake.

**Indications for Discontinuation** – Failure to respond or resolution.

**Strength of Evidence** – **Recommended, Insufficient Evidence (I)**

**Level of Confidence** – Low

**Rationale for Recommendation**

There is one moderate-quality RCT evaluating wrist splints for extensor compartment tenosynovitis with full-time compared with PRN use and found no differences. (1092) Wrist splints are not invasive, have few adverse effects, and are not costly; thus, they are recommended.

**Evidence for the Use of Splints**

There are 3 moderate-quality RCT incorporated into this analysis. (1119-1121) (Mardani-Kivi 14; Mehdinasab 10)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Splinting, thumb spica, Extensor Compartment Tenosynovitis (Including De Quervain’s Stenosing Tenosynovitis and Intersection Syndrome); controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 7 articles in PubMed, 3 in Scopus, 3 in CINAHL, 295 from Google Scholar, and 51 in Cochrane Library. We considered for inclusion 3 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 2 from other sources. Of the 359 articles considered for inclusion, 3 randomized trials and 6 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Menendez 2015</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>5.0</td>
<td>N = 83 (49 females, 9 males in final randomization) with clinically diagnosed extensor compartment</td>
<td>Forearm-based thumb spica splint to be worn full-time (n = 43) vs. forearm-based thumb spica splint to be worn as desired (n = 40).</td>
<td>No significant differences reported between full-time and as-desired groups for grip strength, pain.</td>
<td>&quot;Our study supports the following concepts: (1) there is no difference in patient-reported outcomes and grip strength with full-time and as-desired splinting, and patients can wear the splint as they choose. High dropout rate in full-time splinting group. Data suggest strict splint vs. selective splint wear to treat de Quervain Syndrome.&quot;</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Duration</td>
<td>Sample Size</td>
<td>Description</td>
<td>Intervention</td>
<td>Comparison</td>
<td>Outcomes</td>
<td>Comments</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>----------</td>
<td>-------------</td>
<td>-------------</td>
<td>--------------</td>
<td>------------</td>
<td>----------</td>
<td>----------</td>
<td></td>
</tr>
<tr>
<td>Mardani-Kivi 2014</td>
<td>Randomized prospective trial</td>
<td>4.0</td>
<td>N = 67 patients (12 males, 47 females) with extensor compartment tenosynovitis, or de Quervain tendinopathy, radial pain of the wrist, a positive Finkelstein test, tenderness of the first dorsal compartment and a pain score &gt;6</td>
<td>Both groups received allocated treatment for 6 weeks. Follow-up at 6 weeks.</td>
<td>At 3 weeks and 6 months follow-up, CSI+TSC group had significantly higher percentages of success compared to TSC alone group: 3 weeks - 97% vs. 76%, (p = 0.027), 6 months - 93% vs 69%, (p = 0.021). At 6 months follow-up, CSI+TSC group had significantly higher percentages of decreased VAS scores vs. CSI-only group: 96% vs. 80%, (p &lt;0.001). At 6 months, CSI+TSC group significantly higher mean (±SD) reduction of QuickDASH score vs. CSI only group: 74 (±15) vs. 66 (±18), (p &lt;0.001).</td>
<td>At 3 weeks and 6 months follow-up, CSI+TSC group had significantly higher percentages of success compared to TSC alone group: 3 weeks - 97% vs. 76%, (p = 0.027), 6 months - 93% vs 69%, (p = 0.021). At 6 months follow-up, CSI+TSC group had significantly higher percentages of decreased VAS scores vs. CSI-only group: 96% vs. 80%, (p &lt;0.001). At 6 months, CSI+TSC group significantly higher mean (±SD) reduction of QuickDASH score vs. CSI only group: 74 (±15) vs. 66 (±18), (p &lt;0.001).</td>
<td>The results of this study indicated that the CSI + TSC treatment method was superior to CSI alone with regards to success rate and functional outcomes.</td>
<td></td>
</tr>
<tr>
<td>Mehdinanasab 2010</td>
<td>RCT</td>
<td>4.0</td>
<td>N= 73 patients (9 males, 64 females) with de Quervain’s tenosynovitis. Mean age was 32.6 years.</td>
<td>Injection Group: Injection of methylprednisolone acetate in first dorsal compartment of wrist followed by wrist thumb spica cast (n = 37) vs. Casting Group: Casting only (n =</td>
<td>Overall success rate at final follow-up (6 months) 86.4% in injection group and 36% in casting group. Difference significant (p &lt;0.001) with regards to final</td>
<td>&quot;Support of the wrist with casting alone had less favorable outcome in de Quervain’s tenosynovitis. Adding methylprednisolone acetate injection into the rst dorsal compartment of the wrist is necessary for more optimal results.&quot;</td>
<td>Differences in success percentages at follow up due to dropout. Data suggest a combination of spica casting and corticosteroid injection was superior to injection alone.</td>
<td></td>
</tr>
</tbody>
</table>

† tenosynovitis, or de Quervain tendinopathy. Mean (±SD) age 50 (±13) for full-time group and 50 (±15) for as-desired group.

Both groups received allocated treatment for 6 weeks. Follow-up at 6 weeks.

Intensity, disability and satisfaction with treatment.

“Support of the wrist with casting alone had less favorable outcome in de Quervain’s tenosynovitis. Adding methylprednisolone acetate injection into the rst dorsal compartment of the wrist is necessary for more optimal results.”

Data suggest casting the wrist plus methylprednisolone injections was beneficial in the treatment of de Quervain’s tenosynovitis over casting alone measured by improvement in functional outcomes.
Follow-up visits are generally required every 1 or 2 weeks to evaluate efficacy of interventions until resolution of the condition.

**Medications**

There are few quality studies on use of medications for this condition, although they are frequently prescribed. Medications are more frequently needed compared with trigger digits, as these conditions are typically more painful.

**NSAIDs**

*Recommendation: NSAIDs for Acute, Subacute, or Chronic Extensor Compartment Tenosynovitis*

NSAIDs (oral or topical) are recommended to control pain associated with acute, subacute, or chronic extensor compartment tenosynovitides.

*Indications* – Patients with wrist compartment tendinoses.

*Dose* – Optimal dose is unknown and there are no quality studies comparing different NSAIDs. Regularly scheduled dosing is recommended for acute, significantly symptomatic presentations.

*Indications for Discontinuation* – Failure to respond, development of adverse effects, resolution.

*Strength of Evidence* – **Recommended, Evidence (C)**

*Level of Confidence* – Low

**Rationale for Recommendation**

NSAIDs are often used to treat pain associated with wrist compartment tendinoses.(42, 43, 45, 349, 1091, 1093, 1094) There is one quality study demonstrating efficacy of a ketoprofen patch versus placebo.(1094) However, another study failed to demonstrate efficacy of injectable nimesulide as an adjuvant treatment to triamcinolone acetonide 10mg injection(1093) and another study of diclofenac gel for treating marathon kayakers prior to racing also was negative,(1095) although applicability to occupational populations is questionable. As an NSAID patch has been demonstrated to be efficacious compared to placebo, it is assumed that other topical forms are also efficacious. NSAIDs are not invasive, have low adverse effects in employed populations, and are low cost, thus they are recommended.

**Evidence for the Use of NSAIDs for Extensor Compartment Tenosynovitis**

There are 2 high-(1093, 1094) and 1 moderate-quality(1095) RCTs incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Extensor Compartment Tenosynovitis, De Quervain Disease, De Quervain Stenosing Tenosynovitis, Intersection Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 3 articles in PubMed, zero in Scopus, 2 in CINAHL, zero in...
Cochrane Library, 163 in Google Scholar, and zero from other sources. We considered for inclusion 3 from PubMed, zero from Scopus, zero from CINAHL, and zero from Cochrane Library, zero Google Scholar, and zero from other sources. Of the 2 articles considered for inclusion, 3 randomized trials and zero systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Diclofenac Gel vs. Placebo</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>May 2007 RCT</td>
<td>6.5</td>
<td>N = 42 (36 males/ 6 females) with Kayakers in 5-day marathon. Mean age: 36±12 years</td>
<td>Diclofenac 2.5g 1% gel vs. placebo gel applied 3 times before each day's race. All received ice, massage, stretches, night bandage.</td>
<td>Pain higher on diclofenac than placebo gel especially in days 2 and 3. Comparisons with day 1: 2 (1.7), 3 (0.5), 4 (-0.1), 5 (-0.9).</td>
<td>&quot;[S]tandard treatment appears to be sufficient for the management of wrist extensor tenosynovitis during competition.&quot;</td>
<td>Applications from kayaking marathon to occupational settings unclear. May be more analogous to acute, unaccustomed forceful use. Applications not throughout day may limit conclusions.</td>
</tr>
<tr>
<td><strong>NSAIDs vs. Placebo</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mazieres 2005 RCT</td>
<td>10.0</td>
<td>N = 172 (98 female/74 male) with tendinitis of upper or lower limbs. Age 18-70 years.</td>
<td>Ketoprofen patch (n = 87) vs placebo (n = 85).</td>
<td>Changes from baseline in pain on daily activity (100mm VAS) in ketoprofen vs. placebo: D0: 69.1±12.9 vs 70.1±11.5 p = 0.5876; D3: 48.6±23.2 vs. 56.1±20.0 p = 0.0491; D7±1: 30.8±23.8 vs. 44.3±25.6 p = 0.0013; D14±2: 25.1±25.9 vs. 36.4±27.6 p = 0.0146.</td>
<td>&quot;This trial suggested that a 3-14 day course of treatment by ketoprofen patch is useful in nonarticular rheumatisms, the duration of treatment depending on the results obtained.&quot;</td>
<td>Many diagnoses included and results not stratified by diagnosis.</td>
</tr>
<tr>
<td><strong>Injection with vs. without NSAID</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jirarattanaphochai 2004 RCT</td>
<td>9.0</td>
<td>N = 160 (144 female/16 male) with de Quervain disease, positive Finkelstein test, radial styloid tenderness, pain on first extensor compartmen t with thumb abduction or extension. Mean (±SD) age 48.98 (±9.10) for nimesulide group: 46.87</td>
<td>Injection 10mg of triamcinolone acetonide and 0.5mL of 1% lidocaine and either 200mg daily oral nimesulide group (n = 80) vs. placebo control group (n = 80). Follow-up at 1 week, 6, 12, 18 and 24 months.</td>
<td>No significant differences reported between the nimesulide and placebo groups for VAS pain scores, success rates, adverse reactions and probability of recurrence.</td>
<td>&quot;[S]teroid injection alone was safe and effective in the treatment of de Quervain’s disease, but the oral administration of nimesulide did not provide any additional benefit beyond that of the injection.&quot;</td>
<td>Data suggest nimesulide does not enhance effectiveness of a single triamcinolone injection in de Quervain’s disease treatment. Disease recurrence was correlated to the presence of crepitation in the first dorsal compartment at thumb extensor abduction.</td>
</tr>
</tbody>
</table>
**Physical Methods/Rehabilitation**

**EXERCISE**

Exercise is not generally indicated acutely and most patients with extensor tendon entrapment do not require an exercise program. For those with residual deficits, particularly post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

*Evidence for the Use of Exercise*

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following term Exercise, Physical Activity, Extensor Compartment Tenosynovitis, De Quervain Disease, De Quervain's Stenosing Tenosynovitis, Intersection Syndrome, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion zero articles in PubMed, zero in Scopus, 1 in CINAHL, 1 in Cochrane Library, zero in Google Scholar and zero in other sources. Zero articles met the inclusion criteria.

**IONTOPHORESIS**

**Recommendation: Iontophoresis for Acute and Subacute Extensor Compartment Tenosynovitis**

Iontophoresis treatments using glucocorticosteroids and sometimes NSAIDs are recommended for extensor compartment tenosynovitis.

**Indications** – Patients with wrist compartment tendinoses. Generally those who either fail to respond adequately to NSAIDs, splints, and activity modifications or decline injection.

**Dose** – Glucocorticosteroid is generally used; however, quality studies have documented successful treatment of lateral epicondylalgia with NSAIDs administered via iontophoresis (see Elbow Disorders Guideline), thus they appear reasonable for this indication as well.

**Frequency/Duration** – Generally 2 or 3 appointments to ascertain efficacy; an additional 4 to 6 appointments may be scheduled if efficacious. If improvements continue at 6 appointments, additional 4 to 6 appointments are reasonable.

**Indications for Discontinuation** – Failure to respond, development of adverse effects, resolution.

**Strength of Evidence** – Recommended, Insufficient Evidence (I)

**Level of Confidence** – Low

**Rationale for Recommendation**

There are no quality studies evaluating iontophoresis for extensor compartment tenosynovitis. Iontophoresis is not invasive, has low adverse effects, but is moderate to high cost depending on the number of treatments. Iontophoresis with either a glucocorticoid or NSAID is recommended for select patients who fail to respond to other treatments or who decline injection.

**Evidence for the Use of Iontophoresis**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Iontophoresis, Extensor Compartment Tenosynovitis, De Quervain Disease, De Quervain’s Stenosing Tenosynovitis, Intersection Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective,
and prospective studies. We found, reviewed and considered for inclusion Zero articles in PubMed, Zero in Scopus, Zero in CINAHL, Zero in Cochrane Library, 25 in Google scholar and zero in other sources. Zero articles met the inclusion criteria.

**OTHER NON-OPERATIVE INTERVENTIONS**

**Recommendation: Other Non-operative Interventions Including Manipulation and Mobilization, Massage, Deep Friction Massage, or Acupuncture for Acute, Subacute, or Chronic Extensor Compartment Tenosynovitis**

There is no recommendation for or against the use of other non-operative interventions (i.e., manipulation and mobilization, massage, deep friction massage, or acupuncture) for the treatment of acute, subacute, or chronic extensor compartment tenosynovitis as other interventions have proven efficacy and are preferentially indicated for initial and subsequent treatment options.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)

Level of Confidence – Low

**Rationale for Recommendation**

There are no quality studies evaluating other non-operative interventions for extensor compartment tenosynovitis. Manual therapy has been attempted;(1096) however, there are no quality studies available to assess its efficacy. Deep friction massage has been used and does not appear successful.(1097)

Evidence for the Use of Acupuncture

There is 1 moderate-quality RCT on acupuncture.(1127) (Hadianfard 14) There are no quality studies incorporated into this analysis for manipulation and mobilization or massage.

**Manipulation & Mobilization:**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Manipulation and Mobilization, Extensor Compartment Tenosynovitis, De Quervain Disease, De Quervain's Stenosing Tenosynovitis, Intersection Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion Zero articles in PubMed, Zero in Scopus, Zero in CINAHL, Zero in Cochrane Library, 169 in Google Scholar, and zero other sources. Zero articles met the inclusion criteria.

**Massage:**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Massage, Massage Therapy, Extensor Compartment Tenosynovitis, De Quervain Disease, De Quervain Stenosing Tenosynovitis, Intersection Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion Zero articles in
Glucocorticosteroid injections are frequently used for the wrist compartment tendinoses.(43, 45, 280, 349, 1091, 1093, 1098-1104) Techniques vary slightly(1101, 1105) and have included attempted selective injection of the extensor pollicis brevis tendon,(1106) although there are no quality studies to compare techniques. Estimates of efficacy in case series and active treatment arms of trials range from 54-100%.(1100, 1101, 1105-1110)

**Recommendation:** **Glucocorticosteroid Injections for Acute, Subacute, or Chronic de Quervain’s or Other Wrist Compartment Tendinosis**

*Glucocorticosteroid injections are recommended for treatment of acute, subacute, or chronic de Quervain’s or other wrist compartment tendinosis.*

**Indications** – Wrist compartment symptoms of pain over a compartment. Generally at least 1 week of non-invasive treatment to determine if condition will resolve without invasive treatment. It is reasonable to treat cases with an initial injection although there is no quality evidence to support that approach. Failure or suboptimal results with an initial injection result in a need for additional injection(s) in a minority of patients that is (are) usually successful.(1103, 1105, 1106)

**Dose** – Optimal dose is unknown. Studies have utilized methylprednisolone acetate 40mg,(1064, 1105, 1109) and triamcinolone acetonide 10mg.(1103, 1106) An adjuvant injectable anesthetic is typically used.(1093, 1105, 1106) Some providers splint the wrist afterwards, however, there is no quality evidence this improves efficacy or duration of benefits. Two low-quality studies suggest no greater efficacy with splinting; however, greater costs and lost time were incurred.(1099, 1111)

**Frequency/Duration** – It is recommended that a single injection be scheduled and the results evaluated to document improvement.(1103) Failure of a response within 1 or 2 weeks should result in reanalysis of the diagnosis and consideration of repeat injection.(1103) Recurrence of symptoms months later should result in consideration of re-injection.(1101, 1105) There is no maximum number of injections to treat an episode or over a lifetime demonstrated in quality studies. Therapeutic injection failures are reportedly strongly associated with the presence of a separate compartment for the extensor pollicis brevis tendon in 73% of cases.(1109)

**Indications for Discontinuation** – If a partial response, consideration should be given to repeating the injection, typically at a modestly higher dose.

**Strength of Evidence** – **Recommended, Evidence (C)**
Level of Confidence – High

Rationale for Recommendation
There is 1 moderate-quality study comparing glucocorticosteroid injections with placebo for treatment of de Quervain’s stenosing tenosynovitis. (1103) The trial showed considerable benefits from active treatment that persisted for 12 months and allows for an evidence-based recommendation. One trial found steroid injection superior to acupuncture. (1112) Ultrasound-guidance has been suggested to be moderately superior. (1113) Two trials have found inconclusive evidence regarding whether splint use is required in addition to steroid injection. (1114, 1115) A high-quality trial found the steroid flare was unrelated to pH; (1064) however, there was no placebo control group. Another high-quality trial found no additive benefit of NSAID in addition to injection to prevent recurrence but did not assess reductions in pain immediately after injection thus appears to have no bearing on use of NSAIDs for those purposes. (1093) A low-quality trial found glucocorticosteroid injection superior to splinting in pregnant and lactating females. (1098) These injections are minimally invasive, have low adverse effects, and are moderately costly; thus, they are recommended to treat de Quervain’s or other wrist compartment tendinosis.

Evidence for the Use of Glucocorticosteroid Injections for Wrist Compartment Tendinoses
There are 2 high- (1064, 1093) and 5 moderate-quality (1103, 1112-1115) RCTs incorporated in this analysis. There are 3 low-quality RCTs and 1 longitudinal study (1098, 1099, 1109, 1116) in Appendix 2.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Glucocorticosteroid injection, corticosteroid injection, glucocorticoids, extensor compartment tenosynovitis, de Quervain’s stenosing tenosynovitis, and intersection syndrome, de Quervain disease; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 10 articles in PubMed, 43 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 19 in Google Scholar, and 2 from other sources. We considered for inclusion 7 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 2 from other sources. Of the 7 articles considered for inclusion, 7 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jirarattanaphochai 2004 RCT</td>
<td>10.5</td>
<td>N = 160 (144 female/16 male) with de Quervain disease, positive Finkelstein test, radial styloid tenderness, pain on first extensor compartment with thumb abduction or extension. Mean (±SD) age 48.98 (±9.10) nimesulide group, 46.87 (±12.79) placebo.</td>
<td>Injection of 10mg of triamcinolone acetonide and 0.5mL 1% lidocaine and either 200mg daily oral nimesulide group (n = 80) vs. placebo control group (n = 80). Both groups received allocated treatment for 7 days. Follow-up at 1 week, 6, 12, 18 and 24 months.</td>
<td>No significant differences reported between the nimesulide and placebo groups for VAS pain scores, success rates, adverse reactions and probability of recurrence.</td>
<td>No significant differences reported between the nimesulide and placebo groups for VAS pain scores, success rates, adverse reactions and probability of recurrence.</td>
<td>Data suggest nimesulide does not enhance effectiveness of a single triamcinolone injection in de Quervain's disease treatment. Also, disease recurrence was correlated to the presence of crepitation in the first dorsal compartment at thumb extensor abduction.</td>
</tr>
<tr>
<td>Peters-Veluthamamangal BMC Musculo-skelet Disord 2009 RCT</td>
<td>7.5</td>
<td>N = 21 (13 female/21 male) clinical diagnosis of de Quervain's with Finkelstein's or crepitations on exam. Mean age 52.3 (12.6) for NaCl; 51.2 (20.2) for TCA group.</td>
<td>NaCl, 1-2 injection 1ml triamcinolonacetonide (n = 12) vs. placebo or TCA. 1mL NaCl at site of maximal tenderness. Second injection by different MD at 2 weeks if not satisfied with results; 12 month follow-up (n = 9).</td>
<td>Short-term results of mean pain severity in the past week of saline 4.3 vs. corticoid 1.3. Patients much better or better: 2/12 (33%) saline vs. 7/9 (77.8%), p = 0.047. Maintained improvement over 12 months.</td>
<td>Short-term results of mean pain severity in the past week of saline 4.3 vs. corticoid 1.3. Patients much better or better: 2/12 (33%) saline vs. 7/9 (77.8%), p = 0.047. Maintained improvement over 12 months.</td>
<td>Under enrollment. Small sample size. Considerable differences nevertheless suggest efficacy.</td>
</tr>
<tr>
<td>Goldfarb 2007 RCT</td>
<td>8.0</td>
<td>N = 125 (93 female/32 male) with trigger finger or de Quervain's. Average age was 59 years.</td>
<td>Balanced group, methylprednisolone acetate 40mg, lidocaine, bupivacaine alone (n = 68) vs. Standard group or injection except balanced solution and neutral pH (n = 57); 6 week follow-up.</td>
<td>All immediately responded to injection. Pain rebounded at one day, and then gradually decreased. 23/68 (33.8%) in balanced group vs. 18/57 (31.6%) in acidic pH group had flare reactions (NS).</td>
<td>All immediately responded to injection. Pain rebounded at one day, and then gradually decreased. 23/68 (33.8%) in balanced group vs. 18/57 (31.6%) in acidic pH group had flare reactions (NS).</td>
<td>No placebo group. Some trends in baseline differences of unclear significance. Purpose to assess steroid flare and whether normal pH could reduce this adverse effect. Study suggests steroid flare unrelated to pH.</td>
</tr>
</tbody>
</table>

**Glucocorticosteroid vs. Saline Injections**
### Glucocorticosteroid with vs. without NSAID

**Jirarattanaphochai 2004**

- **RCT**
- No sponsorship. One or more authors received grants or outside funding from the Faculty of Medicine, Khon Kaen University.

| N = 160 (144 female/16 male) with de Quervain disease, positive Finkelstein test, radial styloid tenderness, pain on first extensor compartment with thumb abduction or extension. Mean (±SD) age 48.98 (±9.10) nimesulide group and 46.87 (±12.79) placebo group. | Injection of 10mg of triamcinolone acetonide and 0.5mL of 1% lidocaine and either 200mg daily oral nimesulide group (n = 80) vs. Placebo control group (n = 80). Both groups received allocated treatment for 7 days. Follow-up at 1 week, 6, 12, 18, and 24 months. | Success rates after 1 injection: 67% nimesulide vs. 68% placebo (NS). Overall success 95% both groups. Risk for recurrence doubles with crepitation (RR = 2.13, 95% CI 1.19-3.8). | “Supplemental oral administration of the nonsteroidal anti-inflammatory drug nimesulide does not improve the effectiveness of a single injection of triamcinolone acetonide in the treatment of de Quervain disease.” | No placebo; no recording of pain scores for purposes of evaluating reduced pain after injection. Variable follow-up. Data suggest NSAID provides no incremental benefit to prevent recurrence in addition to steroid injection. |

### Injection vs. Other Treatments

**Hadianfard 2014**

- **RCT**
- Supported by Vice-Chancellery of Research and Technology of Shiraz University of Medical Sciences, Shiraz, Iran. No mention of COI.

| N= 35 patients (6 males, 24 females) with clinical diagnosis of De Quervain’s tenosynovitis. Mean age was 40.7 years. | Acupuncture group: Received 5 acupuncture sessions of 30 minutes duration (n = 18) vs. Injection Group: 1 methylprednisolone acetate injection in first dorsal compartment of wrist (n = 17). Follow-up for 6 weeks. | At last follow-up Q-DASH score decreased by 55.1 in injection group vs. 54.6 in acupuncture group. No significant differences between groups. Difference between baseline and final VAS score decreased significantly between groups, but not significant between groups (p> 0.05). | “We demonstrated short-term improvement of pain and function in both groups. Although the success rate was somewhat higher with corticosteroid injection, acupuncture can be considered as an alternative option for treatment of De Quervain’s Tenosynovitis.” | Data suggests methylprednisolone injections somewhat better than acupuncture for improved pain and function in deQuervain’s tenosynovitis although both groups improved from baseline at 2 and 6 weeks. |

**Kume 2012**

- **Randomized prospective trial**
- No sponsorship or COI.

<p>| N = 44 wrists (5 males, 39 females) with diagnosed de Quervain’s disease | Ultrasound guided injection group (n = 22) vs. Manual injection group (n = 22). Both groups received 20 mg of triamcinolone and 1 ml of 1% lidocaine. Follow-up at 4 weeks. | Reduction in mean VAS pain from baseline to 4 weeks significantly higher in ultrasound guided group vs. manual injection group: 80.3 to 25.6 vs. 78.0 to 58.2. (p = 0.0007). No adverse reactions related to treatment for either group. | “[U]S-guided injection targeting the EPB of dQD with septation was found to be more effective than clinically guided manual injection.” | Data suggest US guided injection targeting EPB in deQuervain’s patients with septation is better than manual injection although both groups showed improvement in pain on VAS. |</p>
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Participants</th>
<th>Intervention</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mardani-Kivi 2014</td>
<td>Randomized prospective trial</td>
<td>No sponsorship or COI</td>
<td>N = 67 patients (12 males, 47 females) with extensor compartment tenosynovitis, or de Quervain tendinopathy, radial pain of the wrist, a positive Finkelstein test, tenderness of the first dorsal compartment and a pain score &gt;6. Mean (±SD) age 42 (±13) for CSI+TSC group and 45 (±12) for CSI only group.</td>
<td>Corticosteroid injection (CSI) and thumb spica cast (TSC) (3 weeks casted) group (n = 33) vs. Corticosteroid injection only group (n = 34). Both groups 40mg of methylprednisolone acetate with 1cc of lidocaine 2%. Follow-up at 3 weeks and 6 months.</td>
</tr>
<tr>
<td>Mehdinasab 2010</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>N= 73 patients (9 males, 64 females) with de Quervain's tenosynovitis. Mean age was 32.6 years.</td>
<td>Injection Group- Injection of methylprednisolone acetate in first dorsal compartment of wrist followed by wrist thumb spica cast (n = 37) vs. Casting Group- Casting only (n = 36). Follow-up for 6 months,</td>
</tr>
</tbody>
</table>
**Surgery**

**Recommendation:** Surgical Release for Subacute or Chronic Extensor Compartment Tenosynovitis

Surgical release is recommended for patients with subacute or chronic extensor compartment tenosynovitis who fail to respond to injection. (1101)

**Indications** – Wrist compartment tenosynovitis that fails to respond to non-operative interventions generally including at least 2 glucocorticosteroid injections.

**Strength of Evidence** – Recommended, Insufficient Evidence (I)

**Level of Confidence** – Moderate

**Rationale for Recommendation**

There are no quality studies evaluating the use of surgical release for extensor compartment tenosynovitis. (346, 1117) While surgery release is invasive, has moderate adverse effects, and is costly, it is usually clinically effective and recommended for patients who have failed glucocorticosteroid injection(s) and other non-invasive treatments.

**Evidence for the Use of Surgery**

There is 1 moderate-quality RCT incorporated into this analysis. (1145) (Abrisham 11)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: extensor compartment tenosynovitis, de Quervain’s stenosing tenosynovitis, and intersection syndrome, de Quervain disease; Surgical release; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 1 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 30 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 Google Scholar, and 0 from other sources. Of the 31 articles considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abrisham 2011 RCT</td>
<td>5.5</td>
<td>N = 120 (24 males, 96 females). Mean age is 45.3 years.</td>
<td>Transverse Incision (N = 60) vs Longitudinal Incision (N =60). Followed for three months. An additional follow up of 2 weeks to remove sutures and finally three months for final assessment.</td>
<td>Complications of surgical treatment with longitudinal incision were lower than the transverse incision. Longitudinal incision had five hypertrophic scars and no injury to nerve or vein reported. Transverse incision had 3 lesions to superficial branch of radial nerve, five injuries to vein in snuffbox area, and five hypertrophic scars.</td>
<td>Longitudinal incision can be recommended for surgical treatment of De Quervain disease.</td>
<td>Data suggest longitudinal incision is superior to transverse incision for treatments of De Quervain tenosynovitis in terms of post-op complication. After a period of 3 months, 14 patients (8 transverse and 6 longitudinal) did not cooperate in follow up from the first time.</td>
</tr>
</tbody>
</table>
Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome)

Diagnostic Criteria
There is no quality evidence comparing diagnostic testing for this disorder. Most case series report electrodiagnostic testing assisted in making a diagnosis. The characteristic finding is a prolonged distal motor latency. One report opined that idiopathic or “cumulative stress” cases have no characteristic pattern.(50) Electrodiagnostic calculations are complicated by the curvilinear course of the deep motor nerve. Witmer described a technique reducing the complexity that may be useful to the electromyographer.(1118)

Special Studies and Diagnostic and Treatment Considerations
ELECTRODIAGNOSTIC STUDIES
Recommendation: Electrodiagnostic Testing for Ulnar Nerve Entrapment at the Wrist
Electrodiagnostic testing is recommended to confirm clinical suspicion of ulnar nerve entrapment at the wrist.

   Strength of Evidence – Recommended, Evidence (C)
   Level of Confidence – Moderate

Rationale for Recommendation
There are 3 moderate studies supporting the use of electrodiagnostic testing.(1119-1121) However, studies need to be performed by well-trained electrodiagnosticians, preferably certified by the American Board of Electrodiagnostic Medicine.

Evidence for the Use of Electrodiagnostic Studies
There are 4 moderate-quality studies incorporated into this analysis.(1119-1122)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Electrodiagnostics nerve conduction study, electromyography, Ulnar Nerve Entrapment at the Wrist (Including Guyon's Canal Syndrome and Hypothenar Hammer Syndrome) diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 0 articles in PubMed, 48 in Scopus, 2 in CINAHL, 3 Cochrane Library, and 350 from Google Scholar. We considered for inclusion 0 from PubMed, 2 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 2 from other sources. Of the 4 articles considered for inclusion 4 diagnostic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI) Score</th>
<th>Population/Case Definition</th>
<th>Investigative Test</th>
<th>Gold Standard/Comparative Test</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lander 2007</td>
<td>Cross-sectional study</td>
<td>6.0</td>
<td>N = 162 referred for hand-arm vibration syndrome or HAVS assessment at specialist occupational health clinic, plus history of exposure to vibrating tools. Mean age onset of symptoms 38.4 (9.0).</td>
<td>Nerve conduction studies (NCS) and current perception threshold (CPT). Using Stockholm sensorineural or SSN scale and quantitative sensory tests (QSTs) measuring vibration and temperature perception.</td>
<td>NCS vs. CPT tests for both upper extremities. Perception measured at 5 Hz, 250 Hz and 2 kHz at index finger for median nerve and at little finger for ulnar nerve.</td>
<td>160 (99%) complained of numbness and/or tingling. CPT in left hand abnormal in 99 subjects, In left hand, overall CPT results ($\chi^2 = 9.87, p = 0.007$) and results from ulnar nerve ($\chi^2 = 11.27, p = 0.004$); significantly associated with SSN staging. CPT and NCS results significantly associated for each of ulnar, median and overall nerve results in right hand and left hand, (p = 0.0001).</td>
<td>Workers being assessed for HAVS should have nerve conduction testing to detect neuropathies proximal to the hand.</td>
<td>Data suggests NCS and CPT significantly associated for the overall results and for ulnar and median results in each hand.</td>
</tr>
<tr>
<td>Hirata 2007</td>
<td>Age-matched</td>
<td>5.0</td>
<td>N = 75 males and controls with hand-arm vibration syndrome (VS). Mean age 58.7 years.</td>
<td>Sensory nerve conduction velocities (SCVs); 0.1-ms rectangular electric pulses at 1 Hz</td>
<td>Associations between frequency of slowed SCV and reduced AMP and frequency of neuropathy types</td>
<td>In median nerve, SCVfp-fd, SCVw-e, AMPw-fp and AMPw-fd significantly reduced vs. controls, (p = 0.005, 0.011, 0.024, 0.013). In ulnar nerve, SCVfp-′fd, SCVw-′fp, AMPw-′fp, AMPw-′fd, AMPfo-′fp and AMPup-′fp significantly reduced in VS patients vs controls (p = 0.000, 0.015, 0.007, 0.000, 0.027 and 0.008). In radial nerve, AMPfo-′th significantly reduced in VS patients vs controls, (p = 0.003).</td>
<td>These findings suggest that VS affects all three nerves in the hand. According to classification results, the main disorders of peripheral nerves comprise digital neuropathy.</td>
<td>Small sample size. Data suggests that vibration syndrome affects all three hand nerves and neuropathy due to VS may in fact represent a multi-focal neuropathy.</td>
</tr>
<tr>
<td>Alaranta 1977</td>
<td>4. 5</td>
<td>An automatic analysis of the electromyographic activity.</td>
<td>N = 38 forest workers and pneumatic-tool operators. Male workers aged 26 to 61 years.</td>
<td>Velocity of lower motor fibers (CVSF) of ulnar nerve and motor distal latency (DL) of median nerve</td>
<td>Subgroup 0 = normal conduction velocity of CVSF and distal latency DL Subgroup 1 = Only one CVSF of ulnar nerve or DL Subgroup 2 = polyneuropathic findings vs vibration syndromes, at least 2 abnormal CVSF or DL findings.</td>
<td>Exposed workers had statistically lower CVSFs of ulnar nerve (p &lt;0.001) and dSCVs of median nerve (p &lt;0.001), longer DLs of median nerve (p &lt; 0.01), and slightly slower dSCVs of ulnar nerve (p &lt;0.05) and SCVs of median nerve (p &lt;0.05) vs. none exposed, as a group.</td>
<td>“In accordance with previous reports the CVSF of the ulnar nerve was a potent factor in differentiating the vibration exposed workers from those nonexposed.”</td>
<td>Data suggests conduction velocity of slower motor fibers of ulnar nerve, distal sensory conduction velocity and motor distal latency of median nerve most sensitive measurement for separation of those with traumatic vasospastic disease from those not exposed.</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Chatterjee 1982</td>
<td>4. 0</td>
<td>Exploratory/observational</td>
<td>N = 31 rock-drillers and controls. Age range 24-57, mean age 37.9 (9.6).</td>
<td>Exploratory electromyography; Disa-type 14 A 30 3-channel electromyograph y with a 14 G 01 digital average capable of averaging up to 1024 successive stimuli.</td>
<td>Motor and sensory conduction velocities in median and ulnar Nerves; and latency, duration, and the amplitude of evoked action potentials measured.</td>
<td>Significant difference between groups, in 1st (p &lt;0.05), 2nd (p &lt;0.01), fourth (p &lt;0.05) digits supplied by median nerve in right hand and first and fourth digits, (p &lt; 0.05). Sensory duration varied from 2.1msec to 2.6 msec in right hand and 1.9 msec to 2.2 msec in left hand vs. controls 1.8 msec and 2.1 msec. Vibration groups significant in first (p &lt; 0.05), second (p &lt; 0.05), third (p &lt; 0.01), fourth (p &lt; 0.01) digits supplied by median nerve and other half of fourth digit (p &lt; 0.05) supplied by ulnar nerve in right hand vs. controls.</td>
<td>The results showed that apart from sensory duration the control group had values that were closest to the students while the vibration group had values furthest away.”</td>
<td>Small sample. Data suggests neurophysiological changes are frequent in those who regularly use vibrating tools and with the exception sensory duration, the median nerve is affected more than the ulnar nerve.</td>
</tr>
</tbody>
</table>
MRI/ULTRASOUND

**Recommendation: MRI or Ultrasound to Diagnose Ulnar Nerve Entrapment at the Wrist**

There is no recommendation for or against the use of MRI or ultrasound to diagnose ulnar nerve entrapment at the wrist. Use of MRI for a suspected soft-tissue mass may be reasonable.

*Strength of Evidence – No Recommendation, Insufficient Evidence (I)*

*Level of Confidence – Low*

**Evidence for the Use of MRI and Ultrasound**

There are no quality studies incorporated into this analysis.

MRI:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Magnetic resonance imaging, MRI, Ulnar Nerve Entrapment, Guyon’s Canal Syndrome, Hypothenar Hammer Syndrome, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 88 articles in PubMed, 0 in Scopus, 0 in CINAHL, 3 in Cochrane Library, 85 from Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Ultrasound:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ultrasound, Ultrasonography, Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome), diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 69 articles in PubMed, 2 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 95 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**CT**

**Recommendation: CT to Diagnose Ulnar Nerve Entrapment at the Wrist**

CT is recommended to diagnose ulnar nerve entrapment at the wrist if a hook of the hamate fracture is suspected based upon the history, a mechanism of potential fracture, focal pain at the hamate and where there are ulnar nerve symptoms.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*

*Level of Confidence – Low*

**Rationale for Recommendations**

There are no quality studies evaluating the use of ultrasound or MRI for ulnar nerve entrapment at the wrist and therefore, there is no recommendation for or against the use of these tests. The use of ultrasound, MRI, or CT has also been reported useful in identifying suspected hamate fractures or mass lesions such as a ganglion cyst.(50, 57, 1118) MRI is generally preferable for soft tissue masses and CT is preferable for boney masses. These tests are moderate to high cost, but are recommended for evaluation of select patients suspected of having occult fractures of the hamate or mass lesions.

**Evidence for the Use of CT**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: CT, CAT, X-Ray CT, Ulnar Nerve Entrapment, Guyon’s Canal Syndrome, Hypothenar Hammer Syndrome, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0
Cochrane Library, and 300 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Initial Care
Ulnar neuropathy at the wrist that is not related to trauma, such as from the use of wheelchair, crutches, or other equipment may benefit initially from non-invasive therapies and activity adjustments including elimination or mitigation of significant pressure points (e.g., using padding, etc.) and splinting.

ACTIVITY MODIFICATION
Recommendation: Activity Modification for Ulnar Nerve Compression at the Wrist
Activity modification, with particular avoidance of significant localized mechanical compression of the nerve or use of the hand as a hammer is recommended for treatment of ulnar nerve compression at the wrist.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Evidence for the Use of Activity Modification
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Rest, resting, Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 1 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 0 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 1 articles considered for inclusion, 0 randomized trials and 1 systematic study met the inclusion criteria.

SPLINTS
Recommendation: Neutral Wrist Splinting for Acute, Subacute, or Chronic Ulnar Nerve Compression at the Wrist
Neutral wrist splinting is recommended as a first-line treatment for acute, subacute, or chronic ulnar nerve compression at the wrist.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Evidence for the Use of Splints
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: splint, splints, splinting; ulnar nerve compression syndromes, ulnar nerve entrapment, wrist, guyon’s canal syndrome, guyon syndrome, ulnar tunnel syndrome, hypothenar hammer syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 68 articles in PubMed, 6 in Scopus, 0 in CINAHL, 9 in Cochrane Library, 283 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.
Follow-up Visits
Follow-up visits are generally required every 2 to 4 weeks to evaluate efficacy of interventions until resolution of the condition.

Medications

NSAIDS
Recommendation: NSAIDS for Acute, Subacute, or Chronic Ulnar Nerve Compression at the Wrist
There is no recommendation for or against the use of NSAIDs to control pain associated with acute, subacute, or chronic ulnar nerve compression at the wrist.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There is no recommendation for or against the use of NSAIDs for acute or subacute ulnar nerve compression at the wrist as evidence of efficacy is lacking. NSAIDs do not work particularly well for other neuropathies (see Chronic Pain Guideline and CTS section), thus other options are generally preferable.

Evidence for the Use of NSAIDs
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDs, acetaminophen Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 150 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

GLUCOCORTICOSTEROIDS
Recommendation: Glucocorticosteroids for Acute, Subacute, or Chronic Ulnar Nerve Compression at the Wrist
There is no recommendation for or against the use of oral and injected glucocorticosteroids for treatment of acute, subacute, or chronic ulnar nerve compression at the wrist.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendations
There are no quality studies evaluating these treatments for ulnar nerve compression at the wrist. Activity modification to avoid focal mechanical compression and the use of the hypothenar area as a hammer are thought to be important and are recommended. NSAIDs have been utilized. However, evidence of efficacy for treatment of CTS and other neuropathic pain (see Chronic Pain Guideline) is lacking, thus other options are generally preferable. The mechanism(s) of efficacy of glucocorticosteroids for treatment of CTS and other disorders is unclear. If the mechanism involves tendon sheaths and related structures, then these medications would be predicted to be ineffective for ulnar neuropathy at the wrist. However, if through another mechanism of action directly involving the nerve sheath, then these injections could be effective. These treatments are not invasive to low invasive, have few adverse effects and are low to moderate cost. They are recommended with the exceptions of NSAIDs and injections for which there is no evidence of efficacy and concerns that the available literature does not support those treatments as efficacious.

Evidence for the Use of Glucocorticosteroids
There are no quality studies incorporated into this analysis.
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Glucocorticosteroids, glucocorticoids, Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 3784 in Scopus, 0 in CINAHL, 1 in Cochrane Library, 150 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 1 articles considered for inclusion, 0 randomized trials and 1 systematic studies met the inclusion criteria.

**Physical Methods/Rehabilitation**

*Recommendation: Physical Methods/Rehabilitation for Acute, Subacute, or Chronic Ulnar Neuropathy at the Wrist*

There is no recommendation for or against the use of physical methods/rehabilitation (i.e., iontophoresis, self-application of ice or heat, manipulation, mobilization, massage, friction massage, or acupuncture) for treatment of acute, subacute, or chronic ulnar neuropathy at the wrist.

*Strength of Evidence – No Recommendation, Insufficient Evidence (I)*

*Level of Confidence – Low*

**Rationale for Recommendation**

There are no quality studies evaluating the efficacy of physical methods/rehabilitation (i.e., iontophoresis, ice, heat, manipulation, mobilization, massage, friction massage, and acupuncture) for ulnar neuropathy at the wrist and therefore, there is no recommendation for or against the use of these treatments.

**Evidence for the Use of Physical Methods/Rehabilitation**

There are no quality studies incorporated into this analysis.

**Iontophoresis**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: iontophoresis; ulnar nerve compression syndromes, ulnar nerve entrapment, wrist, guyon’s canal syndrome, guyon syndrome, ulnar tunnel syndrome, hypothenar hammer syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 0 in Scopus, 0 in CINAHL, 2 in Cochrane Library, 41 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Ice**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ice; Self Application, Ulnar Nerve Compression Syndromes, Ulnar Nerve Entrapment, Wrist, Guyon’s Canal Syndrome, Guyon Syndrome, ulnar tunnel syndrome, Hypothenar Hammer Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 1 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 350 in Google Scholar and 0 in other sources. Zero articles met the inclusion criteria.

**Heat**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Heat; Self Application, Ulnar Nerve Compression Syndromes, Ulnar Nerve Entrapment, Wrist, Guyon’s Canal Syndrome, Guyon Syndrome, ulnar tunnel syndrome, Hypothenar Hammer Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 1 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 350 in Google Scholar and 0 in other sources. Zero articles met the inclusion criteria.
Compression Syndromes, Ulnar Nerve Entrapment, Wrist, Guyon’s Canal Syndrome, Guyon Syndrome, ulnar tunnel syndrome, Hypothenar Hammer Syndrome, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 1 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 730 in Google Scholar, and 0 in other sources. Zero articles met the inclusion criteria.

Manipulation/Mobilization

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: manipulation, mobilization, Ulnar Nerve Entrapment at the Wrist including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, and 0 in Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 0 articles considered for inclusion, 0 randomized trials and 0 systematic studies met the inclusion criteria.

Massage

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Massage, Ulnar Nerve Compression Syndromes OR Ulnar Nerve Entrapment, Wrist, Or Guyon Syndrome or Guyon’s Canal Syndrome or ulnar tunnel syndrome or Hypothenar Hammer Syndrome ;controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, and 0 in Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 0 articles considered for inclusion, 0 randomized trials and 0 systematic studies met the inclusion criteria.

Acupuncture

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: acupuncture, Ulnar Nerve Entrapment at the Wrist (Including Guyon’s Canal Syndrome and Hypothenar Hammer Syndrome);controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, and 0 in Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar and 0 from other sources. Of the 0 articles considered for inclusion, 0 randomized trials and 0 systematic studies met the inclusion criteria.

EXERCISE

Exercise is not generally indicated acutely. Many patients with chronic findings and functional deficits and post-operative patients require some appointments to at minimum help institute a home exercise program. For patients with residual deficits, particularly post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

Evidence for the Use of Exercise

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: exercise, exercising, physical activity; ulnar nerve compression syndromes, ulnar nerve entrapment, wrist, guyon’s canal syndrome, guyon syndrome, ulnar tunnel syndrome, hypothenar hammer syndrome; controlled clinical trial, controlled
trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 9 articles in PubMed, 3 in Scopus, 0 in CINAHL, 16 in Cochrane Library, 468 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Surgery**

Space occupying lesions with significant motor or sensory deficits generally have been reported in the literature as requiring surgical decompression (or needle aspiration of ganglia) with excellent results and rapid recovery of deficits. In addition to lesion type, consideration may be influenced by the presence of diabetes mellitus. Although there are not quality studies, there may be a stronger indication for decompression of peripheral nerve entrapment syndromes in diabetic patients. In a case series of diabetics with peripheral neuropathy, decompression surgery improved sensory function in 88% of upper extremities and 69% of lower extremities compared with 32% of patients that were treated non-operatively. (1123) Another case series demonstrated similar results, also favoring recovered function in the upper extremities. (1124)

**Recommendation: Surgical Decompression for Subacute or Chronic Ulnar Nerve Compression at the Wrist**

Surgical decompression is recommended for subacute or chronic ulnar nerve compression at the wrist after failure of non-operative treatment or if space-occupying lesions are present.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*

*Level of Confidence – Moderate*

**Rationale for Recommendation**

There are no quality studies evaluating the efficacy of surgical intervention for ulnar nerve compression at the wrist. However, clinically many patients respond well to surgery. Surgery is invasive, has adverse effects and is costly. It is recommended for select patients who failed trials of other non-operative treatments or if space occupying lesions are present. It may also be preferential in those with diabetes mellitus.

**Evidence for the Use of Surgery**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: surgery, surgeries, surgical decompression; Ulnar Nerve Compression Syndromes, Ulnar Nerve Entrapment, Wrist, Guyon’s Canal Syndrome, Guyon Syndrome, ulnar tunnel syndrome, Hypothenar Hammer Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 224 articles in PubMed, 12 in Scopus, 3 in CINAHL, 12 in Cochrane Library, 628 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Radial Nerve Entrapment**

**Special Studies and Diagnostic and Treatment Considerations**

**ELECTRODIAGNOSTIC STUDIES**

Electrodiagnostic studies can confirm the diagnosis of a radial nerve motor neuropathy. (58)

**Recommendation: Electrodiagnostic Testing for Radial Nerve Motor Neuropathy**

Electrodiagnostic testing is recommended to confirm clinical suspicion of a radial nerve motor neuropathy.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*
Level of Confidence – Moderate

**Rationale for Recommendation**

There is no quality evidence available for the use of electrodiagnostic testing; however, it is recommended as an objective test to evaluate radial nerve motor neuropathy. (58, 59, 1125) However, studies need to be performed by well-trained electrodiagnosticians, preferably certified by the American Board of Electrodiagnostic Medicine.

**Evidence for the Use of Electrodiagnostic Studies**

There are no quality studies incorporated into this analysis. There are 2 low-quality studies in Appendix 2. (1154, 1155) (Spindler 90; Verhaar 91)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: electrodiagnostic study, nerve conduction study, electromyography, radial nerve entrapment, radial tunnel syndrome, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 6 articles in PubMed, 86 in Scopus, 0 in CINAHL, 1 in Cochrane Library, and 160 from Google Scholar. We considered for inclusion 2 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 3 articles considered for inclusion 2 diagnostic studies met the inclusion criteria.

**ULTRASOUND** (DIAGNOSTIC)

Ultrasound has been used as an adjunct to electrophysiological studies for evaluation of radial nerve neuropathy. (446)

**Recommendation: Diagnostic Ultrasound for Radial Nerve Neuropathy**

There is no recommendation for or against ultrasound to confirm clinical suspicion of a radial nerve neuropathy.

**Strength of Evidence – No Recommendation, Insufficient Evidence (I)**

**Level of Confidence – Moderate**

**Rationale for Recommendation**

There is no quality evidence available that diagnostic ultrasound materially alters the ability to diagnose radial nerve entrapments and thus there is no recommendation for or against diagnostic ultrasound.

**Evidence for the Use of Ultrasound**

There is 1 moderate-quality study incorporated into this analysis. (446)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ultrasound, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 7 articles in PubMed, 93 in Scopus, 0 in CINAHL, 0 in Cochrane Library, and 8540 from Google Scholar, and 0 from other sources. We considered for inclusion 2 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. One article met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score</th>
<th>Area</th>
<th>Diagnoses</th>
<th>Type of Ultrasound</th>
<th>CT used</th>
<th>MRI used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Neurosurgery Performed</th>
<th>Clinical outcomes assessed</th>
<th>Long-term follow-up</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lo 2008</td>
<td>Diagnostic</td>
<td>No mention of sponsorship or COI</td>
<td>7.0</td>
<td>10 (3 female/7 male) with suspected radial neuropathy</td>
<td>HWF</td>
<td>Radial nerve entrapment</td>
<td>Medtronic Keypoint EMG Machine</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
<td>-</td>
<td>Ultrasound correctly identified all 6 with radial neuropathy. Significantly less mean (SD) time for US exam time vs. NCS/EMG: 6.1 (1.1) minutes vs. 30.3 (2.7), p &lt;0.001.</td>
<td>“US is of value as a rapid diagnostic adjunct for the localization of radial nerve entrapment.”</td>
</tr>
</tbody>
</table>

**Initial Care**

Overall, the literature suggests patients most often appear to respond to non-operative treatments including no treatment; avoidance of exposures thought to be contributing (if present); avoidance of wearing a watch, tight jewelry or shirt sleeves on the affected side; corticosteroid injection;(1126) and temporary thumb spica splinting.(66, 1127)

**SPLINTS**

**Recommendation: Wrist Extension or Thumb Spica Splint for Acute, Subacute, or Chronic Radial Nerve Compression Neuropathy**

The use of a wrist extension or thumb spica splint is recommended for treatment of acute, subacute, or chronic radial nerve compression neuropathy.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*

*Level of Confidence – Low*

**Rationale for Recommendation**

Splints appear to be helpful for many cases and thus are recommended, particularly wrist extension splints.

**Evidence for the Use of Splints**

There are no quality studies incorporated into this analysis.
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: splint, splinting, thumb spica, radial nerve entrapment, radial tunnel syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 3 in Scopus, 2 in CINAHL, 7 in Cochrane Library, 180 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Follow-up Visits
Follow-up visits are generally required every 2 to 4 weeks to evaluate efficacy of interventions until resolution of the condition.

Medications
NSAIDs
Recommendation: NSAIDS for Acute, Subacute, or Chronic Radial Nerve Compression Neuropathy
There is no recommendation for or against the use of NSAIDs to control pain associated with acute, subacute, or chronic radial nerve compression neuropathy.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Evidence for the Use of NSAIDs
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDs, acetaminophen, non-steroidal anti-inflammatory, radial nerve entrapment, radial tunnel syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 10 in Scopus, 0 in CINAHL, 2 in Cochrane Library, 170 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

GLUCOCORTICOSTEROIDS
Recommendation: Glucocorticosteroids for Acute, Subacute, or Chronic Radial Nerve Compression at the Wrist
There is no recommendation for or against the use of oral and injected glucocorticosteroids for treatment of acute, subacute, or chronic radial nerve compression at the wrist.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendations
Although there are no quality studies on which to rely for the treatment of distal radial neuropathies, non-invasive options are available and have few adverse effects and are low cost. NSAIDs are not unreasonable and are recommended by some(60) however, evidence of efficacy is lacking, NSAIDs do not work particularly well for other neuropathies (see Chronic Pain guideline); thus, other options are generally preferable. The mechanism(s) of efficacy of glucocorticosteroids is unclear. If the mechanism involves tendon sheaths and related structures, then these medications would be predicted to be ineffective for distal radial neuropathies. However, if through another mechanism of action directly involving the nerve sheath, then these injections could be effective. These treatments are not invasive to low invasive, have few adverse effects, and are low to moderate cost. They are recommended, with the
exception of NSAIDs and injections for which there is no evidence of efficacy and concerns that the available literature does not support those treatments as efficacious.

Evidence for the Use of MRI and Ultrasound
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: oral, injection, intravenous, glucocorticosteroid, corticosteroids, steroid, radial nerve entrapment, radial tunnel syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, random; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 53 in Scopus, 2 in CINAHL, 5 in Cochrane Library, 236 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 2 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 3 articles considered for inclusion, 0 randomized trials and 3 systematic studies met the inclusion criteria.

Physical Methods/Rehabilitation

Recommendation: Physical Methods (Iontophoresis, Self-application of Ice or Heat, Manipulation and Mobilization, Massage, Friction Massage, or Acupuncture) for Acute, Subacute, or Chronic Radial Neuropathy at the Wrist

There is no recommendation for or against the use of physical methods for treatment of acute, subacute, or chronic radial neuropathy at the wrist including iontophoresis, self-application of ice or heat, manipulation and mobilization, massage, friction massage, or acupuncture.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There are no quality studies evaluating the use of iontophoresis, self-application of ice and heat, manipulation and mobilization, friction massage, or acupuncture for radial neuropathy at the wrist and therefore, there is no recommendation for or against these treatments. There are reports of benefits from massage, but no quality studies, thus there is no recommendation for massage.

Evidence for the Use of Physical Methods/Rehabilitation
There are no quality studies incorporated into this analysis.

Ice:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ice; Self Application of Ice, Radial Nerve Entrapment, Radial Tunnel Syndrome, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, random; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 6 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 5670 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Heat:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Heat; Self Application of Heat, Radial Nerve Entrapment, Radial Tunnel Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, random; systematic, systematic review, retrospective, and prospective studies. We found and reviewed
0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 2384 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Manipulation & Mobilization:**
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Manipulation, mobilization, Radial Nerve Entrapment, Radial Tunnel Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, and 0 in Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 0 articles considered for inclusion, 0 randomized trials and 0 systematic studies met the inclusion criteria.

**Massage:**
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Massage, friction massage, Radial Nerve Entrapment, Radial Tunnel Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 0 from Google Scholar and 0 in other sources. Zero articles met the inclusion criteria.

**Acupuncture:**
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Acupuncture, Radial nerve entrapment, Radial tunnel syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 0 from Google Scholar and 0 in other sources. Zero articles met the inclusion criteria.

**Iontophoresis:**
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Iontophoresis, Radial Nerve Entrapment, Radial Tunnel Syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 34 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 60 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**EXERCISE**
Exercise is not generally indicated acutely. Many patients with chronic findings, functional deficits and post-operative patients require some appointments to at minimum help institute a home exercise program. For patients with residual deficits, particularly post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

_Evidence for the Use of Exercise_
There are no quality studies incorporated into this analysis.
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: exercise, exercising, physical activity, radial nerve entrapment, radial tunnel syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 94 in Scopus, 0 in CINAHL, 7 in Cochrane Library, 16,630 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Surgery**

**Recommendation:** Surgical Release for Subacute or Chronic Radial Nerve Compression Neuropathy

Surgical release is recommended for subacute or chronic cases of radial nerve compression neuropathy that persist despite other interventions.(60)

- **Strength of Evidence** – Recommended, Insufficient Evidence (I)
- **Level of Confidence** – Moderate

**Rationale for Recommendation**

There are no quality studies available on the efficacy of surgical intervention. There are no quality studies evaluating the efficacy of surgical intervention for distal radial neuropathies. However, clinically many patients respond well to surgery. Surgery is invasive, has adverse effects and is costly. It is recommended for select patients who failed trials of other non-operative treatments or if space occupying lesions are present.

**Evidence for the Use of Surgery**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: surgical release, surgery release, surgery, surgical procedures, radial tunnel release, radial nerve entrapment, radial tunnel syndrome; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 6 articles in PubMed, 97 in Scopus, 8 in CINAHL, 10 in Cochrane Library, 423 in Google Scholar, and 0 from other sources. We considered for inclusion 2 from PubMed, 2 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 4 articles considered for inclusion, 0 randomized trials and 2 systematic studies met the inclusion criteria.

---

**Non-Specific Hand, Wrist, and Forearm Pain**

**Diagnostic Criteria**

Non-specific pain is not a discrete diagnosis, per se, but the absence of a discrete diagnosis.

**Special Studies and Diagnostic and Treatment Considerations**

**RHEUMATOLGICAL STUDIES AND JOINT ASPIRATION**

1. **Recommendation:** Rheumatological Studies for Arthralgias

   Rheumatological studies are recommended for evaluation of patients with persistent unexplained arthralgias or tenosynovitis.

   - **Indications** – Persistent unexplained arthralgias or tenosynovitis.
   - **Frequency/Duration** – Repeat studies may be required after passage of time as some patients, particularly those with less severe diseases, tend to develop positive anti-bodies after months to years.

   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – High
2. **Recommendation: Arthrocentesis for Joint Effusions**

   **Arthrocentesis (joint aspiration)** of inexplicable joint effusions, particularly for evaluation of infections and crystalline arthropathies is recommended.

   **Indications** – Joint effusions without a clear diagnosis including suspected infection or crystalline arthropathies.

   **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
   **Level of Confidence** – High

**Rationale for Recommendations**

There are no quality studies evaluating rheumatological studies for evaluation of arthralgias; however, these studies have been helpful in diagnosing numerous rheumatological disorders. Arthrocentesis is also helpful for securing important diagnoses, such as septic arthritis and crystalline arthropathies.

**Evidence for the Use of Rheumatological Studies and Joint Aspiration**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Non-specific hand, wrist, and forearm pain, Arthrocentesis, Joint Effusion, Nonspecific, Hydrarthrosis, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 0 articles in PubMed, 9 in Scopus, 1 in CINAHL, 6 in Cochrane Library, 50 from Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**ELECTRODIAGNOSTIC STUDIES**

**Recommendation: Electrodiagnostic Studies to Evaluate Non-specific Hand, Wrist, or Forearm Pain in Patients with Paresthesias or Other Neurological Symptoms**

**Electrodiagnostic studies are recommended to evaluate non-specific hand, wrist, or forearm pain for patients with paresthesias or other neurological symptoms.**

**Indications** – Persistent tingling and pain, particularly symptoms characteristic of radiculopathies and entrapment neuropathies. Providers are cautioned that the prevalence rate of abnormal electrodiagnostic studies in asymptomatic populations are high (see CTS section above) and interpretations of abnormal findings should be cautious.

**Frequency/Dose** – Should generally be performed at least 3 weeks after symptom onset.

   **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
   **Level of Confidence** – Moderate

**Rationale for Recommendation**

There is 1 low-quality study evaluating electrodiagnostic studies for non-specific pain. However, electrodiagnostic studies may assist in diagnosing and treating the condition and thus are recommended.

**Evidence for the Use of Electrodiagnostic Studies**

There is 1 low-quality study in Appendix 2. (1128)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Electrodiagnostic, studies, Nerve conduction, study, NCS, Electromyography, EMG, Non-specific, hand, wrist, forearm, paint controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 31 articles in PubMed, 10870 in Scopus, 298 in CINAHL, 183 from Google Scholar, and 7 in Cochrane Library. We considered for inclusion 1 from PubMed, 0
from Scopus, 1 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 11358 articles
considered for inclusion, 1 randomized trials and 1 systematic studies met the inclusion criteria.

X-RAYS
Recommendation: X-rays for Evaluation of Non-specific Hand, Wrist, or Forearm Pain
X-rays are recommended for evaluation of cases in which non-specific hand, wrist, or forearm
pain persists.

Indications – Persistent non-specific hand, wrist, or forearm pain.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – High

Rationale for Recommendation
There is 1 moderate-quality study evaluating x-ray studies for non-specific pain. X-rays may assist in
diagnosing and treating the condition and thus are recommended.

Evidence for the Use of X-rays
There is 1 moderate-quality study incorporated into this analysis.(1129)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library
without date limits using the following terms: X-ray, Non-specific, HWF, pain; controlled clinical trial,
controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*,
randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective
studies. We found, reviewed and considered for inclusion 7 articles in PubMed, 332343 in Scopus, 0 in
CINAHL, 0 in Cochrane Library and 277000 in other sources. We considered for inclusion 1 from PubMed,
zero from Scopus, zero from CINAHL, zero from Cochrane Library and zero from other sources. Of the 1
articles considered for inclusion, 0 randomized trials and 0 systematic studies met the inclusion criteria.
## Initial Care

### RELATIVE REST

**Recommendation:** Relative Rest for Acute Non-specific Hand, Wrist, or Forearm Pain

Relative rest is a recommended treatment in select cases of acute non-specific hand, wrist, or forearm pain particularly where there are high ergonomic exposures (high force or high force combined with other risk factors).

**Strength of Evidence** – **Recommended, Insufficient Evidence (I)**

**Level of Confidence** – **Low**

**Rationale for Recommendation**

There are no quality studies and treatment is empiric. For patients with high ergonomic exposures, relative rest may be helpful. This intervention is not invasive, has low adverse effects, and for short periods is low to moderate cost, thus it is recommended.

**Evidence for the Use of Relative Rest**

There are no quality studies incorporated into this analysis.

---

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score</th>
<th>Number</th>
<th>Area of upper extremity</th>
<th>Diagnoses</th>
<th>Type of X-rays</th>
<th>CT used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Huellner 2013</td>
<td>Diagnostic</td>
<td>6.0</td>
<td>32</td>
<td>Hand and wrist</td>
<td>Non-specific hand or wrist pain.</td>
<td>Plain radiographs</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td></td>
<td></td>
<td>20 months and 16 months (group dependent)</td>
<td>x</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| | | | | | | | | | | | | | | | | |

Plain radiographs accuracy (25%-31%), sensitivity (24%-30%), and specificity (20%-60%). PPV (66%-76%). SPECT/CT diagnostic results in 44%-77% accuracy, 41%-74% sensitivity, and 60%-90% specificity. PPV (88%-98%). SPECT/CT resulted in the best imaging modality for non-specific hand and wrist pain. MRI showed better result when comparing typification of lesion. Data suggest interobserver agreement for imaging non-specific wrist pain via SPECT/CT good and only MRI better.
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: rest or relative rest, bed rest, nonspecific, non-specific, hand pain, wrist pain, and forearm pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 314 in Scopus, 0 in CINAHL, 2 in Cochrane Library, 34029 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**SPLINTS**

*Recommendation: Splinting for Acute or Subacute Non-specific Hand, Wrist, or Forearm Pain*

There is no recommendation for or against the use of splinting for treatment of acute or subacute non-specific hand, wrist, or forearm pain.

**Strength of Evidence – No Recommendation, Insufficient Evidence (I)**

**Level of Confidence – Low**

**Rationale for Recommendation**

There are no quality studies and treatment is empiric. Splinting may at times be helpful, but enforces debility, thus there is no recommendation for or against its use. It is generally not recommended for chronic use.

**Evidence for the Use of Splints**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: splint, splints or splinting; nonspecific, non-specific, hand pain, wrist pain, forearm pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 43 in Scopus, 0 in CINAHL, 9 in Cochrane Library, 8,360 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**ICE/HEAT**

*Recommendation: Self-application of Ice or Heat for Acute or Subacute Non-specific Hand, Wrist, or Forearm Pain*

Self-application of ice or heat is recommended for treatment of acute or subacute non-specific hand, wrist, or forearm pain.

**Strength of Evidence – Recommended, Insufficient Evidence (I)**

**Level of Confidence – Low**

**Rationale for Recommendation**

There are no quality studies and treatment is empiric. Self-applications of heat or ice are sometimes helpful. These interventions are not invasive, have low adverse effects, and are low cost, and thus are recommended.

**Evidence for the Use of Ice/Heat**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: ice, icing; nonspecific, non-specific, hand pain, wrist pain, forearm pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 11 in Scopus, 0 in CINAHL, 18 in Cochrane Library, 32,300 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: heat, heating, heat therapy, hot temperature; nonspecific, non-specific, hand pain, wrist pain, forearm pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 75 in Scopus, 0 in CINAHL, 45 in Cochrane Library, 269 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Follow-up Visits
Patients may require 1 to 3 appointments depending on the severity or the pain and need for workplace limitations.

Medications

**NSAIDS/ACETAMINOPHEN**

Recommendation: NSAIDs or Acetaminophen for Acute or Subacute Non-specific Hand, Wrist, or Forearm Pain

**NSAIDs or acetaminophen are recommended for control of pain associated with acute or subacute non-specific hand, wrist, or forearm pain.**

**Indications** – Acute or subacute non-specific hand, wrist, or forearm pain.

**Frequency/Duration** – Scheduled dosage rather than as needed is generally preferable.

**Indications for Discontinuation** – Resolution of pain, lack of efficacy, development of adverse effects particularly gastrointestinal.

- **Strength of Evidence** – Recommended, Evidence (C) – NSAIDs
- **Strength of Evidence** – Recommended, Insufficient Evidence (I) – Acetaminophen
- **Level of Confidence** – Low

**Rationale for Recommendation**

There are two moderate-quality studies evaluating the use of NSAIDs or acetaminophen for treatment of non-specific lower extremity pain(1038, 1039) which is presumably analogous to upper extremity pain and showed benefits (see Ankle and Foot Disorders Guideline). These medications may relieve pain and increase function. They are not invasive, have few adverse effects in employed populations, and are low cost. Thus, they are recommended.

**Evidence for the Use of NSAIDs/Acetaminophen**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDs, acetaminophen, non-steroidal anti-inflammatory, acetaminophen, ibuprofen, non-specific, hand, wrist, forearm, pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 83 in Scopus, 0 in CINAHL, 9 in Cochrane Library, 420 in Google Scholar, and 1 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 1 from other sources. Of the 3 articles considered for inclusion, 2 randomized trials and 1 systematic studies met the inclusion criteria.

**OPIOIDS**

See Opioids recommendations in Carpal Tunnel Syndrome section.
**Physical Methods**

**Recommendation:** Physical or Occupational Therapy for Acute, Subacute, or Chronic Non-specific Hand, Wrist, or Forearm Pain

There is no recommendation for or against the use of physical or occupational therapy for treatment of acute, subacute, or chronic non-specific hand, wrist, or forearm pain.

- **Strength of Evidence – No Recommendation, Insufficient Evidence (I)**
- **Level of Confidence – Low**

**Rationale for Recommendation**

There are no quality studies evaluating any of the physical or occupational therapy modalities for treatment of non-specific hand, wrist, or forearm pain. (A case series of hand rehabilitation with occupational therapy services suggested benefits of occupational therapy for patients with heterogenous disorders.) Thus, treatments administered are empiric. These treatments are not invasive, have few adverse effects, but are moderate to high cost depending on number of treatments. They are generally not indicated for initial treatment. They may be more reasonable for more persistent cases. Trials of these modalities may be helpful in cases that do not resolve with initial treatment methods outlined above. However, these treatments are empiric and thus the success may be limited. Thus, there is no recommendation for or against these modalities.

**Evidence for the Use of Physical or Occupational Therapy**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms physical therapy, occupational therapy, nonspecific, non-specific, hand pain, wrist pain, forearm pain; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 13 articles in PubMed, 172 in Scopus, 8 in CINAHL, 3 in Cochrane Library, 150 in Google Scholar and 0 in other sources. Zero articles met the inclusion criteria.

**EXERCISE**

Exercise is not generally indicated acutely. One moderate quality study of mostly chronic patients found no differences between two types of exercise programs, but had no control group.\(^\text{(1130)}\) Many patients with chronic findings, functional deficits and post-operative patients require some appointments to at minimum help institute a home exercise program. For those with residual deficits, particularly post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

**Evidence for the Use of Exercise**

There is 1 moderate-quality RCT incorporated into this analysis.\(^\text{(1130)}\)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library and Google Scholar without date limits using the following terms exercise, physical activity, non-specific Hand, Wrist, Forearm Pain, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 14 articles in PubMed, 38 in Scopus, 1 in CINAHL, 3 in Cochrane Library, and 437 in Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 in Google Scholar and 0 from other sources. Of the 1 articles considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.
van Eijsden-Besseling 2008  
RCT  
Sponsored by Research Stimulation Fund of University Hospital Maastricht, Institute for Rehabilitation Research, Hoensbroek, The Netherlands. No mention of COI.

<table>
<thead>
<tr>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.0</td>
<td>N = 88 with non-specific upper limb disorders; Mean age PE group 33.3±7.7 and SFE group 34.8±7.7. PE group Gender, M:F (19:25); SFE group Gender M:F (19:25)</td>
<td>Postural exercise group. Received 6 postural therapy sessions first 3 weeks, then tapered to 3 sessions in 3 weeks, 2 sessions in 2 weeks, then home exercise (n = 44) vs. strength/fitness exercise group. Received 9 strength/fitness therapy sessions first 3 weeks, then tapered to 6 sessions in 3 weeks, 2 sessions in 2 weeks, and finally home exercise (n = 44). Follow-up at baseline and months 3, 6, and 12.</td>
<td>No significant difference in decrease in pain between the groups at 3 months (0.6 cm, 95% CI 0.0 to 1.2), 6 months (0.2, 95% CI – 0.3 to 0.7), or at 12 months (0.1, 95% CI –0.6 to 0.8)</td>
<td>“Postural exercises showed no additional benefits to recovery when compared to strength and fitness exercise. Roughly 55% of patients reported being complaint free after one year.”</td>
<td>Data suggest no significant differences between types of exercises (comparable efficacy). Some baseline differences in groups for potentially compromising comparability.</td>
</tr>
</tbody>
</table>

**Surgery**  
Not applicable.

**Scaphoid Fracture**

**Diagnostic Criteria**

A clinical impression is made upon history of appropriate injury mechanism, physical examination findings of substantial tenderness particularly over the scaphoid tubercle. Findings of snuffbox tenderness, positive axial compression of thumb test, and effusion in the wrist (possibly echarmosis) should be sought. A fracture identified on imaging that includes a “scaphoid view” confirms that diagnostic impression. Fracture is not always confirmed on initial standard wrist x-rays, although those fractures identified later are by definition non-displaced and have good clinical outcomes with subsequent non-operative treatment. The differential diagnosis includes wrist sprain, undisplaced epiphyseal fractures of the distal portion of the radius in children, fracture of the hook of the hamate, avulsion fracture of the triquetrum, carpal instability, distal ulna subluxation, de Quervain’s tenosynovitis, radioscaphoid arthrosis, scapholunate dislocation, and tri-scaphoid arthrosis.

**Special Studies and Diagnostic and Treatment Considerations**

**X-RAY**

X-rays have been widely used as the first diagnostic test for scaphoid fractures.(285, 290, 1035)

1. **Recommendation: X-rays for Diagnosing Scaphoid Fractures**

   X-rays are recommended for diagnostic purposes that include at least 3 to 4 views including a “scaphoid view.”(71)

   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – High

2. **Recommendation: Follow-up X-rays for Scaphoid Fractures**

   Follow-up x-rays in 2 weeks are recommended for evaluation of potential scaphoid fractures,(1131) particularly for patients with a high clinical suspicion of fracture, but negative initial x-rays.
Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – High

Rationale for Recommendations
There are no quality studies evaluating x-rays for scaphoid fractures. However, x-rays have been used for decades to evaluate these fractures, identify those requiring surgical treatment, and to evaluate healing; thus, they are recommended to diagnose scaphoid fracture.

Evidence for the Use of X-rays
There are 7 moderate-quality studies incorporated into this analysis. (1164-1170) (Herneth 01)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: x-ray, scaphoid fracture, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, predictive value of tests, efficacy, efficiency, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 2 articles in PubMed, 934 in Scopus, 2 in CINAHL, 9 Cochrane Library, and 0 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 1 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 2 from other sources. Of the 3 articles considered for inclusion 3 diagnostic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score</th>
<th>Number</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>X-ray used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical Outcomes</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mallee 2011</td>
<td>Diagnostic</td>
<td></td>
<td>6.5</td>
<td>N = 34</td>
<td>Wrist</td>
<td>Suspected scaphoid fracture</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>Follow-up for 6 weeks. CT imaging resulted in a diagnosis of 20 fractures in 17 patients. For scaphoid fractures there was a sensitivity of 67% and specificity of 96% with an accuracy of 91% in depicting scaphoid fractures. MRI showed sensitivity of 67% for scaphoid fracture, specificity 89% and accuracy 85%.</td>
<td>“CT and MRI had comparable diagnostic characteristics. Both were better at excluding scaphoid fractures than they were at confirming them, and both were subject to false-positive and false-negative interpretations. The best reference standard is debatable, but it is now unclear whether or not bone edema on MRI and small unicortical lines on CT represent a true fracture.”</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Memarsadeghi 2006</td>
<td>Diagnostic</td>
<td></td>
<td>5.5</td>
<td>N = 29, mean age 34 years</td>
<td>Wrist</td>
<td>Wrist trauma accompanied by severe pain over scaphoid with negative radiograph.</td>
<td>Multi-detector with 4-detector row scanner</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>At 6-week follow-up with radiographs, 11 of 29 (38%) had scaphoid fracture; 8 had cortical fracture; 3 had trabecular involvement. MR imaging identified all 11 scaphoid fractures: 100% sensitivity and 100% specificity. 2 of 8 cortical fractures could be seen: 38% sensitivity; “Multi-detector CT is highly accurate in depicting occult cortical scaphoid fractures but appears inferior to MR imaging in depicting solely trabecular injury. MR imaging is inferior to multidetector CT in depicting cortical involvement.”</td>
<td>Small sample. Data suggest similar performance efficacy between CT and MRI for occult scaphoid fracture detection, but CT superior for cortical involvement detection.</td>
<td></td>
</tr>
</tbody>
</table>
| Fotiadou 2011 | Diagnostic | 5.0 | N = 34
mean age 23 years | Wrist trauma, both acute and chronic. | 16 multislice rows CT scanner | + | + | - | - | - | - | - | In 21 of 22 general hospital patients, MRI method of choice following x-rays. CT performed in 1 case. At university hospital CT solely performed in 5/12 cases and was first method of choice in another 3 cases, followed by MRI. Bone injury detected in 17/34 cases. In 7/9 (77.8%) fracture not detected on initial radiographs. Ligament trauma identified solely on MRI in 11 patients. In 4 patients with both MRI and CT, CT revealed 2 fractures not | 100% specificity and 55% accuracy. Multidetector CT identified 8 cortical scaphoid fractures: 100% sensitivity/100% specificity. No trabecular fractures detected. MRI vs. CT p = 0.25 scaphoid fractures; p = 0.03 cortical involvement. | Small sample. Data suggest similar efficacy between CT and MRI but both with limitations. |
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Subjects</th>
<th>Fracture Type</th>
<th>Imaging Technique</th>
<th>Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temple 2005</td>
<td>4.5</td>
<td>Experimental</td>
<td>N = 11 cadavers</td>
<td>Wrist</td>
<td>Cadaveric wrists</td>
<td>Sagittal CT scans performed in longitudinal axis of scaphoid. Used GE LightSpeed 16-slice helical scanner. Slice thickness 0.625mm with reconstructions every 0.50mm (120 per kilovoltage, 80 milli-amps, and 0.5 seconds per rotation).</td>
<td>Median time from injury to CT scan 6 months and median time from injury to surgery 6 months. 20 had histologic avascular necrosis according to criteria established by Ficat. With CT increased radiodensity of proximal pole had strongest correlation with avascular necrosis (p = 0.004). Increased radio density of proximal pole significantly correlated with post-op union rates.</td>
</tr>
<tr>
<td>Smith 2009</td>
<td>4.5</td>
<td>Diagnostic</td>
<td>N = 31 mean age 29 at time of injury</td>
<td>Wrist</td>
<td>Scaphoid fracture</td>
<td>Pre-op CT scans performed in longitudinal axis of scaphoid. Used GE LightSpeed 16-slice helical scanner. Slice thickness 0.625mm with reconstructions every 0.50mm (120 per kilovoltage, 80 milli-amps, and 0.5 seconds per rotation).</td>
<td>&quot;Preoperatively longitudinal CT of scaphoid nonunion is of great value in identifying avascular necrosis and predicting subsequent fracture union.&quot;</td>
</tr>
<tr>
<td>Ilica 2011</td>
<td>4.0</td>
<td>Diagnostic</td>
<td>N = 54; mean age 22 years</td>
<td>Wrist</td>
<td>Clinically suspected scaphoid fracture with negative radiograph</td>
<td>MDCT with a 64-detector multislice system.</td>
<td>In 20 of 55 (36%) wrists, MRI identified 22 fractures: 16 scaphoid fractures. MDCT identified 19 fractures in 17 of 55 (30%) wrists. 3 fractures missed: 2 scaphoid</td>
</tr>
<tr>
<td>Herneth 2001</td>
<td>Diagnostic</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------------</td>
<td>------------</td>
<td>----------------------------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0</td>
<td></td>
<td>N = 15 (7 male and 8 female) with acute wrist trauma had scaphoid fractures. Age range 15.8 – 55.2.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wrist trauma and scaphoid fractures</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>High-spatial resolution 10-5-MHz probe</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>9 or 60% of the 15 patients with acute wrist trauma had scaphoid fractures. At high-spatial-resolution US, 7/9 or 78% had positive results, and 22% false negative. 8/9 or 89% had clinical signs of scaphoid fractures, 3/6 or 50% had false positive results, and 1/9 or 11% had false-negative results. Sensitivity of high-spatial-resolution US in depicting scaphoid fractures was 78%, and the specificity was 100% vs with 56% and 100% obtained for conventional</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>fractures. MDCT 100% specificity, 86% sensitivity, 100% PPV, and 91% NPV.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>“High-spatial-resolution US is a reliable diagnostic tool for the evaluation of occult scaphoid fractures and should be considered an adequate alternative diagnostic tool prior to computed tomography or MR imaging.”</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Small sample size. Data suggest high spatial resolution US “may” assist in diagnosing scaphoid fractures when conventional radiography is negative for fractures.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
MRI
MRI has been used for the diagnosis of scaphoid fractures.(1138-1140)

Recommendation: MRI for Diagnosing Scaphoid Fractures
MRI is moderately recommended for diagnosis of occult scaphoid fractures when clinical suspicion remains high despite negative x-rays.(1133, 1141-1145)

Indications – Clinical suspicion of scaphoid fracture but negative x-rays.

Strength of Evidence – Moderately Recommended, Evidence (B)
Level of Confidence – Low

Rationale for Recommendation
MRI is not required for the majority of scaphoid fractures. However, for patients with a clinical suspicion of scaphoid fracture, but negative x-rays, current treatment recommendations are generally to splint the hand, thus often necessitating prolonged lack of use and lost productivity. A moderate-quality study has reported cost effectiveness of MRI to diagnose occult scaphoid fractures and reduce lost productivity for those without x-ray imaging evidence of fractures.(1146) Two moderate quality studies have suggested comparable results between CT and MRI,(1133, 1137) although two other studies suggested CT was better to evaluate cortical involvement.(1132, 1134) Thus, as there is evidence to support its use among these select patients, MRI is recommended.

Evidence for the Use of MRI for Scaphoid Fracture
There are 30 moderate-quality studies incorporated into this analysis.(1164, 1165, 1169, 1171, 1172) Beeres, 2008 #3210, 1179-1203 (Mallee 11; Fotiadou 11; Tiel-van Buul 96; Bergh 15; Ilica 11; Bretlau 99; Hunter 97; Jorgsholm 13; Kitsis 98; Kusano 02; Moller 04; Raby 01; Lozano-Calderon 06; Larribe 14) There are 6 low-quality studies in Appendix 2.(1027, 1173, 1204-1207) (Imaeda 92; Sharifi 15; Gaebler 96; Seneviratna 13; Schmitt 11)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Scaphoid Fracture, Magnetic Resonance Imaging, MRI, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 267 articles in PubMed, 762 in Scopus, 22 in CINAHL, 2 in Cochrane Library, and 1940 from Google Scholar. We considered for inclusion 10 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 29 from other sources. Of the 40 articles considered for inclusion 36 diagnostic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brooks 2005</td>
<td>RCT</td>
<td>6.5</td>
<td>N = 37 (24 female/13 male) suspected scaphoid fractures in 5 hospitals. Age for MRI and Control: 35.0 (27-41) and 29.0 (24.75-50).</td>
<td>MRI group (n = 11) vs. Control group (n = 17).</td>
<td>$44.37 (Australian) per day saved from unnecessary immobilization by use of MRI. Early MRI improved date of confirming diagnosis by 7 days, Day 3 vs. Day 10 (p = 0.003). When only subjects diagnosed as having no fracture included in analysis, median number of days unnecessarily in plaster in MRI group 3 days, which is significantly less than median of 10 days in control group (p = 0.006).</td>
<td>&quot;Use of MRI in the management of occult scaphoid fracture reduces the number of days of unnecessary immobilisation and use of healthcare units.&quot;</td>
<td>Study may be biased toward justification of early MRI in universal health care models.</td>
</tr>
<tr>
<td>Ng 2013</td>
<td>Diagnostic</td>
<td>7.0</td>
<td>N=35 patients (34 male, 1 female) Mean age: 27.4±9.4 years</td>
<td>Scaphoid fracture delayed-union or non-union who underwent surgery within 12 months of imaging.</td>
<td>Unenhanced MRI vascularity at surgery (sensitivity/specificity/PPV/NPV/accuracy): impaired 70/48/35/80/54; fair 25/74/11/89/69; poor 67/76/36/92/74. Contrast MRI vascularity at surgery (sensitivity/specificity/PPV/NPV/accuracy): impaired 56/64/36/80/62; fair 25/73/11/88/68; poor 60/93/60/93/88. DCE MRI vascularity at surgery (sensitivity/specificity/PPV/NPV/accuracy):</td>
<td>&quot;Comparative relative enhancement in the proximal scaphoid fragment with that in the distal fragment of DCE MRI improved diagnostic accuracy for assessment of proximal fragment vascularity in scaphoid delayed and non-union compared to non-contract and contrast-enhanced MRI examination.&quot;</td>
<td></td>
</tr>
</tbody>
</table>
Low 2005
Diagnostic study:

- 70
- N=50 patients (40 males, 10 females)
- Mean age: 29 years
- Hand: Scaphoid fracture
- 0.2T dedicated extremity system
- Observer agreement: 2 fractures identified by all 4 observers; 3 fractures by 3 observers; 5 fractures by 2 observers; incorrectly judged as normal 13 times by 4 observers. Observers saw poor sensitivities (11/9/43/49) and low NPV (31/30/39/40) but good specificities (93/93/87/80). 

Gäbler 2001
Diagnostic study:

- 65
- N=121 patients (77 males, 44 females)
- Mean age: 30.3±13.2 years
- Hand: Occult scaphoid fracture
- 1.0 T unit and circular surface coil
- MRI injury detection: none in 39 patients, injuries detected 112 in 82 patients. 10 days after injury: of 62 patients with MRI detectable injuries 39 diagnosed correctly and another 7 partially correct. 24 days after injury: 14 patients with MRI-detectable injuries, correct diagnosis in 6 cases and partially correct in 2, another 6 cases were diagnosed as negative which was incorrect. All 28 scaphoid fractures were correctly identified.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Study Design</th>
<th>Participants</th>
<th>Symptoms</th>
<th>Imaging</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unay 2009</td>
<td>Diagnostic 6.5</td>
<td>187 (29 males, 12 females)</td>
<td>History of fall on outstretched hand and tenderness upon palpation of anatomical snuffbox and scaphoid tubercle without angulation</td>
<td>1.5 T superconductor</td>
<td>diagnosed correctly; occult fractures diagnosed after a mean of 14.9±9.3 days. Negative diagnosis correctly achieved after mean 12.2±5.12 days. No false-positives in study</td>
<td></td>
</tr>
<tr>
<td>Mallee 2011</td>
<td>Diagnostic 6.5</td>
<td>N = 34 patients (25 males, 15 women) with suspected scaphoid fracture</td>
<td>Presence of sharp lucency line within trabecular bone pattern, break in continuity of cortex, sharp step in cortex, or dislocation</td>
<td>Follow-up for 6 weeks. CT imaging resulted in a diagnosis of 20 fractures in 17 patients. For scaphoid fractures there was a sensitivity of 67% and specificity of 96% with an accuracy of 91% in depicting scaphoid</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Data suggest comparable between CT and MRI for suspected scaphoid fractures.

Follow up include only 34 patients of original 40.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Patients</th>
<th>Hand</th>
<th>Fracture Type</th>
<th>MRI Field Strength</th>
<th>MRI Coil</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>Accuracy</th>
<th>Interval (days)</th>
<th>Fractures Represented</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patel 2013</td>
<td>6.0</td>
<td>N=91</td>
<td>Hand</td>
<td>Occult scaphoid fractures</td>
<td>1.0T Philips Inter a using C3 surface coil</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>42 days</td>
<td>Scaphoid fractures: MRI 3, control group 4. Normal MRI scan: MRI 28.9% vs. control 84.6% (p=0.03). Mean±SD clinical fracture appointment: MRI 1.1±0.5 vs. control 2.3±0.8 (p=0.001). Mean±SD plain radiographs: MRI 1.2±0.8 vs. control 1.7±1.1 (p=0.03). Mean±SD perceived effect of injury (MRI vs. control): day 42, work 0.6±0.9 vs. 1.2±1.6 (p=0.03); hindrance 6.3 vs. 4.9 (p=0.03). Early MRI in occult scaphoid fractures is marginally cost saving compared with conventional management and may reduce potentially large societal costs of unnecessary immobilisation.</td>
</tr>
<tr>
<td>Fox 2010</td>
<td>6.0</td>
<td>N=29</td>
<td>Wrist Scaphoid fracture</td>
<td>1.5 tesla MRI scan</td>
<td>1.5 tesla MRI scan</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>42 days</td>
<td>The mean interval from the date of MRI to the date of surgery was 54 days. When comparing the MR and surgical findings, there were 6 true positive results, 17 true-negative results, 1 false-negative result, 0 false-positive results. T1-weighted unenhanced MRI is an acceptable alternative to delayed contrast-enhanced MRI in the preoperative assessment of the vascular status of the proximal pole of the scaphoid in patients with chronic fracture nonunions.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Type</td>
<td>Study Setting</td>
<td>MRI Results</td>
<td>E-MRI Results</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>------</td>
<td>---------------</td>
<td>-------------</td>
<td>---------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fowler 1998</td>
<td>98</td>
<td>Diagnostic</td>
<td>N=45 patients (21 males, 22 females) with acute trauma and clinical symptoms of scaphoid fractures. Mean age: 32 years.</td>
<td>MRI results showed 100% sensitivity and 100% specificity while Bone Scintigraphy showed 83% sensitivity and 95% specificity. MRI was found to be more effective than Bone Scintigraphy for the diagnostic potential for scaphoid fractures. MRI has increased convenience for the patient and no use of radiation.</td>
<td>MRI results showed 100% sensitivity and 100% specificity while Bone Scintigraphy showed 83% sensitivity and 95% specificity. MRI was found to be more effective than Bone Scintigraphy for the diagnostic potential for scaphoid fractures. MRI has increased convenience for the patient and no use of radiation.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bretlau 1999</td>
<td>99</td>
<td>Diagnostic</td>
<td>N=52 patients (27 males, 25 females) Mean age: 44</td>
<td>E-MRI detected occult fractures of the scaphoid in 9 patients, and of the distal radius in a further 6 patients. All these fractures were confirmed at follow-up radiographs. Furthermore, E-MRI revealed a fracture of the capitate bone in 1 patient, and of the triquetrum in 2 patients, and in 8 patients, bone bruise in 1 or more of the carpal bones. However, these fractures and bone lesions could not be confirmed by the follow-up radiographs.</td>
<td>E-MRI seems to be better than radiographs in the early diagnosis of occult</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

MRI results showed 100% sensitivity and 100% specificity while Bone Scintigraphy showed 83% sensitivity and 95% specificity. MRI was found to be more effective than Bone Scintigraphy for the diagnostic potential for scaphoid fractures. MRI has increased convenience for the patient and no use of radiation.
The agreement between the two examiners was high (kappa = 0.8) for E-MRI detection of fractures.
<table>
<thead>
<tr>
<th>Study</th>
<th>Gender and age not mentioned</th>
<th>Wrist</th>
<th>Scaphoid fracture</th>
<th>Not mentioned</th>
<th>+</th>
<th>-</th>
<th>+</th>
<th>-</th>
<th>-</th>
<th>Not mentioned</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lozano-Calderon 2006</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>CT scans had a interobserver reliability value of 0.44 (95% CI = 0.16 – 0.44, p&lt;0.001) compared to the radiography value of 0.16 (95% CI = 0 – 0.25, p&lt;0.01). CT had a sensitivity of 72% (95% CI = 58-87%), specificity of 80% (95% CI =72%-87%) and an accuracy of 77% (95% CI = 70%-83%). Radiography had values of 75% (95% CI = 67%-88%), 64% (95% CI =52%-70%), 68% (95% CI = 60%-74%), respectively. However, when both viewed at the same time, the sensitivity increased (80% (95% CI = 70%-94%) while the specific and accuracy decreased (73% (95% CI = 65%-89%) and 75% (95% CI = 67% - 82%), respectively).</td>
</tr>
<tr>
<td>De Zwart 2012</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Among 319 rated MRI scans 247 were diagnosed with no injury, 13 with scaphoid fracture, 23 with other fracture and 36 as a bone</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>“This study suggests that computed tomography scans are useful for ruling out displacement but not for diagnosing it.”</td>
</tr>
</tbody>
</table>
Mean age: 28 years.

Based on these data, the specificity of MRI was estimated as 95.9%.

scans from healthy volunteers. MRI is not an adequate reference standard for true fractures among patients with suspected scaphoid fractures.

The mean interval between MRI and surgery was 0.7 days. 4 of the 6 necrotic fragments were correctly classified into the necrotic group and 2 patients into the viable group. The was a sensitivity of 67%, specific of 67%, positive predictive value of 50% and a negative predictive value of 80%.

"Our data are consistent with previously reported data supporting contrast-enhanced MRI for assessment of viability, and showing that dynamic imaging with time–intensity curve analysis does not provide additional predictive value over standard delayed enhanced imaging for acute scaphoid fracture."

"[I]t may be worth considering early application of MRI in the diagnostic algorithm of skeletally immature patients sustaining wrist trauma. A normal initial MR has a negative predictive value of 100% as early as 2 days after injury, whereas clinical and radiographic findings are not as reliable; also, scaphoid fractures may be identified on MR earlier than on radiographs in many patients. Additionally, MRI identified a large number of other injuries of both osseous and soft tissue structures."
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Sample Size</th>
<th>Mean Age</th>
<th>Wrist Location</th>
<th>Imaging Modality</th>
<th>Fracture Line</th>
<th>Fracture Evidence on MRI</th>
<th>Follow-up CT</th>
<th>Treatment Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kusano 2002 Diagnostic</td>
<td>5.5</td>
<td>N=52 patients (32 males, 20 females) with suspected scaphoid fracture. Mean age: 36.7</td>
<td>MRI (0.2 T)</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>In 18 of the 53 wrists, fracture was detected on MRI. Fracture was also found in the distal end of the radius in 11 patients and in the capitate in one patient. A bone contusion was found in the distal end of the radius in two patients. A fracture line was found in 13 of 16 diagnosed scaphoid fractures via CT.</td>
</tr>
<tr>
<td>Fotiadou 2011 Diagnostic</td>
<td>5.0</td>
<td>N = 34 mean age 23 years</td>
<td>Wrist trauma, both acute and chronic.</td>
<td>MRI method of choice following x-rays.</td>
<td>CT performed in 1 case. At university hospital CT solely performed in 5/12 cases and was first method of choice in another 3 cases, followed by MRI. Bone injury detected in 17/34 cases. In 7/9 (77.8%) of patients with fracture evidence on MRI but without a fracture line on the initial CT, did well without surgery and demonstrated evidence of a healed fracture on the follow-up CT. The drawback of MRI and CT examination is its high cost; however, it may avoid unnecessary treatment or decrease treatment period and thus reduce total expense.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Small sample. Data suggest similar efficacy between CT and MRI but both with limitations.

No mention of gender.
Ligament trauma identified solely on MRI in 11 patients. In 4 patients with both MRI and CT, CT revealed 2 fractures not found on MRI.

Of 195 patients, 99 (51%) had normal MRI results, 20 (10%) showed carpal or distal radius bone bruising. 74 patients (38%) were diagnosed with fractures, 37 (19%) with scaphoid fractures and 28 (14.4%) with distal radius fractures.

"MRI can now justifiably be regarded as the gold standard investigation for clinical scaphoid fracture. Using MRI we have determined that the incidence of occult scaphoid fracture is 19%. MRI enables the correct diagnosis to be reached early and by directing appropriate patient management prevents the unnecessary overtreatment of the majority of patients thus bringing both health and economic benefits."

Low-field MRI showed a high incidence of fractures in patients with posttraumatic radial wrist tenderness and demonstrated more fractures than radiographs and CT. A scaphoid fracture was by far the most common injury. However, it is not clear whether diagnosis of subtle injuries only...
found in 71 wrists. The most commonly found fracture combinations were that of the scaphoid and distal radius, followed by scaphoid and capitae fracture. The sensitivity of radiographs for visualization of scaphoid fractures was 70% and the specificity was 98%. Radiographic sensitivity for other fractures was less than 60%. The sensitivity of CT for visualization of scaphoid fractures was 95%, and between 75% and 100% for other fractures. MRI revealed 9 wrists with bone edema in the scaphoid and capitate.

Møller 2004
Diagnostic
No mention of sponsorship or COI.

<table>
<thead>
<tr>
<th>WRIST Scaphoid Fracture T1w and STIR coronal 3 mm thickness</th>
<th>+</th>
<th>-</th>
<th>+</th>
<th>-</th>
<th>+</th>
<th>-</th>
<th>-</th>
</tr>
</thead>
</table>

The MRI radiographers reported 43 scaphoid fractures, whereas the radiologist ultimately diagnosed only 36 scaphoid fractures (16.1% of patients) (sensitivity, 100%). It is possible to provide an acute MRI service to patients with clinically suspected fracture of the scaphiod and a normal plain radiograph. The MR images can be primarily read by sufficiently trained MR radiographers. This new work-up protocol reduces the cost for society.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Gender</th>
<th>Diagnosis</th>
<th>Wrist</th>
<th>Imaging Procedure</th>
<th>Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tiel-vanBuul 1996</td>
<td>4.5</td>
<td>16</td>
<td>(11 males, 5 females)</td>
<td>Clinical suspected scaphoid fracture</td>
<td>Wrist</td>
<td>Clinical 3-phase radio nuclide bone scintigraphy was obtained after 72 hours following trauma using 200 MBq 99m Tc-methylene diphosphonate</td>
<td>MRI only available for 16 of 19 patients. X-ray also performed. Bone scintigraphy positive in 7 for scaphoid fractures while MRI only positive in 5.</td>
<td>Small sample size. Data suggest MRI not superior to 3-phase bone scan for scaphoid fracture detection.</td>
</tr>
<tr>
<td>Tibrewal 2012</td>
<td>4.5</td>
<td>137</td>
<td>(79 males, 57 females)</td>
<td>Suspected scaphoid fracture</td>
<td>Wrist</td>
<td>1.5 T scanner N/A</td>
<td>37 (27%) MRI exams normal, 59 (43.4%) diagnosed with soft tissue injuries. 17 (12.5%) resulted in scaphoid fractures and 30 (22%) resulted in...</td>
<td>&quot;MRI should be regarded as the gold standard investigation for patients in whom scaphoid fracture is suspected clinically.&quot;</td>
</tr>
<tr>
<td>Hunter 1997</td>
<td>Diagnostic No mention of sponsorshi p or COI.</td>
<td>4. 5</td>
<td>N=36 patients (28 males, 8 females) with wrist trauma injury suspected of scaphoid fracture. Mean age: 26</td>
<td>WR IST</td>
<td>Scaphoi d Fractur e</td>
<td>Sign a 1.5-T MR imag er with a phas ed-array coil.</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

MR imaging revealed 22 occult fractures in 20 patients. Thirteen of these 22 fractures were in the scaphoid bone, and 9 were in the distal radius. On MR images, 16 patients had no evidence of fracture. Follow-up radiographs were available in 17 of the 20 patients who had occult fracture revealed by MR imaging. Eleven of the 13 occult fractures of the scaphoid bone were followed up (2 lost to follow-up), and three of these showed evidence of healing fracture. Three patients without MR evidence of a fracture had follow-up radiographs that showed no fracture. Three patients had findings consistent with bone contusion on MR images; in two patients, the contusion was associated with other
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>n</th>
<th>Gender</th>
<th>Mean Age</th>
<th>Wrist Location</th>
<th>Imaging Modality</th>
<th>Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beeres 2008</td>
<td>4.0</td>
<td>Diagnostic</td>
<td>79</td>
<td>43 males, 36 females</td>
<td>41 years</td>
<td>Wrist</td>
<td>Scaphoid fractures</td>
<td>1.5 Tesla MR scanner</td>
<td>The pairwise and overall k statistic was 0.67 (0.44-0.90) for inter-observer variation for a scaphoid fracture. The intra-observer variation was calculated for 38 patients, and the k statistic was 0.96 (0.69-1.0) for a scaphoid fracture.</td>
</tr>
<tr>
<td>Ilica 2011</td>
<td>4.0</td>
<td>Diagnostic</td>
<td>54</td>
<td>54 males, 0 females</td>
<td>22 years</td>
<td>Wrist</td>
<td>Scaphoid fracture with negative radiograph</td>
<td>MDC T with a 64-detector multi slice system</td>
<td>In 20 of 55 (36%) wrists, MRI identified 22 fractures: 16 scaphoid fractures. MDCT identified 19 fractures in 17 of 55 (30%) wrists. 3 fractures missed: 2 scaphoid fractures. MDCT 100% specificity, 86% sensitivity, 100% PPV, and 91% NPV.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Diagnostic</td>
<td>N</td>
<td>Patients</td>
<td>Age</td>
<td>Wrist trauma or fracture</td>
<td>Imaging Protocol</td>
<td>Results</td>
<td></td>
</tr>
<tr>
<td>------------</td>
<td>------</td>
<td>------------</td>
<td>---</td>
<td>----------</td>
<td>-----</td>
<td>--------------------------</td>
<td>-----------------</td>
<td>---------</td>
<td></td>
</tr>
<tr>
<td>Querellou</td>
<td>2014</td>
<td>4.0</td>
<td>57</td>
<td>patients (26 males, 31 females) with unilateral acute carpal trauma, hand pain or wrist pain. Mean age: 34 years</td>
<td>H/W/F</td>
<td>Wrist trauma occult fractures</td>
<td>1.5-T Scanner (Magnetom Avento 1.5 T; Siemens)</td>
<td>+ - + - + - + +6 months</td>
<td></td>
</tr>
<tr>
<td>Bergh</td>
<td>2015</td>
<td>4.0</td>
<td>125</td>
<td>patients (68 males, 56 females) with clinically suspected scaphoid fracture. Mean age: 30 years</td>
<td>Wrist</td>
<td>Scaphoid fractures</td>
<td>Dual-headed gamma camera with built-in CT</td>
<td>- ? - - + - + -</td>
<td></td>
</tr>
</tbody>
</table>

26 presented wrist and hand fractures through SPECT/CT; 26 presented positive results for wrist and hand fractures or bruising during MRI scans; 17 had discordant results between MRI and SPECT/CT in regards to bruising vs. fracture diagnoses.

“This study highlights that bone scintigraphy associated with SPECT/CT is a very useful and sensitive imaging technique to depict occult wrist fracture in patients with carpal trauma. Its interest is to allow the detection of these specific fractures and reduces the secondary risks such as nonunion. When carpal occult fracture is strongly suspected clinically, SPECT/CT might be proposed as a sensitive follow-up examination.”

7 diagnosed scaphoid fractures in MIR group vs. 4 in control group. For patients without fractures, those in MRI group used cast for fewer days (1 day) vs. control group (mean 14 days) (p <0.001). MRI group also had less days on sick leave than controls; 7 vs. 15 (p = 0.002).

“In a Norwegian setting, an early MRI was of value in patients with clinically suspected scaphoid fracture and normal plain radiographs.”
| Bhat 2004 | 4.0 | N=50 with fractures of waist of scaphoid. Age not given. | Wrist | Isolated fracture of waist of scaphoid. | 1.5 Tesla | N=50 | - | - | - | + | + | Assessments of both observers showed: sensitivity of 100%, specificity of 74%-87%, negative predictive value of 100%, and accuracy of 76%-88% for predicting nonunion, but less satisfactory positive predictive values (20% and 33%). Assessment of displacement on scaphoid series of radiographs had sensitivity between 33%-47% and positive predictive value between 27%-86%. Correct identification of displaced fractures from plain radiographs by both observers no more than 33%-47%. |
| Breitenseher 1997 | 4.0 | N=42 patients (23 males, 19 females) with clinical suspicion of scaphoid fracture after acute wrist injury. Mean age: 30.5±13.8 years | Wrist | Acute wrist injury | 1.0-T unit | N/A | + | + | - | - | + | MI depicted occult fractures of scaphoid bone in 14 or 33%; capitate bone in 4 or 10%; and trapezium in 1 patient (5%). Sensitivity and specificity for detection of radiographically occult fractures of wrist; 100%, and 95% and 100% for second |

"[T]he assessment of displacement of scaphoid fractures on MRI can probably be used to assess the likelihood of union although the small number of nonunions limits the power of the study."
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Patients (Sex)</th>
<th>Methodology</th>
<th>Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kitsis 1998</td>
<td></td>
<td>0</td>
<td>N=22 patients (9 males, 13 females) Mean age:34</td>
<td>WRIST Scaphoid Fracture</td>
<td>The MRI scan was on a Picker Vista 0.5 tesla knee coil</td>
<td>Eight patients had no bone injury in either the MRI or the bone scan. Three scaphoid fractures were found on the MRI and the bone scan and one scaphoid fracture was diagnosed with bone scanner and not on the MRI.</td>
</tr>
<tr>
<td>Raby 2001</td>
<td></td>
<td>0</td>
<td>N=56 patients</td>
<td>WRIST Scaphoid Fracture</td>
<td>0.2T extremity MR system. Spin echo T1 and STIR T1 70</td>
<td>The early MR group had seven scaphoid, six radial and four other fractures. Management was altered in 89%. The late MR group had 14 scaphoid, nine radial and three other fractures. Management was altered in 69%. A cost model showed that overall costs are less with early rather than late scanning.</td>
</tr>
</tbody>
</table>

Sensitivities for detection of cortical fracture: 21%, 100%, and 14% (T1 and T2* sequences, respectively). Sensitivities for detection of bone marrow abnormality: 100%, 100%, and 59%, respectively.
HIGH-SPATIAL RESOLUTION SONOGRAPHY
High-spatial resolution sonography has been used to diagnose scaphoid fractures.(1162)

Recommendation: High-spatial Resolution Sonography for Diagnosing Scaphoid Fractures
High-spatial resolution sonography is recommended to diagnose occult scaphoid fractures when clinical suspicion remains high despite negative x-rays.

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – Moderate

Rationale for Recommendation
There are a few quality studies regarding the use of high-spatial resolution sonography to diagnose scaphoid fractures, with data suggesting reasonable reliability.(1162)

Evidence for the Use of High-Spatial Resolution Sonography
There are 4 moderate-quality studies incorporated into this analysis.(1170, 1177, 1208, 1209) (Fusetti 05; Hauger 02; Herneth 01; Tiel Van-Buul 93)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: high spatial resolution sonography, scaphoid bone, fractures, bone or scaphoid fractures, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 2 articles in PubMed, 2 in Scopus, 1 in CINAHL, 0 from Cochrane Library, and 418 from Google Scholar. We considered for inclusion 2 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 5 from Google Scholar, and 0 from other sources. Of the 7 articles considered for inclusion 3 diagnostic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest</th>
<th>Score</th>
<th>Number</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>Type of CT</th>
<th>X-ray Used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fusetti 2005</td>
<td>Diagnostic</td>
<td>No mention of sponsorship or COI.</td>
<td>6.5</td>
<td>N = 24 (11 female and 13 male) with clinically suspected fracture and normal radiographs.</td>
<td>Hand</td>
<td>Occult scaphoid fractures</td>
<td>MX-8000 16 Slices; High-spatial-resolution sonography (HSR-S)</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>10 (42%) presented high index of suspicion, 7 (29%) moderate index, and 7 (29%) a low index. RS effusion observed in 16 or 66% and STT effusion in 8 or 33%. Sensitivity / specificity / PPV/and NPV of HSR-S for early detection of occult SFs 100% (5/5), 79% (15/19), 56% (5/9), and 100% (15/15).</td>
<td>“HSR-S is a reliable, available, and cost-effective method in early diagnosis of occult fractures of the scaphoid.”</td>
</tr>
</tbody>
</table>

| Tiel-Van Buul 1993 | 5.5 | 160 male | Wrist | Scaphoid fracture | Scaphoid Radiography | - | - | + | - | - | + | Patients were reviewer | 35 patients showed evidence for a scaphoid fracture on the | “We advise scaphoid radiography” |

Copyright © 2016 Reed Group, Ltd.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Age Range</th>
<th>Gender</th>
<th>Diagnostic Information</th>
<th>Imaging</th>
<th>Results</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hauger 2002</td>
<td>2002</td>
<td>54 (35 males and 19 females) with clinically suspected scaphoid fracture and normal findings on initial radiographs, including specific scaphoid images. Age range 10 – 75.</td>
<td>10 – 75.</td>
<td>female</td>
<td>Mean age = 38.6</td>
<td>initial radiographs. Overall, 21 patients were positive for a scaphoid fracture, 24 was positive for other bone fractures, and 80 were negative. The bone scan revealed 41 patients with a scaphoid fracture, 49 with other bone fractures, and 41 negative results. No information about sensitivity and specificity were mentioned.</td>
<td>after at least one year. Initially, using at least four views.</td>
<td>High-resolution sonography is a reliable and accurate method of evaluating occult fractures of the scaphoid waist. Data suggest high spatial resolution sonography can be beneficial in diagnosing scaphoid fractures when plain radiographs are negative when there is a high index of suspicion for scaphoid fracture. However, findings support that cortical disruption is key in making the diagnosis.</td>
</tr>
</tbody>
</table>
CT imaging has been used to diagnose scaphoid fractures. (1163)

**Recommendation: CT Imaging for Diagnosing Scaphoid Fractures**

CT imaging is moderately recommended for diagnosing occult scaphoid fractures when clinical suspicion remains high despite negative x-rays. (1162, 1164, 1165) Quality studies include multiplanar reconstructive CT. (1165)

**Strength of Evidence** – Moderately Recommended, Evidence (B)

**Level of Confidence** – Moderate

**Rationale for Recommendation**

There are some quality studies regarding the use of CT to diagnose scaphoid fractures, although false positives occur. (1164) One comparative trial was unable to confirm CT as superior to bone scan. (1166) A retrospective case series study reported that 22% (n = 118) of patients with negative x-rays, but with clinical suspicion of scaphoid fracture, were confirmed positive by CT imaging. (1167) There are no studies comparing MRI with CT with bone scanning and no recommendation is made for one over the other. Two moderate quality studies have suggested comparable results between CT and MRI, (1133, 1137) although two other studies suggested CT was better to evaluate cortical involvement. (1132, 1134) For patients with continuing symptoms suggestive of scaphoid fracture, but absence of findings on repeat x-ray, CT scan has been reported to be an effective imaging technique. (1168, 1169) Therefore, CT imaging for those with clinical impression of fracture but negative x-rays is recommended.
Evidence for the Use of CT Imaging
There are 10 moderate-quality studies incorporated into this analysis.(1164-1166, 1208, 1211-1213, 1217-1219) (Mallee 11; Memarsadeghi 06; Ilica 11; Cruickshank 07)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: CT imaging, CT, CAT, scaphoid fracture, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, predictive value of tests, efficacy, efficiency, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 0 articles in PubMed, 20 in Scopus, 20 in CINAHL, 3 Cochrane Library, and 20 from Google Scholar. We considered for inclusion 0 from PubMed, 4 from Scopus, 3 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 4 from other sources. Of the 11 articles considered for inclusion, 10 diagnostic studies met the inclusion criteria.(1162, 1164-1166, 1170, 1171)

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Conflict of Interest (COI)</th>
<th>Number</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>Type of CT</th>
<th>X-ray Used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical Outcomes</th>
<th>Long-term follow-up</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adey 2007</td>
<td>Diagnostic</td>
<td>Sponsored by unrestricted research grants from AO Foundation, Small Bone Innovations, Smith and Nephew, Wright Medical, Biomet, and Joint Active Systems. No mention of COI.</td>
<td>7.0</td>
<td>N = 13 (gender not specified) with nondisplaced scaphoid fractures and 17 diagnosed with suspected fractures, average age 33 years</td>
<td>Han d</td>
<td>Non-displaced scaphoid waist fracture s</td>
<td>GE Lightspeed Qx/i CT Scanner; GE Medical Systems, Pewaukee, WI</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>Average sensitivity/ specificity/ and accuracy of CT for nondisplaced scaphoid fracture, for 1st round: 89% / 91% / and 90% 2nd round: 97% / 85% / and 88%. Positive predictive value or PPV for detection of radiographically occult scaphoid fractures with tomography of wrist 0.28 (95% CI, 0.23-0.32); NPV 0.99 (95% CI, 0.97-0.99).</td>
<td>Data suggest CT as better for ruling out fractures that result in due to relative infrequency of time fractures in patients with suspected scaphoid fractures.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fusetti 2005</td>
<td>Diagnostic</td>
<td></td>
<td>6.5</td>
<td>N = 24 (11 female and 13 male) with clinically suspected</td>
<td>Han d</td>
<td>Occult scaphoid fractures</td>
<td>MX-8000 16 Slices; High-spatial-resolution</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>10 (42%) presented high index of suspicion, 7 (29%)</td>
<td>HSR-S is a reliable, available, and cost-effective</td>
<td>Small sample size. Data suggest (HSR-S)</td>
<td></td>
</tr>
</tbody>
</table>
No mention of sponsorship or COI.

| Hanneman n 2013 | Diagnostic Sponsored by a research grant from the Netherlands Organisation for Health Research and Development. No COI. | 6.  | 5 | N = 44 (10 female/34 male) with radiologically proven unilateral scaphoid fracture. Age over 18. | Hand | Proven unilateral scaphoid fracture | Multiplanar reconstruction CT | n sonography (HSR-S) | moderate index, and 7 (29%) a low index. RS effusion observed in 16 or 66% and STT effusion in 8 or 33%. Sensitivity / specificity / PPV and NPV of HSR-S for early detection of occult SFs 100% (5/5), 79% (15/19), 56% (5/9), and 100% (15/15). | method in early diagnosis of occult fractures of the scaphoid. * is reliable as well as cost effective method in early diagnosis of occult fractures of the scaphoid and this method is not without problems and CT is still superior. |

| Hanneman n 2014 Diagnostic | | 6.  | 5 | N = 102 ≥18 years | Hand | Randomized to: Group A Multiplanar reconstructed | - - + - - - + All views combined (transversal, coronal, and sagittal) for: no union, partial union, or union was moderate overall inter-observer agreement (k = 0.576) (95% CI: 0.396–0.753). Overall inter-observer agreement (k = 0.699, 95% CI: 0.529–0.870). Average sensitivity of multiplanar reconstruction CT was 73% and average specificity 80%. | Time to clinical union; median of 6 | [T]he addition of PEMF bone. * Data suggest multiplanar reconstruction CT is accurate and reliable in the diagnosis of union and non-union scaphoid. Wrist fractures with respect to partial union fractures is significant variation between observers. |
RCT Double-blind
No sponsorship or COI.

or active PEMF (n = 51) vs. Group B, or placebo (n = 51) Assessed functional and radiological outcomes (multiplanar reconstructed CT scans) at 6, 9, 12, 24 and 52 weeks.

CT (MRCT)

weeks (6-24, IQR 6-9) in group A vs. median of 6 weeks (6-52, IQR 6-9) in group B. The range of movement returned to normal at 12 week in both groups. Weighted mean inter observer agreement for union ($\kappa$ = 0.683, 95% CI 0.473 - 0.893) and nonunion ($\kappa$ = 0.791, 95% CI 0.599 - 0.984) for all CT scans, (p < 0.002).

Median time to radiologically confirmed union in group A was six weeks vs 12 weeks in group B, (p = 0.30).

Waist fractures proceeded to union earlier in group A vs B (median 12 weeks (6 to 12) vs 52 weeks (6-52), chi-squared test = 4.156, (p = 0.04).

growth stimulation to the conservative treatment of acute scaphoid fractures does not accelerate bone healing."

Mallee 2011

Diagnostic

6.5

N = 34

Wrist

Suspected scaphoid fracture

Presence of sharp lucent line within trabecular bone

- + + + - - +

Follow-up for 6 weeks. CT imaging resulted in a diagnosis of 20 CT and MRI had comparable diagnostic characteristics.

Data suggest comparable between CT and MRI for bone growth stimulation did not accelerate bone healing when compare to placebo.
| Cruicksand 6. 5 | 47 patients with suspected scaphoid fractures | Wrist | Scaphoid fracture | Siemens Somatom Volume Zoom (4 slice) for first 13 patients. Rest of patients were scanned with Siemens 64 slices machine. | + if patient continued to have stuff box tenderness and normal | - | - | - | + | 10 - 14 days post injury. Again at 7 days and 6-8 weeks if x-ray shows | CT had a positive predictive value of 100% (95% CI = 78%-100%) and the specificity was 100% (95% CI = 87%-100%). The negative predictive value for fractures are 96.7% (95% CI = 82%-100%) with a sensitivity of 94.4% (95% CI = 72%-100%) One fracture had a positive predictive value of 100% (95% CI = 78%-100%) and the specificity was 100% (95% CI = 87%-100%). The negative predictive value for fractures are 96.7% (95% CI = 82%-100%) with a sensitivity of 94.4% (95% CI = 72%-100%) One fracture had a positive predictive value of 100% (95% CI = 78%-100%) and the specificity was 100% (95% CI = 87%-100%). The negative predictive value for fractures are 96.7% (95% CI = 82%-100%) with a sensitivity of 94.4% (95% CI = 72%-100%) One fracture had a positive predictive value of 100% (95% CI = 78%-100%) and the specificity was 100% (95% CI = 87%-100%). The negative predictive value for fractures are 96.7% (95% CI = 82%-100%) with a sensitivity of 94.4% (95% CI = 72%-100%) One fracture had a positive predictive value of 100% (95% CI = 78%-100%) and the specificity was 100% (95% CI = 87%-100%). The negative predictive value for fractures are 96.7% (95% CI = 82%-100%) with a sensitivity of 94.4% (95% CI = 72%-100%) One fracture had a positive predictive value of 100% (95% CI = 78%-100%) and the specificity was 100% (95% CI = 87%-100%). The negative predictive value for fractures are 96.7% (95% CI = 82%-100%) with a sensitivity of 94.4% (95% CI = 72%-100%) One fracture had a positive predictive value of 100% (95% CI = 78%-100%) and the specificity was 100% (95% CI = 87%-100%). The negative predictive value for fractures are 96.7% (95% CI = 82%-100%) with a sensitivity of 94.4% (95% CI = 72%-100%) One fracture had a positive predictive value of 100% (95% CI = 78%-100%) and the specificity was 100% (95% CI = 87%-100%). The negative predictive value for fractures are 96.7% (95% CI = 82%-100%) with a sensitivity of 94.4% (95% CI = 72%-100%) One fracture | Both were better at excluding scaphoid fractures than they were at confirming them, and both were subject to false-positive and false-negative interpretations. The best reference standard is debatable, but it is now unclear whether or not bone edema on MRI and small unicortical lines on CT represent a true fracture. Data suggest early CT is reliable for diagnosing scaphoid fractures and other fractures of the wrist and carpal.

"CT has the potential to limit the need for immobilization for the majority of patients with clinical Scaphoid fracture, who do not actually have a fracture."
| Study       | Year | Participants | Intervention | Treatment 1 | Treatment 2 | | | | | |
|------------|------|--------------|--------------|-------------|-------------| | | | | |
| Clementson 2015 | 5    | N = 65       | Hand         | Operative    | Cast        | Fracture type            | x-ray | + | - | + | + | + | 24 fractures immobilized 5-8 weeks, 11 for 10-12 weeks, 4 for 13-16 weeks. 6-week CT scan demonstrated 27/30 or 90% of non- or minimally displaced fractures had united, linear association, (p = 0.47). In operatively treated group, 17 fractures immobilized in plaster for 2 weeks, 2 for 3-4 weeks, 5 for 6 weeks, 2 for 10 weeks; union rate at 6 weeks for non- or minimally displaced fractures 82%, dropping to 40% for severely displaced fractures. CT scan demonstrated 80% united at 6 weeks, increasing to 94% after 10 weeks. | "The majority of non- or minimally displaced scaphoid waist fractures are sufficient treated with 6 weeks in a cast." |

Data suggests most non- or minimally displaced scaphoid wrist fractures adequately treated for 6 weeks in cast. Screw fixation did not appear to shorter time to fracture union. Conservative treated fractures with prolonged time to union comminuted.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Age</th>
<th>Fracture Location</th>
<th>Injury Type</th>
<th>Methodology</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>Reference</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Memarsadghi 2006</td>
<td>5.5</td>
<td>29</td>
<td>34 years</td>
<td>Wrist</td>
<td>Wrist trauma</td>
<td>Multi-detector CT with 4-detector row scanner, 11 of 29 (38%) had scaphoid fracture: 8 had cortical fracture; 3 had trabecular involvement. MR imaging identified all 11 scaphoid fractures: 100% sensitivity and 100% specificity. 2 of 8 cortical fractures had positive bone scintigraphy and negative CT scan. CT galse negative in 5 and false positive in 1 patient. Bone scintigraphy has sensitivity of 93% (13/14) and a specificity of 91% (78/86). CT has sensitivity of 64% (9/14) and specificity of 99% (85/86).</td>
<td>13/14</td>
<td>85/86</td>
<td>At 6-week follow-up with radiographs, 11 of 29 (38%) had scaphoid fracture: 8 had cortical fracture; 3 had trabecular involvement. MR imaging identified all 11 scaphoid fractures: 100% sensitivity and 100% specificity. 2 of 8 cortical fractures had positive bone scintigraphy and negative CT scan. CT galse negative in 5 and false positive in 1 patient. Bone scintigraphy has sensitivity of 93% (13/14) and a specificity of 91% (78/86). CT has sensitivity of 64% (9/14) and specificity of 99% (85/86).</td>
<td>“Multi-detector CT is highly accurate in depicting occult cortical scaphoid fractures but appears inferior to MRI imaging in depicting solely trabecular injury. MR imaging is inferior to multidetector CT.”</td>
</tr>
</tbody>
</table>

Rhemrev 2010
Diagnostic
No sponsorship or COI.

5.5
N = 100 with clinically suspected scaphoid fracture.
Hand evaluated with CT within 24 hours after injury and bone scintigraphy between 3 and 5 days after injury.
Lightspeed Qx/I CT Scanner, Pewaukee, WI | - + + - + + | Significant difference in union rate between treatment groups at any measure point, (p = 1.00). 13 had positive bone scintigraphy and negative CT scan. CT galse negative in 5 and false positive in 1 patient. Bone scintigraphy has sensitivity of 93% (13/14) and a specificity of 91% (78/86). CT has sensitivity of 64% (9/14) and specificity of 99% (85/86). “In conclusion, this study confirms that bone scintigraphy remains the gold standard to date.” | Data suggest bone scan 3-5 deep laser superior to CT within 24 of accident. Timing is different not a head to head comparison. |
**Recommendation: Bone Scanning for Diagnosing Scaphoid Fractures**

Bone scanning is recommended to diagnose occult scaphoid fractures when clinical suspicion remains high despite negative x-rays. (1141-1145, 1173, 1174)

| Ilica 2011 Diagnostic | 4.0 | N = 54; mean age 22 years | Wrist | Clinicaly suspected scaphoid fracture with negative radiograph. | MDCT with a 64-detector multislice system. | + | + | - | - | - | - | In 20 of 55 (36%) wrists, MRI identified 22 fractures: 16 scaphoid fractures. MDCT identified 19 fractures in 17 of 55 (30%) wrists. 3 fractures missed: 2 scaphoid fractures. MDCT 100% specificity, 86% sensitivity, 100% PPV, and 91% NPV. | "MDCT offers highly accurate results, especially concerning cortical involvement, and is a useful alternative in facilities lacking MRI." | Data suggest MDCT useful in detecting cortical involvement, but not superior to MRI for scaphoid fracture detection. |
**Indications** – At least 48 hours after the injury with continuing clinic suspicion of scaphoid fracture. (1175)

**Strength of Evidence** – **Recommended, Evidence (C)**

**Level of Confidence** – Moderate

**Rationale for Recommendation**

There are few quality studies on bone scanning for scaphoid fracture and suggesting utility. (1173, 1175-1177) Bone scans are not required for evaluation of the majority of patients with scaphoid fractures; however, in those patients with a clinical suspicion of scaphoid fracture, but negative x-rays, bone scans may assist in securing an earlier diagnosis that may obviate prolonged splinting in those without a fracture. Thus, bone scans are recommended for these select patients.

**Evidence for the Use of Bone Scans**

There are 9 moderate-quality studies incorporated into this analysis. (1162, 1194, 1221-1223, 1225-1228) (Tiel van Buul 93; Murphy 95; Hiscox 14; Beeres 05; Beeres 07) There is 1 low-quality study in Appendix 2. (1224)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: bone scan, scaphoid fracture, scaphoid bone fracture, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 42 articles in PubMed, 85 in Scopus, 2 in CINAHL, 1 in Cochrane Library, and 96 from Google Scholar. We considered for inclusion 10 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and from 0 other sources. Of the 11 articles considered for inclusion 10 diagnostic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score</th>
<th>N</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>Type of Bone Scans</th>
<th>CT Used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes assessed</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rolfe 1981</td>
<td>Diagnostic</td>
<td>5.0</td>
<td>99</td>
<td>Hand</td>
<td>Recent history of carpal trauma, clinical signs suggestive of scaphoid fracture, no identifiable fracture on initial radiographic.</td>
<td>Isotope bone imaging (IBI)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Nielsen 1983 Diagnostic</td>
<td>4.5</td>
<td>100 (101 wrists)</td>
<td>Wrist Scaphoid fracture. Mean age 33 years.</td>
<td>99m-Tc_MDP wrist scintigraphy performed with a Nuclear-Chicago Pho/Gam ma 3 scanner.</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>2 months.</td>
<td>Scintigram result: 54 positive, 13 inconclusive, among which 25 fractures detected (11 were scaphoid).</td>
<td>99m-Tc-MDP wrist scintigraphy appears expedient to exclude scaphoid bone fracture, if performed in case of doubt after secondary clinical and radiographic assessment and guided by negative scintigrams, the number of clinical examination, radiographic scans and superfluous casting days are reduced.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td>Data suggest wrist bone scan highly sensitive but low specificity for scaphoid fractures.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Clinical suspect ed scaphoid fracture

3-phase radionuclide bone scintigraphy was obtained after 72 hours following trauma using 200 MBq 99mTc-methylenediphosphonate

-  +  +  -  -  +

72 hours after injury

MRI only available for 16 of 19 patients. X-ray also performed. Bone scintigraphy positive in 7 for scaphoid fractures while MRI only positive in 5.

We conclude that in the diagnostic management of patients with suspected scaphoid fracture and negative initial radiographs, the use of MRI may be promising, but is not superior to three-phase bone scintigraphy.

Small sample size. Data suggest MRI not superior to 3-phase bone scan for scaphoid fracture detection.

Recent history of carpal trauma, clinical signs suggestive of scaphoid fracture, no identifiable fracture on initial radiograph.

Three phase radionuclide bone scintigraphy (72 hours after injury)

-  -  +  +  -

1 day, 2 weeks, 6 weeks

A total of 152 scaphoid radiographs were available for interpretation. In 18 patients the initial radiographs were judged positive for scaphoid fracture, where as 60 patients had negative initial radiographs.

The best diagnostic strategy in the management of clinically suspected scaphoid fractures consists of initial radiography followed by bone scintigraphy in patients with negative radiographs.

Data suggest bone scan should be used only after failed radiographs. Bone scans should be used instead of multiple radiographs after a failing initial radiograph.
After 2 weeks, two more scaphoid fractures were recognized, and one additional scaphoid fracture was identified after 6 weeks. Bone scintigraphy was obtained in the 60 patients with initially negative radiographs and in 15 patients a “hot-spot” in the scaphoid region was seen.

<table>
<thead>
<tr>
<th>Murphy</th>
<th>1995</th>
<th>Diagnostic 7.0</th>
<th>99 males, 44 females</th>
<th>Male, Female</th>
<th>Mean age = 36</th>
<th>Hand and wrist</th>
<th>Clinical scaphoid fracture was defined as presence of “snuffbox tenderness” or pain on direct palpation of the anatomic</th>
<th>Three-phase technetium methylene diphosphonate bone scan</th>
<th>-</th>
<th>-</th>
<th>+</th>
<th>+</th>
<th>-</th>
<th>-</th>
<th>+</th>
<th>4 days, 14 days</th>
<th>Day 4 bone scans, when compared to the diagnosis made with a radiograph on day 14, had a sensitivity of 100%. “Day 4 bone scans are an accurate means of ruling out scaphoid fracture. However, because of a significant</th>
</tr>
</thead>
</table>

Data suggest bone scans performed on day 4 detect more wrist fractures of all types not just scaphoid fractures.
##### Wrist Patients with clinical scaphoid fractures based on acute wrist injury and snuffbox tenderness and normal radiographs.

| Mean age | 6.0 | 27 males, 11 females | Three-phase bone scan | 10 to 14 days, then 6 weeks, then 12 months | Mean number of days immobilized was 26 in radiograph/radiotracation group while the mean was 29 for bone scan/early diagnosis group. The current study suggests that the use of bone scans to help diagnose occult scaphoid fractures does not reduce the number of casted days for occult scaphoid fracture.

| Specifity of 92%, positive predictive value of 65%, negative predictive value of 100%, accuracy of 93%. | Small sample so study aim cannot be adequately answered. Data suggest comparable efficacy and bone scans do not appear to reduce the number of casted days for occult scaphoid fracture.
<table>
<thead>
<tr>
<th>Beeres 2005 Diagnosti c</th>
<th>5. 5</th>
<th>56 ma le, 26 fe ma le</th>
<th>Mean age = 38</th>
</tr>
</thead>
</table>

| Patients with suspect ed scaphoi d fracture that did not show on plain radiogra phs. Clinical signs of fracture include swollen and tender anatomical snuffbox. | Three-phase bone scan. Technetium-diphosph onate, Tc99m-HDP | - | - | + | + | - | + |

| Week 1, then wee k 6, and then mon th 3 | Bone scans show ed a fractur e in 38/56 patien ts. 15 fractur es were at the scaph oid bone. | “If there is a strong clinica l suspic ion of a scaph oid fractur e, which canno t be confir med by conve ntiona l radiol ogy, BS is a valua ble diagn |

Kapla n-Meier surviv al analy sis using the log- rank test reveal ed that there was no statistically signifi cant differe nce betw en days immo bilized betwe en the radiog raph and bone scan group s (p = 0.38). immo bilized and that the differe ntial diagn osis of occult scaph oid fractur es shoul d remai n broad becau se other injure s are comm on.”
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Gender</th>
<th>Mean Age</th>
<th>Location</th>
<th>Signs of Fracture</th>
<th>Imaging Procedure</th>
<th>Follow up</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beeres 2007 Diagnosti c</td>
<td>5.5</td>
<td>50</td>
<td>29 male, 21 female</td>
<td>Wrist</td>
<td>Acute trauma and suspected scaphoid fracture, Tender anatomical snuffbox and pain when applying axial pressure</td>
<td>Palmar and dorsal images after injection of 500 MBq of Technetium-99m diphosphonate (Tc99m-HDP)</td>
<td>- - + + - - +</td>
<td>Depending on injury and grouping – between two weeks and 24 weeks</td>
</tr>
<tr>
<td>Stordahl 1984 Diagnosti c Articles</td>
<td>4.0</td>
<td>30</td>
<td>31</td>
<td>Wrist</td>
<td>Clinical signs of fracture and either negative or non-diagnostic initial x-rays.</td>
<td>Radionuclide imaging, administration of 10-15 mCi 99mTc Dimethyl Phosphonate. We used a Pho/Gamma 4 Camera with 9 had focal increased activity on bone scan located on the scaphoid bone, 4 of these had</td>
<td>- - + + - - - -</td>
<td>Foll ow up at 2 and 6 weeks</td>
</tr>
</tbody>
</table>

Data suggest bone scintigraphy in combination with physical examination is the gold standard for diagnosing suspected scaphoid fractures when scaphoid radiographs cannot confirm the scaphoid fracture.
Initial Care

SPLINTS

Recommendation: Wrist Splinting for Scaphoid Tubercle Fractures

Wrist splinting is recommended for treatment of scaphoid tubercle fractures.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – High

Rationale for Recommendation

There are no quality studies evaluating wrist splinting for treatment of scaphoid tubercle fractures. However, clinical experience suggests splinting may suffice, as these fractures heal well due to adequate blood supply. (1229) (Symes 11) Splinting is not invasive, has few adverse effects, is low cost, and thus is recommended.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Splint, splinting, scaphoid fracture, Scaphoid Bone, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 8 articles in PubMed, 68 in Scopus, 3 in CINAHL, 1 in Cochrane Library, 95 in Google Scholar, and 0 from other sources. We considered for inclusion 3 from PubMed, 1 from Scopus, 0 from CINAHL, 1 from Cochrane Library, 1 from Google Scholar, and 0 from other sources. Of the 6 articles considered for inclusion, 0 randomized trials and 1 systematic studies met the inclusion criteria.

CAST IMMOBILIZATION

Casting has long been traditionally used as a primary intervention, with successful union being achieved 88 to 95% of the time. (1178) Typically, a Colles’ cast is recommended with the wrist in approximately 20° anatomic extension (functionally neutral posture), although many practitioners prefer a thumb spica cast. (402, 1179, 1180) High-risk scaphoid fractures should be promptly referred to hand or orthopaedic surgical specialists for definitive treatment because of the higher risk of these fractures developing a nonunion, malunion, or degenerative joint disease.

1. Recommendation: Wrist Casting for Stable Scaphoid Fractures

Immobilization of the wrist with casting is moderately recommended for treatment of documented stable scaphoid fractures which are displaced less than 1mm, are non-oblique, and do not include the proximal 1/3 of the scaphoid.

Indications – Stable documented scaphoid fractures that include fractures with any of these properties:
• Fragments displaced less than 1mm;
• Fragments are non-oblique;
• Fragment does not include the proximal 1/3 of the scaphoid.

_Frequency/Duration_ – Casting should be performed for 6 to 8 weeks, and then with the cast removed, imaging taken to assess healing.\(^{(402, 1131)}\)

_Strength of Evidence_ – Moderately Recommended, Evidence (B)
_Level of Confidence_ – High

**Rationale for Recommendation**
There is no quality evidence comparing casting to no immobilization for scaphoid fractures. However, in cadaveric studies there is a significant difference in angulation and rotation when comparing casting with no-casting. There are 6 moderate-quality studies that include casting as a treatment with effective results in achieving successful union reported.\(^{(401, 402, 1178, 1181-1183)}\) Casting is not invasive, has some associated stiffness, decreased grip strength, and atrophy due to disuse, and is of moderate cost; however, it is believed to be essential to healing. It also has been associated with lower rates of subsequent development of osteoarthrosis than operative fixation.\(^{(1184)}\) Thus, casting is recommended for treatment of stable scaphoid fractures.

2. **Recommendation: Use of Thumb Immobilization with Casting for Scaphoid Fractures**

There is no recommendation for or against concurrent immobilization of the thumb with the wrist for treatment of scaphoid fractures.

_Strength of Evidence_ – No Recommendation, Insufficient Evidence (I)
_Level of Confidence_ – Low

**Rationale for Recommendation**
There is significant debate whether or not the thumb should be immobilized along with the wrist. There is one moderate-quality study that found no advantage to using a thumb spica compared with a Colles’ cast in 392 patients.\(^{(1181)}\) Another study included thumb immobilization in both groups when comparing long and short arm casts to evaluate the effect of pronation and supination.\(^{(1182)}\) The authors concluded inhibition of pronation and supination during the first 6 weeks was beneficial. However, in a cadaveric model study, short arm casting was found to be just as effective as a thumb spica,\(^{(1185)}\) in eliminating displacement and rotation of the fracture. Thus, there is no evidence of improved healing rates or reduced rates of non-union between the two types of cast, although thumb immobilization markedly reduces function.\(^{(1146, 1181, 1186, 1187)}\)

3. **Recommendation: Colles’ Casting or Supportive Bandaging for Suspected but Radiographically Negative Scaphoid Fracture**

Colles’ casting or supportive bandaging is recommended for patients with suspicion of scaphoid fracture, but with negative x-rays.\(^{(1188)}\)

_Duration_ – 2 weeks, followed by cast removal, clinical examination, and re-x-ray.\(^{(1131, 1189)}\) (Leslie 81; Gumucio 89)

_Strength of Evidence_ – Recommended, Insufficient Evidence (I)
_Level of Confidence_ – Low

**Rationale for Recommendation**
The prognosis of occult fractures is thought to be very good as the fragments are by definition, well approximated.\(^{(288, 1131, 1190)}\) For patients with suspicion of fractures, but negative x-rays, either Colles’ casting or supportive bandaging\(^{(1188)}\) is recommended for 2 weeks, followed by cast removal, clinical examination, and repeat x-ray.\(^{(1131, 1189)}\) Reassessment in 2 weeks allows sufficient time for the fracture plane to be identifiable on repeat x-rays. Casting or splinting in 2 weeks is generally sufficient to prevent significant range of motion during the initial time the fracture would be healing and is recommended. If x-rays are again negative and symptoms persist, it is unlikely that there is a fracture and other diagnoses should be sought.

4. **Recommendation: Casting for High-risk Scaphoid Fractures**
Long-arm casting at 90° of elbow flexion is recommended for high-risk scaphoid fractures that are displaced 1mm or more,(1191, 1192) or fractures of the proximal 1/3 of the scaphoid and oblique fractures.(1131, 1189) It is recommended that high-risk scaphoid fractures be evaluated and treated by a specialist experienced in the management of these fractures.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
Scaphoid fractures are at a high risk for non-unions. High-risk scaphoid fractures have been treated surgically for many years as they tend to not heal well, thus fixation is believed to facilitate healing. While there are no quality studies supporting this belief, clinical experiences indicate superior results with this approach. Surgical intervention is invasive, has significant adverse effects including risk of non-union, and is costly. However, the risks of not operating appear higher and surgery is recommended. For non-displaced fractures, non-operative treatment is likely preferable, particularly as the long-term risk of osteoarthritis is lower.

Evidence for Casting with Thumb Immobilization for Scaphoid Fractures
There are 7 moderate-quality RCTs incorporated into this analysis.(1179, 1181-1183, 1186, 1193, 1194)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: cast immobilization, scaphoid fracture, Scaphoid Bone, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 29 articles in PubMed, 110 in Scopus, 11 in CINAHL, 15 in Cochrane Library, 6 in Google Scholar, and 0 from other sources. We considered for inclusion 29 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 0 from other sources. Of the 31 articles considered for inclusion, 7 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herbert Screws</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Saedén 2001</td>
<td>RCT</td>
<td>No sponsorship. No mention of COI.</td>
<td>5.0</td>
<td>N = 61 with 62 (49 males, 13 females) acute fractures of scaphoid. Mean±SD age 29±13 years.</td>
<td>Short arm cast (n = 30) vs. Herbert screws (n = 30). 12-year follow-up.</td>
<td>Patients treated by surgery working at time of injury on sick leave an average of 6-3 weeks vs. 15+10 weeks in conservatively treated group (p = 0.002, t = - 3.77). At 12-year follow-up, 90% surgical and 69% conservative groups reported no pain or wrist discomfort. Grip strength and ROM not different between groups. Radiographic evidence of osteoarthritis more common in surgical group (p = 0.049), but no difference in symptoms.</td>
<td>&quot;In our study the fractures united whether they were treated operatively or conservatively. Internal fixation of an acute fracture of the scaphoid allows early return to normal function and should be regarded as an alternative to conservative treatment in those patients who cannot accept immobilisation in a cast for three months or more, for sport, social or work-related reasons.&quot;</td>
<td>Randomization and allocation methods unclear. Surgery may result in faster recovery times and less time off work. However, surgery resulted in higher risk of arthritis.</td>
</tr>
<tr>
<td>Dias 2008</td>
<td>RCT</td>
<td>No COI. No mention of sponsorship.</td>
<td>4.5</td>
<td>N = 71 (62 males, 9 females) with fractured scaphoid. Mean (SEM) age fixation: 29.3 (16 to 50). Cast: 31.4 (16 to 61).</td>
<td>Herbert screw fixation (n = 35) vs. below elbow plaster cast immobilization (n = 36). Mean follow up was 93 months.</td>
<td>No statistical difference in symptoms and disability as assessed by mean Patient Evaluation Measure (p = 0.4), or mean Patient-Rated Wrist Evaluation (p = 0.9), mean range of movement of wrist (p = 0.4), mean grip strength (p = 0.8), or mean pinch strength (p = 0.4).</td>
<td>&quot;No medium-term difference in function or radiological outcome was identified between the two treatment groups.&quot;</td>
<td>Data suggest comparable efficacy between group outcomes comparing use of casts vs. surgical treatment of acute scaphoid fractures at 93 months.</td>
</tr>
<tr>
<td>Buijze 2014</td>
<td>RCT</td>
<td>No mention of sponsorship. No COI.</td>
<td>7.0</td>
<td>N = 62 (19 female, 43 male) with CT or magnetic resonance image-confirmed nondisplaced or minimally displaced fracture of scaphoid. Mean±SD age no thumb: 42±18 years. Thumb cast 33±14 years.</td>
<td>Below-elbow cast with inclusion of thumb (n = 31) vs below-elbow cast without inclusion of thumb (n = 31). Follow up at 10 weeks and 6 months.</td>
<td>Mean±SD extent of union (%) no thumb vs. thumb cast: 85±24 vs. 70±30, p = 0.048.</td>
<td>&quot;Immobilization of the thumb appears unnecessary for CT or magnetic resonance image-confirmed nondisplaced or minimally displaced fractures of the waist of the scaphoid.&quot;</td>
<td>Data suggest immobilization of thumb via casting for non-displaced and minimally displaced scaphoid wrist fracture is not beneficial as more union occurred in those without thumb casting via CT. Functional measures between groups comparable.</td>
</tr>
</tbody>
</table>
Cohen 2001  
RCT  
No mention of sponsorship or COI.  
4.0  
N = 200 with arm and leg injuries requiring cast support. Age and gender not reported.  
Standard cast consisting of synthetic or plaster of paris, vs. focused rigidity cast of synthetic material.  
Focused rigidity casting superior to traditional techniques for ability score (p = 0.0001), satisfaction score (p = 0.0023), overall impairment of function (p = 0.019), limitation of movement following cast removal (p = 0.024)  
“Compared with the standard technique, focused rigidity casting has been shown to be superior to traditional methods with regard to satisfaction and functional scores without any detriment to clinical results.”  
Data suggest increased patient satisfaction with FRC vs. conventional plaster of Paris cast with comparable efficacy.

<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>No.</th>
<th>Gender</th>
<th>Fracture Location</th>
<th>Cast Type</th>
<th>Methodology</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cohen 2001</td>
<td>RCT</td>
<td>4.0</td>
<td></td>
<td>Arm and leg injuries</td>
<td>Standard cast vs. focused rigidity cast</td>
<td>Superiority to traditional techniques for ability score, satisfaction score, overall impairment, and limitation of movement following cast removal.</td>
<td>Increased patient satisfaction with focused rigidity casting.</td>
</tr>
<tr>
<td>Gellman 1989</td>
<td>RCT</td>
<td>4.0</td>
<td></td>
<td>Scaphoid fractures</td>
<td>Long thumb spica vs. short-thumb spica cast</td>
<td>Fracture healing time shorter with long spica cast.</td>
<td>Compared with conventional plaster of Paris, focused rigidity casting resulted in faster fracture healing.</td>
</tr>
<tr>
<td>Clay 1991</td>
<td>RCT</td>
<td>4.0</td>
<td></td>
<td>Scaphoid fractures</td>
<td>Colles' cast vs. scaphoid cast</td>
<td>No difference in non-unions or cast tolerance or functional outcomes.</td>
<td>Both types of cast were equally well tolerated and rehabilitation did not appear to be adversely affected by immobilisation.</td>
</tr>
<tr>
<td>Hambidge 1999</td>
<td>RCT</td>
<td>4.5</td>
<td></td>
<td>Scaphoid fractures</td>
<td>Colles'-type plaster cast in either 20° flexion vs. 20° extension</td>
<td>Nonunion not influenced by position of immobilization.</td>
<td>Position of wrist before casting is not important, rather, immobilization via casting is important.</td>
</tr>
</tbody>
</table>
Follow-up Visits
Duration of immobilization is typically 6 to 8 weeks to develop resolution of tenderness and for imaging evidence of healing.(402, 1131) After 6 to 8 weeks, the cast should be removed, imaging repeated, and casts reapplied for an additional 3 to 6 weeks, with a repeating process until evidence of fracture healing is documented. The average casting time for non-displaced fractures is 10 weeks,(402) with all expected to heal in 6 months.(401)

Medications
NSAIDs/ACETAMINOPHEN
Recommendation: NSAIDs or Acetaminophen for Scaphoid Fractures
NSAIDs or acetaminophen are recommended to control pain associated with scaphoid fractures.
Indications – Pain due to a scaphoid fracture.
Frequency/Duration – Scheduled dosage rather than as needed is generally preferable.
Indications for Discontinuation – Resolution of pain, lack of efficacy, development of adverse effects, particularly gastrointestinal.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
There is no quality evidence for or against the use of NSAIDs or acetaminophen for scaphoid fractures. These medications have been found useful in other musculoskeletal injuries and by inference may be efficacious for control of swelling and pain in the initial stages of injury, although some concerns about healing of bones have been raised. Other studies have suggested no delayed bone healing (see Distal Forearm Fractures section).

Evidence for the Use of NSAIDs/Acetaminophen
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDs, non-steroidal anti-inflammatory, acetaminophen, ibuprofen, scaphoid bone, scaphoid fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 4 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 80 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Physical Methods/Rehabilitation
1. Recommendation: Education after Cast Removal for Scaphoid Fractures
   Referral of select patients needing education after cast removal for scaphoid fractures is recommended.
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Low

2. Recommendation: Physical or Occupational Therapy after Cast Removal for Scaphoid Fractures for Patients with Functional Debilities
   Referral of patients with functional debilities or those unable to return to work for physical or occupational therapy after cast removal for scaphoid fractures is recommended.
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Moderate
3. **Recommendation: Physical or Occupational Therapy after Cast Removal for Scaphoid Fractures for All Other Patients**

Routine referral for physical or occupational therapy after cast removal for scaphoid fractures of otherwise healthy patients who are able to return to work is not recommended.

**Strength of Evidence – Not Recommended, Insufficient Evidence (I)**

**Level of Confidence – Low**

**Rationale for Recommendations**

There are no quality studies evaluating education or occupational or physical therapy for scaphoid fracture. (However, there are several studies showing this for various MSD outcomes when comparing formal therapy with a self-administered home exercise program – see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.) These therapies are generally unnecessary for most patients. However, some patients may need formal therapy with exercises if there are considerable impairments or a failure to progress after removal of the cast or splint. A few appointments for educational purposes for select patients are recommended. The number of appointments is dependent on the degree of debility, with one or two educational appointments appropriate for mildly affected patients. Patients with severe debility or those unable to return to work may necessitate 8 to 12 appointments that particularly include progressive strengthening exercises. Additionally, while routine use may be of limited benefit, those patients who have muscle weakness or other debilities may also derive benefit from therapy including self-training exercises, particularly if unable to return to work. Therefore, occupational or physical therapy is recommended for select patients.

**Evidence for the Use of Physical Methods/Rehabilitation**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Cast, Casts, Immobilization, Remove, Removal; scaphoid bone, scaphoid fractures, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 105 articles in PubMed, 15 in Scopus, 23 in CINAHL, 1 in Cochrane Library, 112 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 1 article considered for inclusion, 0 randomized trials and 1 systematic studies met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Physical, Therapy, Rehabilitation, scaphoid bone, scaphoid fractures, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 121 articles in PubMed, 65 in Scopus, 21 in CINAHL, 16 in Cochrane Library, 153 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Surgery**

Displaced fractures are believed to require surgical treatment with fixation, but there are no quality studies of displaced fractures. Surgical treatment of non-displaced scaphoid fractures has been evaluated in quality studies and there is no quality evidence of improved long-term outcomes with surgery.(402, 1183, 1195-1200) These studies generally indicate earlier, short-term functional recovery is achieved by surgery compared with prolonged casting.(401, 1183, 1195, 1197, 1198) A Swedish study also found higher costs among manual workers treated with casts due to longer periods of lost time.(1200) However, two quality studies, one with 10-year follow-up, demonstrated an 11-fold increased risk of scaphotrapezial osteoarthritis in those surgically treated with internal fixation compared with those...
Indications to surgically fix a scaphoid fracture are not well defined, and there is a suggestion that some patients are better candidates than others (e.g., widely displaced fragments, or requirement for earlier recovery such as in professional athletes). Quality evidence indicates operative treatment of non-displaced or minimally displaced scaphoid fractures provide no long-term benefit in functional outcomes, and results in significantly higher incidence of scaphotrapezial joint osteoarthritis. Until better quality evidence becomes available, the decision to surgically treat a non-displaced scaphoid fracture is a decision between the orthopedist and patient with a discussion suggested to include the benefits of earlier functional recovery versus the longer term risks of osteoarthrosis.

1. **Recommendation: Surgical Fixation of Displaced Scaphoid Fractures**
   Surgical fixation of displaced scaphoid fractures is recommended.
   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – High

   *Rationale for Recommendation*
   See above.

2. **Recommendation: Surgical Intervention of Non-displaced or Minimally Displaced Scaphoid Fractures for Patients Requiring Early Recovery**
   Surgical intervention of treatment of non-displaced or minimally displaced scaphoid fractures is recommended for patients requiring earlier functional recovery.
   - **Indications** – Patients with non-displaced or minimally displaced scaphoid fractures who cannot or do not wish to be treated with an attempt at non-operative treatment. This includes athletes. It also may include patients who are unable to work until the fracture is healed, thus electing to forego attempted non-operative management and its attended lower risk of later osteoarthrosis but longer course of immobilization in exchange for earlier return to work. There is no significant evidence that one technique, including bone grafting is superior to another. (1201-1205)
   - **Strength of Evidence** – Recommended, Evidence (C)
   - **Level of Confidence** – Low

3. **Recommendation: Surgical Intervention of Non-displaced or Minimally Displaced Scaphoid Fractures for All Other Patients**
   Surgical intervention for treatment of non-displaced or minimally displaced scaphoid fractures is not recommended for all other patients.
   - **Strength of Evidence** – Not Recommended, Evidence (C)
   - **Level of Confidence** – Low

   *Rationale for Recommendation*
   See above.

**Evidence for the Use of Surgery vs. Non-operative Treatment for Scaphoid Fractures**
There are 13 moderate-quality RCTs incorporated into this analysis. (401, 402, 1217, 1236, 1248-1250, 1253-1258) (Drac 14) There is one low-quality trial included in the Appendix 2. (1259) (Jeon 09)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Surgical Fixation, Surgery, Scaphoid fracture, scaphoid bone, fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 132 articles in PubMed, 343 in Scopus, 2 in CINAHL, 4 in Cochrane Library, 657 in Google Scholar, and 0 from other sources. We considered for inclusion 17 from PubMed, 5 from Scopus, 1 from CINAHL, 0 from Cochrane Library, 2 from Google Scholar, and 0 from other sources. Of the 25 articles considered for inclusion, 14 randomized trials and 2 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>McQueen 2008</td>
<td>RCT</td>
<td>7.5</td>
<td>60 (50 males, 10 females) patients between the ages of 17 and 65 with a Herbert type B1 or B2 fracture of the scaphoid. Mean age was 29.4 years.</td>
<td>Percutaneous fixation of the scaphoid within 14 days of injury using a standard Acutrak screw (Group 1, N=30) Vs. Colles cast with the thumb free (Group 2, N=30). Immobilization continued for at least 8 weeks, no patient was treated in a last for longer than 12 weeks. Follow-up for 1 year.</td>
<td>Mean decrease grip strength (%) (8 weeks/12 weeks/26 weeks/52 weeks): operative (10/3/-1/-2) v. non-operative (42/25/11/5), (p&lt;0.001) at week 8, 12, and 26, NS at 52. Mean decrease pinch strength (5): operative (9/4/0/-5) v. non-operative (29/15/3/1), (p&lt;0.001) at week 8, (p=0.012) at week 12, NS at week 26 and 52. Mean decrease range of movement (%): operative (11/6/3/2) v. non-operative (52/32/11/6), (p&lt;0.001) at weeks 8 and 12, (p=0.018) at week 26, NS at week 52. Mean Green/O’Brian score: week 8, operative (79) v. non-operative (39), (p&lt;0.001); week 12, operative (88) v. non-operative (56), (p&lt;0.001); week 26 (92 v. 78), (p=0.006); week 52 NS. Percentage good and excellent results: week 8 (52 v. 0), (p&lt;0.001); week 12 (68 v. 15), (p&lt;0.001); week 26 (81 v. 56), (p=0.055); week 52 (100 v. 88), (p=0.025).</td>
<td>“[O]ur study confirms earlier time to union and quicker return to work and sport with percutaneous screw fixation of nondisplaced fractures of the waist of the scaphoid.”</td>
<td>Effects of surgical intervention allowed earlier return to work or sport with faster mean time to union. There were no differences in function at 1 year.</td>
</tr>
<tr>
<td>Study</td>
<td>Follow-up Duration</td>
<td>Participants</td>
<td>Intervention</td>
<td>Outcome Measures</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-----------------------</td>
<td>--------------------</td>
<td>--------------</td>
<td>--------------</td>
<td>----------------------------------------------------------------------------------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vinnars 2008 RCT</td>
<td>10-yr</td>
<td>75 (58 males, 17 females) with acute nondisplaced or minimally displaced scaphoid fracture. Mean age was 30.5 years.</td>
<td>Nonoperative treatment with a cast (N=35) vs. Internal fixation with a Herbert screw (N=40). Follow up over 10 years.</td>
<td>All fractures united. A significant increase in prevalence of osteoarthritis in scaphotrapezial joint found in operatively treated group. No differences in subjective symptoms, as measured with limb-specific outcome scores found between two groups. No significant differences in range of motion, grip strength, changed hand dominance after injury, or return to same work after injury. Scaphotrapezial arthritis occurred in 1 patient in nonoperatively treated group and in 11 in operatively treated group (p = 0.005).</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Radiological outcome resportion: operative 0 v. non-operative 8, (p=0.02). Mean time to union (weeks): operative (9.2) v. non-operative (13.9), (p<0.001). Mean time to normal ADLs (weeks): full sports (6.4 v. 15.5), (p<0.001); full employment (3.8 v. 11.4), (p<0.001).

"This study showed that the primary benefit of operative treatment-(i.e., a short immobilization time and an early return to work) was transient. Our observation of an increased risk of osteoarthritis in the operatively treated group points to the importance of careful selection of patients who may benefit from operative treatment."

10-yr follow-up of non displaced scaphoid fracture suggests conservative management has equal long term functional outcomes and lower risk for scaphotrapezial arthritis.
member of their immediate families received payments or other benefits or a commitment or agreement to provide such benefits from a commercial entity. No commercial entity paid or directed, or agreed to pay or direct, any benefits to any research fund, foundation, division, center, clinical practice, or other charitable or nonprofit organization with which the authors, or a member of their immediate families, are
<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Patients</th>
<th>Method</th>
<th>Follow-up</th>
<th>Findings</th>
<th>Recommendations</th>
<th>Allocation Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dias 2005 RCT</td>
<td>6.5</td>
<td>N = 88 (79 males, 9 females) patients with a bicortical fracture of the scaphoid. Mean age was 29.5 years.</td>
<td>Internal fixation with Herbert screw (no cast) (N=44) vs. Below elbow cast with thumb free (Colles'). (N=44)</td>
<td>Follow up for 52 weeks.</td>
<td>Grip strength and range of motion better in operative group at 8 weeks, but differences disappeared by 12 weeks. No other significant differences in pain, patient evaluation, or return to work.</td>
<td>“Each fracture should be treated non-operatively in a functional cast. Surgical intervention should be offered only to the every few patients who cannot return to work in a cast, and such patients should be made fully aware of the risks and limited gains provided by acute fixation.”</td>
<td>Allocation, randomization details unclear.</td>
</tr>
<tr>
<td>Saedén 2001 RCT</td>
<td>5.0</td>
<td>N = 62 (49 males, 13 females) acute fractures of the scaphoid. Mean age was not provided.</td>
<td>Short arm cast (N=30) vs. Herbert screws group (N=32)</td>
<td>12-year follow-up.</td>
<td>Patients treated by surgery who were working at time of injury were on sick leave an average of 6 + 3 weeks compared with 15 + 10 weeks in conservatively treated group (p = 0.002, t = 3.77). At 12 year follow-up, 90% surgical and 69% conservative groups reported no pain or wrist discomfort. Grip strength and ROM not different between groups. Radiographic evidence of osteoarthritis more common in surgical group (p = 0.049), although no difference in symptoms.</td>
<td>“In our study the fractures united whether they were treated operatively or conservatively. Internal fixation of an acute fracture of the scaphoid allows early return to normal function and should be regarded as an alternative to conservative treatment in those patients who cannot accept immobilisation in a cast for three months or more, for sport, social or work-related reasons.”</td>
<td>Randomization and allocation methods are unclear. Surgery may result in faster recovery times and less time off work, although it may come at the expense of higher radiographic arthritic changes.</td>
</tr>
<tr>
<td>Bond 2001 RCT</td>
<td>5.0</td>
<td>N=25 (22 males, 3 females) full-time military personnel with acute nondisplaced scaphoid fractures</td>
<td>Cast immobilization (N=14) vs. fixation with a percutaneous cannulated Acutrak screw</td>
<td>Average time to fracture union: seven weeks in screw fixation vs. 12 weeks in cast immobilization, p=0.0003. Return to work: 8 weeks fixation group</td>
<td>“Percutaneous cannulated screw fixation of nondisplaced scaphoid fractures resulted in faster radiographic union and return to military duty compared with cast immobilization. The small sample size (n=25). Data suggest average time to fracture union in percutaneous screw fixation group was seven weeks.”</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Participants</td>
<td>Intervention</td>
<td>Outcome</td>
<td>Comments</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>--------------</td>
<td>-------------</td>
<td>---------</td>
<td>----------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Medicine and Surgery, Navy Department, Washington, DC, Clinical Investigatio n program.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>No mention of COI.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ed fracture of the scaphoid waist.</td>
<td>(Acumed, Beaverton, Oregon) (N=11). Follow up for 2 years.</td>
<td>vs. 15 weeks cast immobilization group, p=0.0001.</td>
<td>specific indications for and the risks and benefits of percutaneous screw fixation of such fractures must be determined in larger randomized, prospective studies.”</td>
<td>weeks compared to twelve weeks in cast group. Additionally, the time to return to work in surgical group was eight weeks compared to fifteen week in cast group. Both groups showed comparable results for grip strength, ROM and patients satisfaction at 2 years.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adolfsson 2001 RCT</td>
<td>No mention of COI or sponsorship.</td>
<td>N=53 (39 males, 14 females) with undisplaced fracture of the waist of the Scaphoid. Mean age of 31 (range, 15±75) years.</td>
<td>Immobilization in a below elbow plaster cast for 10 weeks. If no union cast immobilization was continued for another 6 Weeks (N=28) vs. Percutaneous Acutrak screw fixation (N=25). Follow up for up to 16 weeks if nonunion.</td>
<td>No statistically significant differences between the two treatment groups with regard to either the rate of union or the time to union.</td>
<td>“Acute percutaneous internal fixation of undisplaced scaphoid waist fractures using the Acutrak screw allows early mobilisation without adverse effects on fracture healing.”</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Data suggest comparable results between casting versus Acutrak screw insertion in terms of rate of or time to union. Patients with screw insertion had significantly better ROM at 16 weeks but no better grip strength.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clementso n 2015 RCT</td>
<td>Supported by grants from the Swedish Research Council (Medicine) and Funds from Region</td>
<td>N=38 with acute non- or minimally displaced scaphoid waist fracture. Mean age and gender were not provided.</td>
<td>Conservative treatment: below-elbow thumb spica cast, incorporating the thumb up to the interphalangeal joint (N=24) vs. arthroscopic screw fixation (N=14). Follow-up for 3 years.</td>
<td>ROM at 26 weeks: 88% fixation group vs. 97% conservative group; p=0.004.</td>
<td>“Non- and minimally displaced scaphoid waist fractures are best treated conservatively. Operative treatment may provide an improved functional outcome in the short term but at the price of a possible increased risk of arthritis in the long term.”</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Data suggest conservative treatment group (cast) had significantly better ROM at 26 weeks. No significant differences between grip and for pinch strength. Surgery group “may” provide improved short term functional outcomes but at 6 years radiography</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Skåne. No COI.

Vinnars 2008 RCT

In support of their research for or preparation of the article, one or more authors received, in any one year, outside funding or grants in excess of $10,000 from the Folksam research fund (Sweden) and the AFA research fund (Sweden)

7.0

75 (58 males, 17 females) patients with a scaphoid fracture that occurred less than 28 days before being seen and between the ages of 17 and 65. Mean age was 30.5 years.

Non-operative treatment: immobilization in a below-the-elbow scaphoid cast with the thumb in palmar abduction, the interphalangeal joint free, and the wrist in neutral or slight extension; cast worn for 6 weeks with option of an additional cast worn for another 2-4 weeks (N=42) vs.

operative treatment: used volar approach centered over the tubercle of the scaphoid, with minimal incision exposing only the scaphotrapezial joint, dorsal approach, or combined volar and dorsal approach; after surgery, application of well-padded short arm noncircumferential dorsal plaster splint with the thumb left free for 2 weeks (N=43). Follow-up for a median of 10 years.

There were no significant differences between groups for primary outcomes.

"This study did not demonstrate a true long-term benefit of internal fixation, compared with nonoperative treatment, for acute nondisplaced or minimally displaced scaphoid fractures."

10-yr follow-up of non displaced scaphoid fracture suggests conservative management has equal long term functional outcomes and lower risk for scaphotrapezial arthritis.

Bone grafting

Caporrino 2014 RCT

5.0

N=75 (71 males, 4 females) Vascularized bone grafting (VBG) using the 1, 2 intercompartmental

Mean±SD time to union: NVBG 69.7±15.1 days vs. "Although the VBG group attained earlier union, this may not be clinically meaningful, nor justify Patient blinding unclear. Data suggest VBG group achieved an
<table>
<thead>
<tr>
<th>Study</th>
<th>Fixation vs. Bone graft</th>
<th>Follow-up</th>
<th>Fixation</th>
<th>Bone graft</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Braga-Silva 2008</td>
<td>N = 80 (56 males, 24 females) with symptomatic scaphoid non-union pseudarthrosis of single wrist submitted for surgery. Dominant hand involved in 88% of cases. Mean age was 26 years.</td>
<td>Distal radius vascularised bone graft (n = 35) vs. iliac crest non-vascularised bone graft (n = 45). Mean±SD follow up radial grafts: 3.1±1.2 years. Iliac grafts: 2.6±1.6 years.</td>
<td>No statistically significant difference between two groups with regards to ranges of extension, flexion and ulnar deviation movements.</td>
<td>“The use of bone graft in the treatment of scaphoid nonunion has improved the prognosis, allowing an increase in the likelihood of painless bone consolidation and restoration of wrist function.”</td>
<td>Data suggest comparable results between techniques for grip strength and ROM post-op mean time 2.8 years. Union consolidation of fracture quickest in non-vascularized group.</td>
</tr>
<tr>
<td>Garg 2013</td>
<td>N=100 with scaphoid nonunion. Mean age Group 1: 32.4 years (25 males, 17 females). Group 2: 36.8 years (30 males, 16 females).</td>
<td>Internal fixation plus distal radius bone graft (Group 1: N=50) vs. iliac crest bone graft was used instead (Group 2: N=50). Follow up for 3 years.</td>
<td>Bone fusion was achieved in 87.1% of group 1 and 86.5% of group 2 patients. No p-value given. Mean time for union was 4.2 months in group 1 and 4.5 months in group 2. No p-value given.</td>
<td>“There is no advantage of the iliac crest over the distal radius graft to justify its greater morbidity.”</td>
<td>Data suggest comparable results between distal radius bone grafts vs. iliac crest bone graft for scaphoid nonunion.</td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Sample Size</td>
<td>Methods</td>
<td>Results</td>
<td>Conclusion</td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>-------------</td>
<td>---------</td>
<td>---------</td>
<td>------------</td>
</tr>
<tr>
<td>Ribak 2010</td>
<td>RCT</td>
<td>N = 86 with scaphoid nonunion. No mention of age or gender.</td>
<td>Vascularised bone graft from dorsal and distal aspect of radius (n = 46) vs. Conventional non-vascularised bone graft from distal radius (n = 40). Mean follow up in group 1 25.3 months. Group 2 22.5 months.</td>
<td>Vascularized bone graft achieved 89.1% bone fusion compared to 72.5% bone fusion rate in non-vascularised bone graft. (p = 0.024)</td>
<td>&quot;Vascularised bone grafting yields superior results and is more efficient when there is a sclerotic, poorly-vascularised proximal pole in patients in scaphoid nonunion. On the other hand, in patients with well vascularised fragments, either the vascularised or conventional technique can be used, depending upon the surgeon’s experience and preference.&quot;</td>
</tr>
<tr>
<td>Raju 2011</td>
<td>RCT</td>
<td>N=33 (27 males, 6 females) with non-union of the scaphoid. Mean age: 28 years.</td>
<td>Herbert screw fixation (n=11) vs. Matti Russe bone grafting (n=9) vs. Kohlman modification of vascularised muscle pedicle graft procedure (n=13). Mean follow-up duration was 28 Months.</td>
<td>Herbert vs. Matti vs. Kohlman: 8, 6 and 11 patients achieved scaphoid union after mean intervals of 17, 16, and 15 weeks. No p-values given.</td>
<td>&quot;The time to union was earliest in the Kohlman modification of vascularised muscle pedicle graft procedure, which is recommended for patients with old non-union (&gt;1 year) or proximal pole fractures.&quot;</td>
</tr>
<tr>
<td>Drac 2014</td>
<td>RCT</td>
<td>Supported by grant project IGA MZCR NS 9623-4/2008. No COI.</td>
<td>Group A- Palmar Percutaneous approach (N=36) Vs. Group B- Dorsal Limited Approach (N=36) Follow-up for 1 year after surgery.</td>
<td>There were no significant differences between Group A and Group B for flexion, extension, radial and ulnar deviation, grip strength, presence of persisting complaints, patient satisfaction or DASH score at any of the follow-up points (p&gt;0.05).</td>
<td>&quot;We found no advantage to the palmar percutaneous approach in the treatment of nondisplaced and minimally displaced scaphoid fractures type B2 compared to dorsal limited approach.&quot;</td>
</tr>
</tbody>
</table>

Data suggest comparable efficacy at one year.
ULTRASOUND/OSTEOGENIC PROTEIN ADJUVANT

1. Recommendation: Ultrasound with Bone Graft for Scaphoid Fractures

There is no recommendation for or against the use ultrasound to accelerate bone graft healing for scaphoid fractures.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation

Low intensity pulsed ultrasound has been evaluated for the treatment of fractures.(1260-1269) (Rubin 01, Riboh 12, Siska 08, Virk 12, Pounder 08, Barry 15, Nelson 03, Parvizi 05, Smoljanovic 07, Griffin 12) There is one moderate-quality RCT that reported earlier healing of muscle-pediculated bone graft after low intensity ultrasound treatment for 21 patients with scaphoid non-union with healing of a mean 38 days earlier.(1270) While the results are promising, they need replication with a larger sample size prior to recommendation.

Evidence for the Use of Ultrasound with Bone Graft for Scaphoid Fractures

There is 1 moderate-quality RCT incorporated into this analysis.(1270)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ultrasound, Osteogenic Protein Adjuvant, Scaphoid Fractures, Ultrasonography, Ultrasonic, Scaphoid Bone, bone fractures, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 18 articles in PubMed, 80 in Scopus, 0 in CINAHL, 4 in Cochrane Library, and 2,268 in Google Scholar. We considered for inclusion 1 from PubMed, 4 from Scopus, 0 from CINAHL, 1 from Cochrane Library, 5 from Google Scholar, and 0 from other sources. Of the 11 articles considered for inclusion, 1 randomized trials and 10 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Compariso n Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ricardo 2006</td>
<td>RCT</td>
<td></td>
<td>4.5</td>
<td>N = 21 with vascularized bone graft and internal fixation with k-wire</td>
<td>Ultrasound treatment vs. sham ultrasound</td>
<td>Daily 20 minute low intensity ultrasound treatment over scaphoid led to reduced time to overall (clinical and radiographic) healing by 38 days (average 56±3.2 days compared with 94±4.8 days; p&lt;0.0001).</td>
<td>All patients achieved fracture union (active and placebo groups), but compared with the placebo device (11 patients), the active device (ten patients) accelerated healing by 38 days (56±3.2) days compared with 94±4.8 days, p&lt;0.0001, analysis of variance.</td>
<td>Study suggests low intensity ultrasound treatment beneficial in improving healing time in this subset of patients undergoing bone graft with internal fixation.</td>
</tr>
</tbody>
</table>

2. Recommendation: Osteogenic Protein Adjuvant for Scaphoid Fractures

There is no recommendation for or against the use of osteogenic protein-1 for adjuvant treatment with bone grafting for scaphoid fractures.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation

There is a small trial of osteogenic protein-1 (BMP-7) for treatment of 17 patients with scaphoid non-union at the proximal pole included 3 arms comparing: 1) autologous iliac bone graft; 2) autologous iliac bone graft plus osteogenic protein-1; versus 3) allogenic iliac bone graft plus osteogenic protein-1.(1207) The study reported the following healing rates: sclerotic area at 3 months 138.3±15.1 versus 74.0±14.1
versus 103.6±13.2mm² respectively. However, the results need repeating in a larger sample size prior to a recommendation.

**Evidence for the Use of Osteogenic Protein Adjuvant with Bone Graft for Scaphoid Fractures**

There is 1 moderate-quality RCT incorporated into this analysis.(1207)

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bilic 2006</td>
<td>6.0</td>
<td>N = 17</td>
<td>Autologous iliac graft vs. Autologous iliac graft + osteogenic protein-1 (OP-1) vs. Allogenic iliac graft + OP-1</td>
<td>OP-1 improved performance of autologous graft healing (4 vs. 9 weeks in control). OP-1 improved functional performance of both groups vs. autologous graft alone. Sclerotic bone replaced by vascularized bone as assessed by CT 3 months after operation vs. 24 months after operation (sclerotic area mm²): Autograft only: 138.3±15.1* vs. 111.5±8.6; Autograft + OP-1: 74.0±14.1 vs. 31.7±6.8***; Allograft + OP-1: 103.6 ±13.2* vs. 55.6±11.7*** *p &lt;0.05 vs. before operation ***p &lt;0.05 vs. autograft only</td>
<td>“Recombinant human OP-1 supports proximal pole scaphoid non-union healing via increased bone vascularization and replacement of preexisting proximal pole sclerotic bone as a consequence of avascular necrosis. The addition of OP-1 to allogenic bone implant equalised the clinical outcome with the autologous graft procedure. Consequently the harvesting of autologous graft can be avoided.”</td>
<td>Small sample size; study suggests significant potential benefit from using OP-1 in healing time, functional improvement, and avoiding autologous grafting.</td>
</tr>
</tbody>
</table>

**Distal Phalanx Fractures and Subungual Hematoma**

**Diagnostic Criteria**

Diagnosis is evident from clinical suspicion, physical examination findings, and x-ray confirmation.

**Special Studies and Diagnostic and Treatment Considerations**

**X-RAYS**

**Recommendation:** X-rays for Diagnosing Tuft Fractures

**X-rays are recommended to diagnose tuft fractures.**

*Indication* – Clinical tuft fractures that do not involve the DIP joint do not require x-rays as they do not alter treatment. Still, the threshold for obtaining x-rays for those fractures is low in the event they may involve the joint.

*Frequency/Duration* – Obtaining x-rays once is generally sufficient. Follow-up x-rays are rarely indicated aside from complicated healing.

**Strength of Evidence** – **Recommended, Insufficient Evidence (I)**

**Level of Confidence** – **Moderate**

**Rationale for Recommendation**

There are no quality studies evaluating the use of x-rays for distal phalangeal/tuft fractures. X-rays may assist in identifying fractures and the magnitude of the involvement of the DIP joint surface, which if large enough may alter management in favor of surgery (see below). As this section of the digit is readily accessible for physical examination, patients may be treatable without x-rays as x-rays will not change
the management of tuft fractures that do not involve the joint. X-rays are recommended for assessment of fractures thought to involve the DIP joint.

Evidence for the Use of X-rays
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: X-Ray, radiography, radiograph, roentgenogram, Distal Phalanx Fractures, Tuft Fractures subungual hematoma, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 51 articles in PubMed, 46 in Scopus, 0 in CINAHL, 2 in Cochrane Library, and 382 from Google Scholar. We considered for inclusion Zero from PubMed, Zero from Scopus, Zero from CINAHL, Zero from Cochrane Library, Zero from Google Scholar, and Zero from other sources. Zero articles met the inclusion criteria.

MRI/CT/ULTRASOUND/BONE SCAN IMAGING
Recom mendation: MRI, CT, Ultrasound, or Bone Scanning for Diagnosing Tuft Fractures
MRI, CT, ultrasound, or bone scanning is not recommended for diagnosing tuft fractures.

Strength of Evidence – Not Recommended, Insufficient Evidence (I)
Level of Confidence – High

Rationale for Recommendation
There are no quality studies regarding the use of MRIs, CTs, ultrasound, or bone scanning for diagnosing tuft fractures. As x-rays are sufficient for diagnostic purposes, neither MRI, CT, diagnostic ultrasound, nor bone scanning is recommended.

Evidence for the Use of MRI/CT/Ultrasound/Bone Scan Imaging
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: MRI, CT, CAT, Ultrasound, Bone scan imaging, Distal Phalanx Fractures, Subungual Hematoma, Tuft Fractures, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 20 articles in PubMed, 10 in Scopus, 0 in CINAHL, 6 Cochrane Library, and 60 from Google Scholar. We considered for inclusion Zero from PubMed, Zero from Scopus, Zero from CINAHL, Zero from Cochrane Library, Zero from Google Scholar, and Zero from other sources. Zero articles met the inclusion criteria.

Initial Care
Tuft fractures are initially treated by caring for accompanying soft tissue injury and splinting of the finger to prevent further discomfort or injury. Reduction of the relatively uncommon significantly displaced fractures should be attempted with dorsal traction followed by immobilization in a volar splint. In the small percentage of patients, reduction cannot be achieved and referral to an orthopedic surgeon for consideration of pinning may be indicated.(77)

Management of subungual hematoma associated with a tuft fracture varies widely. There are no quality RCTs investigating the treatment ramifications of no treatment, trephination, nail plate removal, nail bed laceration repair, or conversion of closed fracture into open fracture.(75, 76, 1208-1218) As subungual hematoma is often associated with nail bed laceration, many practitioners promote removing the nail and repairing the nail bed to avoid future cosmetic defects.(76) The primary concern for this procedure is the potential to convert an underlying fracture into an open fracture.(75, 76, 1208-1218)

Tuft fractures associated with nail avulsion may require reduction of the nail plate under the eponychium, or removal if reduction cannot be performed. As with the removal of the nail for other conditions, the eponychial space should be preserved by packing with petroleum gauze cut in the shape of the nail to
prevent scarring of the nail bed and stunted nail growth. (1218) The nail or gauze should remain in place for 2 to 3 weeks to allow initial formation of a new nail plate. Full growth of the new nail takes approximately 4 to 5 months. Open fractures other than from subungual hematoma trephination of the distal phalanx require cleansing, debridement, and inspection for foreign bodies. Orthopedic assistance is usually not required for uncomplicated closures. Open fractures with extensive soft tissue damage frequently are associated with chronic pain and disability and generally require assistance from an orthopedic or hand surgeon.

1. **Recommendation: Trephination for Management of Subungual Hematoma**

   Trephination is recommended for management of subungual hematoma.

   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Moderate

2. **Recommendation: Nail Removal or Laceration Repair for Management of Subungual Hematoma**

   Nail removal or laceration repair is not recommended for the management of subungual hematoma.

   - **Strength of Evidence** – Not Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Low

**Rationale for Recommendations**

There are no quality studies regarding trephination or nail removal/laceration repair to manage subungual hematoma. (75, 76, 1208-1218) In a prospective study of 47 patients with subungual hematoma involving 50% or more surface area of the nail, a nail bed laceration was associated with fracture in 80% of patients. Thus, tuft fractures with subungual hematoma have a high likelihood of an associated laceration. (76) However, another prospective study of 48 patients with subungual hematoma, 42% of which had an underlying tuft fracture, were treated exclusively with electrocautery trephination. At 10-months follow-up post-trephination, there were no infectious or cosmetic complications despite not repairing the laceration or by converting a closed fracture into an open fracture. Another prospective study conducted in children with subungual hematoma compared nail removal and laceration repair to trephination over a 2-year follow-up and concluded that there was no justification based on absence of adverse clinical outcomes from trephination to perform nail removal and exploration. (1216) Each participant had a short course of antibiotics. There were no case reports found of osteomyelitis from trephination over hand tuft fracture, nor any reports of adverse cosmetic outcomes (nail scarring, permanent depression) in patients with finger nail bed laceration that were managed without suturing. Thus, the practice of avoiding trephination over distal phalangeal fracture to avoid creating an open fracture, or the practice of exploring and repairing nail bed lacerations associated with subungual hematoma appears unsupported by the available literature.

Trephination is most commonly accomplished with a hot cautery unit. Successful trephining with 29-gauge needle inserted below the nail plate reported, (1219) as well as fine point scalpel blade, surgical drill and laser have also been reported. (1209) Trephining gives good cosmetic and functional results. (1208)

**Evidence for the Use of Trephination and Nail Removal or Laceration Repair**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Trephination; nail removal; laceration repair (subungual hematoma) / Distal Phalanx Fractures and Subungual Hematoma, Tuft Fractures ;controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 6 articles in PubMed, 1 in Scopus, 0 in CINAHL, 1 in Cochrane Library, 180 in Google Scholar, and 1 from other sources. We considered for inclusion 3 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 1 from
other sources. Of the 6 articles considered for inclusion, 0 randomized trials and 2 systematic studies met the inclusion criteria.

**NSAIDs/ACETAMINOPHEN**

**Recommendation: NSAIDs or Acetaminophen for Tuft Fractures**

**NSAIDs or acetaminophen are recommended to control pain associated with tuft fractures.**

**Indications** – Pain due to tuft fracture.

**Frequency/Duration** – Scheduled dosage rather than as needed is generally preferable.

**Indications for Discontinuation** – Resolution of pain, lack of efficacy, development of adverse effects, particularly gastrointestinal.

- **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
- **Level of Confidence** – **Low**

**Rationale for Recommendation**

There is no quality evidence regarding the use of NSAIDs or acetaminophen to control pain associated with tuft fractures. However, these medications are thought to be effective for control of swelling and pain in the initial stages of injury, are not invasive, have low adverse effects, are low cost, thus they are recommended.

**Evidence for the Use of NSAIDs or Acetaminophen**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDS, Anti-Inflammatory Agents, Non-Steroidal, non-steroidal anti-inflammatory Agents, Non-Steroidal agents; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, and 0 in Cochrane Library, 719 in Google Scholar. Zero articles met the inclusion criteria.

**ANTIBIOTIC PROPHYLAXIS**

**Recommendation: Post-trephination Antibiotic Prophylaxis for Open Fractures**

There is no recommendation for or against the use of post-trephination antibiotic prophylaxis for open fractures.

- **Strength of Evidence** – **No Recommendation, Insufficient Evidence (I)**
- **Level of Confidence** – **Low**

**Rationale for Recommendation**

Antibiotic prophylaxis for open fractures is widely used. However, they may not be necessary for open phalangeal fractures as a quality study did not show evidence of improvements upon infection rates compared with aggressive irrigation and debridement as there were equal numbers of soft tissue infections and no cases of osteomyelitis in either group.(1220) However, the study appears underpowered to detect these relatively infrequent events. Use of antibiotics may be more strongly indicated for those with risks for infection, such as patients with diabetes mellitus. Thus, there is no recommendation for or against use of antibiotics and the threshold for use of antibiotics for prophylaxis is suggested to be low.

**Evidence for the Use of Antibiotic Prophylaxis**

There is 1 high-quality RCT incorporated into this analysis.(1287) (Stevenson 03) There is 1 low-quality RCT in Appendix 2.(1288) (Sloan 87)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Antibiotic prophylaxis, Distal Phalanx Fractures and Subungual Hematoma, Tuft Fractures; controlled clinical trial, controlled trials, randomized
controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 12 in Scopus, zero in CINAHL, and 2 in Cochrane Library. We considered for inclusion 2 from PubMed, zero from Scopus, zero from CINAHL, zero from Cochrane Library and zero from other sources. Of the 2 articles considered for inclusion, 2 randomized trials and zero systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stevenson 2003 RCT</td>
<td>8.5</td>
<td>N = 193 (159 males; 34 females) with an open fracture of the distal phalanx; Age range 16 – 88.</td>
<td>Antibiotic four times a day for five days (N = 98) vs Placebo four times a day for five days (N = 95). Follow-up 4 or 5 days, 14 days and 8 weeks following injury unless wound was healed and patient is asymptomatic.</td>
<td>Infection rate (antibiotic vs. placebo): 3% vs 4% (p&gt;0.05).</td>
<td>“[T]he addition of prophylactic flucloxacillin to thorough wound toilet and careful soft-tissue repair of open fracture of the distal phalanx confers no benefit.”</td>
<td>Data suggest no benefit of addition of prophylactic flucloxacillin for treating distal phalanx fractures compared to placebo.</td>
</tr>
</tbody>
</table>

**TETANUS IMMUNIZATION**

Recommendation: Tetanus Immunization Status for Open Fractures

For open fractures, it is recommended that tetanus immunization status to be updated as necessary.

**Indications** – Wounds that are not clean or burns if more than 5 years have elapsed since last tetanus immunization.(1221)

*Strength of Evidence – Recommended, Insufficient Evidence (I)*

*Level of Confidence – High*

**Rationale for Recommendation**

There are no quality studies of tetanus immunization updating for open fractures. However, these immunizations are widely used and believed to have been successful on a population basis in reducing risk of tetanus over many decades. Tetanus immunizations are minimally invasive, have low adverse effects, and are low cost. As the adverse effects of not immunizing may be fatal, tetanus immunization updating for open wounds is recommended. Wounds that are not clean or burns should require immunization if over 5 years since last immunization, rather than 10 years.(1221) Patients without a completed immunization series of 3 injections should receive tetanus immune globulin along with immunization.

**Evidence for the Use of Tetanus Immunization**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Tetanus immunization, Distal Phalanx Fractures and Subungal Hematoma, Tuft Fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 0 articles in PubMed, 10 in Scopus, 0 in CINAHL, 0 in Cochrane Library and 0 in other sources. Zero articles met the inclusion criteria.
IMMOBILIZATION

1. Recommendation: Protective Splinting of Distal Phalanx for Fractures
   Protective splinting of the distal phalanx to the PIP is recommended for fractures.(77, 78, 1222)
   Duration – Approximately 3 weeks.
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Moderate

2. Recommendation: Tight Circumferential Taping for Tuft Fractures
   Tight circumferential taping around the fingertip is not recommended for tuft fractures.
   Strength of Evidence – Not Recommended, Insufficient Evidence (I)
   Level of Confidence – Low

Rationale for Recommendations
There are no quality studies evaluating immobilization for fractures. In the closed crush fracture of the distal phalanx, the L-shaped Alumafoam splint placed on the volar aspect to protect the soft tissues has been considered the best treatment, although quality comparative trials are lacking. Splinting generally is maintained for approximately 3 weeks. Tight circumferential taping is not recommended due to potential to impair circulation. Volar splinting is not invasive, has few adverse effects, is low cost and is recommended.

Evidence for the Use of Immobilization
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Immobilization, Splinting, Tight, circumferential, taping, Distal, Phalanx, Tuft, Fractures, fracture, Subungual, Hematoma; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 0 in Scopus 0 in CINAHL, 1 in Google Scholar, and 1 in Cochrane Library. We considered for inclusion 15 from PubMed, 5 from Scopus, 11856 from CINAHL, 24 in Google Scholar, 91 from Cochrane Library, and 0 from other sources. Of the 11986 articles considered for inclusion, 0 randomized trials and 4 systematic studies met the inclusion criteria.

Follow-up Visits
Uncomplicated closed tuft fractures do not require follow-up, particularly if there is not a need for work and activity limitations. Two or three appointments may be required for gradual reduction in limitations. Patients should be advised that residual tenderness and hypersensitivity to cold temperatures may persist for 6 months in a more than half of all patients with this injury.(1223)

Prescription Medications
There are no quality studies of the use of other prescription medications in tuft fracture than already addressed in the Initial Care section. However, some patients may require pain medication, especially nocturnally, for the first few days.

Physical Methods/Rehabilitation

EXERCISE
Exercise is not indicated acutely. Few patients require exercise after recovery. For those with residual deficits, particularly post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

Recommendation: Routine Use of Physical or Occupational Therapy for Tuft Fractures
There is no recommendation for or against the routine use of physical or occupational therapy for treatment of tuft fractures.
   Strength of Evidence – No Recommendation, Insufficient Evidence (I)
   Level of Confidence – Low
**Rationale for Recommendation**

There are no quality studies of the use of physical or occupational therapy or other methods for tuft fractures, and these injuries rarely require therapy. Joint mobilization therapy may be useful for complicated injuries or post surgical fixation. A few appointments for purposes of teaching range of motion exercises for recovery of full motion may be rarely indicated, particularly for those with more severe injuries or those with a lack of improvement after removal of splints. However, the vast majority of patients with tuft fractures require no further treatment.

**Evidence for the Use of Physical or Occupational Therapy**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Exercise, Physical Therapy, Occupational Therapy, Distal Phalanx Fractures and Subungual Hematoma, Tuft Fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 12 articles in PubMed, 3 in Scopus, 0 in CINAHL, 2 in Cochrane Library, 167 in Google Scholar, and 0 in other sources. Zero articles met the inclusion criteria.

**Surgery**

Distal phalangeal diaphyseal fractures rarely require operative fixation, except those that are extremely displaced, unable to be reduced or are unstable. Retrograde percutaneous Kirschner-wire fixation is the preferred internal fixation technique.\(^{(87)}\)

**Evidence for the Use of Surgery**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Retrograde percutaneous Kirschner-wire fixation, Bone Wires, Distal Phalanx Fractures and Subungual Hematoma, Tuft Fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 6 in Scopus, 0 in CINAHL, and 12 in Cochrane Library, 136 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Of the 1 articles considered for inclusion, 0 randomized trials and 1 systematic studies met the inclusion criteria.

---

**Middle and Proximal Phalangeal and Metacarpal Fractures**

**Diagnostic Criteria**

Diagnosis is determined by clinical suspicion evident from history, physical examination findings and x-ray confirmation.

**Special Studies and Diagnostic and Treatment Considerations**

**X-RAYS**

*Recommendation: X-rays for Diagnosing Phalangeal or Metacarpal Fractures*

X-rays are recommended for diagnosing phalangeal or metacarpal fractures and should include three projections, including a posteroanterior, lateral, and oblique view. A true lateral projection isolating the involved digit is required.

Strength of Evidence – **Recommended, Insufficient Evidence (I)**  
Level of Confidence – **High**
Rationale for Recommendation
There are no quality studies evaluating the use of x-rays for phalangeal and metacarpal fractures. However, x-rays assist in identifying fractures, orientation of fracture plane(s), magnitude of the involvement of the interphalangeal and metacarpal phalangeal joints, which if large enough may alter management in favor of surgery (see below). X-rays are recommended for assessment of fractures of the phalanges and metacarpals.

Evidence for the Use of X-rays
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL and Cochrane Library without date limits using the following terms: X-Ray, Metacarpal, Middle, Proximal, Phalangeal, boxer's, Fracture, Bone, Diagnostic, Diagnosis, Sensitivity, Specificity, positive, predictive, value, negative, predictive, Predictive, Value, of, Tests, efficacy, efficiency. We found, reviewed and considered for inclusion 251 articles in PubMed, 2 in Scopus, 7 in CINAHL, 0 in Cochrane Library, 1080 in Google Scholar and 0 in other sources. Zero articles met the inclusion criteria.

MRI/CT/ULTRASOUND/BONE SCAN IMAGING

Recommendation: MRI, CT, Ultrasound, or Bone Scanning for Diagnosing Phalangeal or Metacarpal Fractures

MRI, CT, ultrasound, or bone scanning is not recommended for diagnosing phalangeal or metacarpal fractures.

Strength of Evidence – Not Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
There are no quality studies evaluating MRIs, CTs, ultrasound, or bone scanning for diagnosing phalangeal or metacarpal fractures. As fracture displacement and rotation are of primary concern, most fractures are readily diagnosed and treatment planned with radiographs. Discovering occult non-displaced fractures on CT would be unlikely to change the management except for delineation of articular impaction injuries. (78) Therefore, MRI, CT, diagnostic ultrasound, or bone scanning is not recommended for diagnosing phalangeal or metacarpal fractures.

Evidence for the Use of MRI/CT/Ultrasound/Bone Scan Imaging
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: MRI, CT, Ultrasound, bone, scan, imaging; Metacarpal, Middle, Proximal, Phalangeal, boxer's, Fracture, Bone, Diagnostic, Diagnosis, Sensitivity, Specificity, positive, predictive, value, negative, predictive, Predictive, Value, of, Tests, efficacy, efficiency. We found and reviewed 90 articles in PubMed, 1 in Scopus, 5 in CINAHL, 647 in Google Scholar, and 1 in Cochrane Library. We considered for inclusion 2 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 744 articles considered for inclusion, 0 randomized trials and 0 systematic studies met the inclusion criteria.

Initial Care

Initial management should include treatment of soft tissue injuries (1224) and pain control following completion of physical examination. Regional anesthesia should be administered to complete diagnostic assessment (passive range of motion, rotational alignment) and to perform closed reduction of the fracture, although not until neurovascular examination is documented.

Regional anesthesia is typically performed through injection of local anesthetic as a digital block through one of many described techniques including digital ring block, palmar subcutaneous block, metacarpal block, and volar thecal block. The traditional digital block technique, also known as dorsal subcutaneous
block, and occasionally referred to as metacarpal block, includes instilling local anesthetic from a dorsal approach into the webspace lateral to each side of the injured finger. A true metacarpal block is similar to ring block, but at the metacarpal head. A volar thecal block, also referred to as transthecal block, is the instillation of local anesthetic into the potential space of the tendon sheath at the distal palmar crease (A-1 pulley) proximal to the injured digit. The palmar subcutaneous block is performed at the same location as the thecal block, but subcutaneously. Other block techniques include ulnar or radial block injuries that are proximal to the phalanx, such as for metacarpal injuries, and hematoma block which is the direct injection of local anesthetic into the fracture hematoma.

DIGITAL BLOCK
Recommendation: Digital Block – Traditional (Ring) Block Technique, Palmar Subcutaneous Block

The ring block technique, followed by volar subcutaneous block, is moderately recommended for digital anesthesia, as it provides more effective coverage of dorsal phalangeal injuries than the other techniques.

Strength of Evidence – Moderately Recommended, Evidence (B)
Level of Confidence – Moderate

Rationale for Recommendation
For phalangeal fractures, there is clear evidence that the three most common digital blocks are similarly effective in onset and depth of anesthesia, although each has advantages and drawbacks particular to the specific technique. However, although it requires two punctures, the traditional digit or ring block has been found to be as effective or more effective than the other two block types as it provides better anesthetic results for the dorsal finger as compared to palmar (subcutaneous) block(1225-1227) and transthecal block.(1228-1230) There is no clear difference in the primary anesthesia outcomes between transthecal and palmar techniques(99, 1231, 1232) although patients preferred the subcutaneous technique and many reported residual pain at the block site 24 hours after injection in the transthecal block group. Subjects in the ring block were also satisfied with the technique compared to transthecal blocks, and were rated very similar to palmar block despite having two injections. Thus, the subcutaneous techniques of ring block palmar subcutaneous block are recommended over transthecal block mainly related to patient preference and residual pain, and ring block is recommended as the first line technique as it is less likely to have incomplete anesthesia of the dorsal finger. There are no quality studies for hematoma block in the hand, but they have been reported effective in distal radius, ulnar, and ankle injuries. Hematoma block may provide advantage for proximal metacarpal injuries over ulnar/radial blocks.

Evidence for the Use of Digital Block for Middle and Proximal Phalangeal or Metacarpal Fractures
There are 2 high- (99, 1227) and 7 moderate-quality(1225, 1226, 1228-1232) RCTs or crossover trials incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Digital block, digital anesthesia, ring block technique, palmar subcutaneous block, middle, proximal, phalangeal, metacarpal, fractures, bone fractures, boxers; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 41 in Scopus, 1 in CINAHL, 0 in Cochrane Library, 60 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Nine articles met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yin 2006</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>8.5</td>
<td>N = 91 (23 female/68 male) with injuries to 1-2 fingers distal to basal crease of finger. Age 14-60.</td>
<td>Traditional digital block (n = 50) vs single subcutaneous palmar block (n = 41). Follow-up for 1 month.</td>
<td>No differences between 2 groups per time to onset of anesthesia and injection pain score with per protocol or ITT analyses.</td>
<td>&quot;The palmar techniques, including single subcutaneous palmar block and transthecal block carry a risk of not anesthetizing the dorsum of the digit adequately, particularly the dorsum of the thumb and the proximal phalanx of the fingers.&quot;</td>
<td>Study included RCT as well as meta-analysis of other digital anesthesia RCTs.</td>
</tr>
<tr>
<td>Hung 2005</td>
<td>RCT</td>
<td>Sponsored by funds from American Foundation of Surgery of Hand, Raymond M. Curtis Research Foundation and MedStar Research Institute. No mention of COI.</td>
<td>8.0</td>
<td>N = 50 (gender not specified) healthy volunteers. Age not given.</td>
<td>Digital (metacarpal) block vs. single subcutaneous palmar block vs. transthecal block</td>
<td>Overall significant difference (p &lt;0.001) between methods evaluated with digital metacarpal block taking significantly longer to abolish sensation (265 seconds vs. 187 seconds vs. 176 seconds) as compared with other 2 methods. No significant difference between average pain scores by patients; 43% chose subcutaneous block as their first choice vs. metacarpal block vs. transthecal block.</td>
<td>&quot;Subcutaneous block is effective and preferred by healthy volunteers for digital anesthesia.&quot;</td>
<td>Study conducted in non-injured hands. Volume of anesthetic was limited to 2ml. All subject received all blocks in different fingers. Results are opposite those found by Knoop.</td>
</tr>
<tr>
<td>Hill 1995</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>7.0</td>
<td>N = 81 (gender not specified) healthy adults. Age 18-45 years.</td>
<td>TT or transthecal block vs TD or traditional digital block or ring block.</td>
<td>Blocks completed with 2ml 1 % lidocaine at each site. All blocks successful without complications. Mean VAS pain scores favored traditional block (1.4 ± .13 vs. 1.7 ± .17, p = 0.02). Time to loss of pinprick sensation was faster for ring block (188 vs. 152 seconds).</td>
<td>Transthecal digital block is clinically equal to the traditional method in terms of time to anesthesia and associated pain.</td>
<td>Study included 162 blocks on 81 subjects. Patients were healthy without injury and served as their own control.</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Mean Age</td>
<td>Gender</td>
<td>Technique</td>
<td>Pain Scores</td>
<td>Comparison</td>
<td>Results</td>
<td>Methodology</td>
</tr>
<tr>
<td>-------</td>
<td>---</td>
<td>----------</td>
<td>--------</td>
<td>-----------</td>
<td>------------</td>
<td>-----------</td>
<td>---------</td>
<td>-------------</td>
</tr>
<tr>
<td>Williams 2006</td>
<td>7.0</td>
<td>N = 27 (16 female/11 male) volunteers. Mean age 31 years.</td>
<td>Digital block vs. single subcutaneous palmar block.</td>
<td>No difference in median pain scores with respect to volar and dorsal injection techniques (VAS 4.06 vs. 4.52). Volunteers preferred palmar block (22 of 27) if required to have another in the future.</td>
<td>&quot;Our results demonstrated that there was more pain experienced with the use of the two-injection dorsal technique, but the difference in pain scores was not statistically significant.&quot;</td>
<td>Lack of blinding; study conducted on healthy volunteer population. Both techniques had incomplete anesthesia in some subjects (palmar – dorsum of phalanges, digit – hemidigit anesthesia).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cummings 2004</td>
<td>6.5</td>
<td>N = 25 Paid volunteers. Mean age of 31 years old. 13 Females, 12 Males</td>
<td>Transthecal (modified) (N=25) vs. Traditional digital block (N = 25)</td>
<td>No difference in pain rating from the block procedures (p = 0.579). Average time to complete block was faster in all measured dermal zones (average of 1.38 to 5.46 minutes faster) for traditional block vs. transthecal block (p &lt;0.05).</td>
<td>&quot;The effect of modified transthecal block is equal to that of traditional block in terms of pain perception. For the dorsal and radial proximal zones, the traditional block appears to have better distribution of anesthesia.&quot;</td>
<td>Subjects served as both comparison groups. Author states study was double-blind, but appears questionable as number and location of puncture was different for each method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cummings 2004</td>
<td>6.5</td>
<td>N = 142 Mean age of 33.5 years old. 14 Females, 128 Males</td>
<td>Transthecal (N = 71) vs. Single injection subcutaneous (superficial to A-1 pulley) digital block (N=71)</td>
<td>Blocks performed with 3cc 1% lignocaine/ bupivacaine mixture. No differences between 2 techniques with regards to effectiveness, distribution, onset, and duration of anesthesia.</td>
<td>&quot;The subcutaneous block would appear to be a better choice as it is easier to administer and has no risk of intraarticular injection.&quot;</td>
<td>Study compared single injection techniques in subjects with actual injuries. Randomization and allocation is unclear.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Knoop 1994</td>
<td>5.5</td>
<td>N = 30 patients that required digital anesthesia.</td>
<td>Digital block (N = 30) vs. Single subcutaneous palmar block</td>
<td>Digital block not statistically less painful than metacarpal block (VAS 2.53±1.98cm vs. 3.35±2.77cm, p = 0.18). Digital block more efficacious as metacarpal block failed anesthesia to pinprick in 23% vs.</td>
<td>&quot;Digital block and metacarpal block, as described in this study, are equally painful procedures. Digital block, however, is more efficacious and requires significantly less</td>
<td>Subjects served as both comparison groups with both procedures being completed on half of same finger, which is major weakness. Lack of methodology details.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Sample Size</td>
<td>Study Design</td>
<td>Interventions</td>
<td>Results</td>
<td>Comments</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-------------</td>
<td>--------------------</td>
<td>--------------------------------------------</td>
<td>-------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Keramidas 2004</td>
<td>N = 50</td>
<td>Crossover Trial</td>
<td>Transthecal Digital Block (N = 50)</td>
<td>Time to anesthesia shorter for digit 2.82 minutes ± 1.01 vs. 6.35 minutes ± 2.94 (p &lt;0.001).</td>
<td>The transthecal digital block is comparable to the traditional subcutaneous infiltration technique with respect to the time and effectiveness of anesthesia but not with respect to the associated pain following anesthesia. Patients seem to prefer the subcutaneous infiltration technique because it is less painful.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Traditional digital block (N = 50)</td>
<td></td>
<td>Randomization and allocation unclear, although patients served as both intervention arms. States study double blinded but only described blinding of assessor.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low &amp; Vartany 1997</td>
<td>N = 20</td>
<td>Crossover Trial</td>
<td>Transthecal (N = 20) vs. Single injection subcutaneous (superficial to A-1 pulley) digital block (N = 20)</td>
<td>Blocks performed with 2ml 1% lidocaine; 40% of transthecal group and 45% of subcutaneous group achieved entire finger anesthesia. No differences based on injection method. No differences in magnitude of sensory nerve action potentials. Injector subjectively rated subcutaneous injections as easier to perform than transthecal.</td>
<td>Transthecal and subcutaneous techniques showed no differences in terms of distribution, onset, and duration of anesthesia. Although both techniques give similar levels of anesthesia, subcutaneous block is believed to be superior because the transthecal technique has more disadvantages.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Follow up 24 hrs after experiment.</td>
<td></td>
<td>Lack of study details, including randomization and allocation methods. Subjects were own control, and had no injuries.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Medications

NSAIDS/ACETAMINOPHEN
Recommendation: NSAIDs or Acetaminophen for Phalangeal or Metacarpal Fractures
NSAIDs or acetaminophen are recommended to control pain from phalangeal or metacarpal fractures.

Indications – Pain due to phalangeal or metacarpal fracture.
Frequency/Duration – Scheduled dosage rather than as needed is generally preferable.
Indications for Discontinuation – Resolution of pain, lack of efficacy, development of adverse effects particularly gastrointestinal.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
There is no quality evidence, however these medications are thought to be effective for control of swelling and pain in the initial stages of injury, are not invasive, have low adverse effects, are low cost, and thus are recommended. While there have been some concerns regarding delayed fracture healing, other studies have suggested no delayed bone healing (see Distal Forearm Fractures section). These concerns appear outweighed by pain management concerns.

Evidence for the Use of NSAIDs or Acetaminophen
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAID, aspirin, acetaminophen, Middle, Proximal, Phalangeal, Metacarpal, Fractures, bone Fractures, boxer’s; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 56 in Scopus, 0 in CINAHL, 4 in Cochrane Library, 60 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

ANTIBIOTIC PROPHYLAXIS
Recommendation: Antibiotic Prophylaxis for Open Phalangeal Fractures

There is no recommendation for or against the use of antibiotic prophylaxis for open phalangeal fractures.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
Antibiotic prophylaxis for open phalangeal fractures are commonly used but may not be necessary based on the results of a prospective (non-randomized) trial of 91 open phalangeal fractures in fingers with intact digital arteries which compared aggressive irrigation and debridement with antibiotics. There were equal numbers of soft tissue infections and no cases of osteomyelitis in either group.(1220) However, the study may have been underpowered for these infrequent complications.

Evidence for the Use of Antibiotic Prophylaxis
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, and Cochrane Library without date limits using the following terms: Anti-bacterial agents, antibiotics, antibiotic prophylaxis, and antibiotic; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly;
systematic, systematic review, retrospective, and prospective studies. We found and reviewed zero articles in PubMed, 1 in Scopus, zero in CINAHL, and 1 in Cochrane Library. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Zero articles met the inclusion criteria.

**TETANUS IMMUNIZATION**

**Recommendation: Tetanus Immunization Status for Open Fractures**

For open fractures, it is recommended that tetanus immunization status be updated as necessary.

**Indication** – Wounds that are not clean or burns if more than 5 years have elapsed since last tetanus immunization.(1221)

*Strength of Evidence* – Recommended, Insufficient Evidence (I)

*Level of Confidence* – High

**Rationale for Recommendation**

There are no quality studies of tetanus immunization updating for these fractures. However, these immunizations are widely used and believed to have been successful on a population basis in reducing risk of tetanus over many decades. Tetanus immunizations are minimally invasive, have low adverse effects and are low cost. As the adverse effects of not immunizing may be fatal, tetanus immunization updating for open wounds is recommended. Wounds that are not clean or burns should require immunization if over 5 years since last immunization, rather than 10 years.(1221) Patients without a completed immunization series of 3 injections should receive tetanus immune globulin along with immunization.

**Evidence for the Use of Antibiotic Prophylaxis**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, and Cochrane Library without date limits using the following terms: Anti-bacterial agents, antibiotics, antibiotic prophylaxis, and antibiotic; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed zero articles in PubMed, 1 in Scopus, zero in CINAHL, and 1 in Cochrane Library. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library and 0 from other sources. Zero articles met the inclusion criteria.

**Physical Methods/Rehabilitation**

**EXERCISE**

Exercise is not indicated acutely. Some patients have considerable, functional deficits after casting and require exercise. For patients with residual deficits, particularly after casting or post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

**Middle and Proximal Phalanx Fractures**

**IMMOBILIZATION AND SURGERY**

There are no quality studies comparing non-operative treatment, percutaneous fixation, bone screws, or plates for middle and proximal phalangeal fractures. There also are no quality studies defining acceptable limits of displacement for non-operative management, determining the ideal splint time or duration of internal or external fixation, making comparisons of fixation techniques or defining ideal post-operative rehabilitation impractical. Immobilization or fixation technique is therefore dictated by the physical and radiographic findings. More than 90% of phalangeal fractures can be managed non-operatively.(83, 1233) Non-operative management techniques include padded aluminum splints, buddy tape, functional splinting, and gutter casting.
1. **Recommendation: Immobilization for Middle and Proximal Phalanx Fractures**
   
   Immobilization is recommended for treatment of middle and proximal phalanx fractures.(83, 1233)
   
   *Frequency/Duration* – When percutaneous fixation with wire is used, supplemental stabilization with splint or casting for 3 to 4 weeks should also be used as the wire does not provide sufficient rigidity.
   
   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – **Moderate**

**Rationale for Recommendation**

For middle and proximal phalangeal fractures that do not fit the criteria addressed in the specific fracture types, splinting for 3 to 4 weeks is recommended.

2. **Recommendation: Immobilization for Non-displaced and Stable Transverse Diaphyseal Fractures of the Middle and Proximal Phalanges**

   Non-operative management (immobilization) of non-displaced and stable transverse diaphyseal fractures of the middle and proximal phalanges is recommended as these fractures do not require fixation and can be managed without surgery.

   *Frequency/Duration* – Immobilization of the affected digit with neighboring digit in 70 to 90° of MCP flexion for 1 to 3 weeks.

   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – **Moderate**

**Rationale for Recommendation**

There are no quality studies that address non-operative management of acute non-displaced and stable transverse diaphyseal fractures of the middle and proximal phalanges. These fractures have good results with non-operative management. The affected digit is immobilized with neighboring digit in 70 to 90° of MCP flexion for 1 to 3 weeks. The tolerance limits for non-operative management after closed reduction are angulation of 10°, shortening less than 2mm, bone apposition of greater than 50%, and no malrotation. Displacement outside these limits should be evaluated for treatment with closed reduction and percutaneous fixation, or upon failure of closed reduction, open reduction and internal fixation.(68, 87)

3. **Recommendation: Non-operative Management of Non-displaced Oblique Fractures of the Middle and Proximal Phalanges**

   Non-operative management of non-displaced oblique fractures of the middle and proximal phalanges is recommended as these fractures are usually stable and require rigid immobilization alone.

   *Frequency/Duration* – Examinations weekly for the first 3 weeks.

   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – **Moderate**

**Rationale for Recommendation**

There are no quality studies for management of oblique fractures. Buddy taping should not be used as rotational correction may not occur. The fracture must be examined weekly for the first 3 weeks. Displaced fractures can be stabilized with closed reduction and percutaneous Kirschner wires or through open reduction with interfragmentary screw or plate-and-screw devices. Long oblique fractures (length double the diameter of bone at fracture site) can be stabilized by closed reduction and percutaneous Kirschner wires(78, 87) or with intramedullary wires.(89)

4. **Recommendation: Closed Reduction with Splinting for Base Phalanx Fractures**

   Closed reduction with splinting is recommended for base phalanx fractures.(1224)

   *Indications* – Involvement of less than 40% of the middle phalanx base.

   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – **Low**
Rationale for Recommendation
There are no quality studies for management of base fractures. However, base fractures are commonly a fracture-dislocation of the PIP joint and consists of an avulsion fracture of the volar lip of the middle phalanx and dorsal subluxation of the remaining base of the middle phalanx. Closed reduction with splinting is recommended if the fracture involves less than 40% of the middle phalanx base. If this fails, treatment is by pin fixation. Dynamic traction is another effective described technique for base fractures and also for the treatment of comminuted intra-articular fractures (Pilon fractures) of the base of the middle phalanx. Unstable displaced articular fractures at the base of the proximal phalanx are treated with percutaneous wires crossing the MCP joint to hold the joint reduced, and a transverse wire holding the fracture alignment similar to Bennett’s fracture of the thumb.(1224)

5. Recommendation: Surgical Management of Condylar Fractures
Surgical management of condylar fractures is recommended as these fractures are unstable.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
Displaced oblique fractures involving a single condyle are unstable, and are stabilized operatively with two transverse pins or screws. Bicondylar fractures are reconstructed with screws and connected to the shaft with a pin or through the use of a condylar plate.(78, 1224)

6. Recommendation: Surgical Management for Malrotated Phalangeal Fractures
Surgical management for malrotated phalangeal fractures is recommended as deformity and impairment may result.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
Surgical management for malrotated phalangeal and metacarpal fractures is recommended, to prevent or reduce rotational deformity that can result in fingers crossing over each other or interfering with hand function, if malrotation cannot be corrected and stabilized by closed reduction.

Metacarpal Fractures
Except for 5th metacarpal neck fractures, there are no quality studies comparing non-operative management, percutaneous fixation, bone screws, or plates. Further, there are no quality studies defining acceptable limits of displacement for non-operative management, determining the ideal splint time or duration of internal or external fixation, making comparisons of fixation techniques or defining ideal post-operative rehabilitation. Metacarpal head fracture in an uncommon fracture, usually intra-articular and frequently results in late traumatic arthrosis.

1. Recommendation: Non-operative Treatment of Distal Metacarpal Head Fracture
Non-operative treatment of distal metacarpal head fractures using closed reduction and protective immobilization with radial or ulnar gutter splint is recommended for fractures with less than 20% of joint involvement.

Indication – Fractures with less than 20% of joint involvement.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation
There are no quality studies that address non-operative treatment of acute distal metacarpal head fractures. Metacarpal neck fractures are common extra-articular fractures at the base of the head, usually the result of axial impaction, resulting in the neck being displaced dorsally and the metacarpal head being displaced volarly. Recommendations are based on prior clinical experience. Cases with greater than 20% joint involvement likely require open reduction and internal fixation followed by nearly immediate motion.(84)
2. **Recommendation: Non-operative Treatment of Distal Metacarpal Head Fracture with Acceptable Angulation**

   **Non-operative treatment of distal metacarpal head fracture using angulation is recommended.**

   **Indication** – Degree of angulation 15° in the ring finger and 10° in the index and long fingers.
   **Frequency/Duration** – These fractures heal quickly in 3 to 4 weeks with a gutter or radial splint maintaining MCP joint flexion. Operative fixation is usually with percutaneous pinning.(84)

   **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
   **Level of Confidence** – Moderate

**Rationale for Recommendation**

Treatment of Boxer’s fracture, or 5th metacarpal neck fracture, varies widely, with proponents of casting, splinting, taping, and operative fixation. There are no quality studies comparing non-operative and operative techniques, although there are two prospective trials with long-term follow-up suggesting non-operative treatment with early mobilization provides comparable outcomes to operative intervention, and perhaps is superior as operative fixation may increase the risk for metacarpohamate joint osteoarthrosis,(1234) although it may result in slightly more cosmetic deformity.(1235)

3. **Recommendation: Non-operative Treatment of Fifth Metacarpal Neck Fractures (Boxer’s Fracture)**

   **Non-operative treatment is recommended before surgical treatment for most 5th metacarpal neck fractures as the outcomes are similar both functionally and anatomically.**

   **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
   **Level of Confidence** – Moderate

4. **Recommendation: Use of Functional Therapies Rather than Casting or Splinting for Fifth Metacarpal Neck Fractures**

   **The use of functional therapies including taping, functional bracing, and strapping is moderately recommended over casting or ulnar splinting for 5th metacarpal neck fractures.**

   **Strength of Evidence** – **Moderately Recommended, Evidence (B)**
   **Level of Confidence** – Moderate

**Rationale for Recommendations**

There are eleven moderate-quality studies available comparing the effectiveness of different non-operative measures and no clear evidence of superiority of one approach over another.(1236-1246) A Cochrane review also concluded that no single non-operative treatment regimen for fracture of the neck of the 5th metacarpal can be recommended as superior to another in results.(1247) However, there is moderate evidence supporting functional therapies in general, with two moderate-quality studies supporting functional therapies over casting or splinting. Functional taping provided better functional outcome with no increase in deformity over casting,(1236) and treatment by compression bandage without reduction or splinting with a mean angulation angle of 48° had equal functional outcomes with closed reduction and splinting.(1238) Another moderate-quality study supports the use of strategic metacarpal bracing (3-point brace), another type of functional therapy, which provided equivalent functional outcomes to neighbor strapping but with less pain.(1237)

Ulnar gutter cast was compared with functional mobilization (pressure bandage for 1 week) in patients with 70° angulation or less and no rotation of the 5th metacarpal. Although the study was limited by small sample size, there were no differences in subjective symptoms of pain, return to work and hobby, or the need for physiotherapy.(1239) Several non-randomized prospective and retrospective trials with long-term follow up (up to 4 years) of patients treated without immobilization support these findings.(1248-1250) Other methods described in the literature for non-operative management with reported efficacy include fracture brace,(1251) modified Thomine brace,(1252) and a glove cast.(1253) However, there is no recommendation for or against any of these interventions as there is insufficient evidence.

There is no consensus on the degree of acceptable volar angulation manageable with non-operative treatment. It is reported as 30° in a small prospective case series of 18 patients(1254) followed for a mean of 20 months, and 60° and 70° in early mobilization trials.(1238, 1239) Intra-articular fractures
have also been reported to be successfully treated non-operatively, although comparison of non-operatively and operative management demonstrated high level of intermittent pain (38%), 49% decreased grip, and 65 radiographic signs of osteoarthritis in both groups. 

Evidence for the Use of Functional Therapies vs. Casting or Splinting for Metacarpal Fractures
There are 11 moderate-quality RCTs incorporated into this analysis. There are 3 low-quality RCTs in Appendix 2.

Taping:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Taping, functional bracing, strapping vs. casting or splinting (fifth metacarpal neck fractures only), Middle and Proximal Phalangeal and Metacarpal Fractures (fifth metacarpal neck fractures, boxer's fracture, shaft metacarpal fractures - transverse, oblique, spiral, comminuted); controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 17 articles in PubMed, 4 in Scopus, zero in CINAHL, zero in Cochrane Library, 27 in Google Scholar, and zero from other sources. We considered for inclusion 11 from PubMed, zero from Scopus, zero from CINAHL, zero from Cochrane Library, zero from Google Scholar, and zero from other sources. Of the 11 articles considered for inclusion, 11 randomized trials and zero systematic studies met the inclusion criteria.

Fixation:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: percutaneous fixation, bone screws, plates, internal fixation, external fixation, closed reduction, middle, proximal, phalangeal, metacarpal, fractures, bone fractures, boxer's, condylar fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 244 articles in PubMed, 301 in Scopus, 11 in CINAHL, 1 in Cochrane Library, 282 in Google Scholar, and 0 from other sources. We considered for inclusion 3 from PubMed, 2 from Scopus, 0 from CINAHL, Cochrane Library, and Google Scholar, and 1 from other sources. Of the 6 articles considered for inclusion, 5 randomized trials and 1 systematic studies met the inclusion criteria.

Immobilization:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms Immobilization: padded aluminum splints, buddy tape, functional splinting, gutter casting, splinting (closed reduction), Middle, Proximal, Phalangeal, Metacarpal, Fractures, bone Fractures, boxer's; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 2 in Scopus, 4 in CINAHL, 19 in Cochrane Library, 100 in Google Scholar, and 1 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 1 from Cochrane Library, 0 from Google Scholar, and 1 from other sources. Of the 2 articles considered for inclusion, 0 randomized trials and 2 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Study Authors</th>
<th>Year</th>
<th>Score</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hofmeister</td>
<td>2008</td>
<td>6.0</td>
<td>N=81 with an acute (&lt;7 days old) isolated fracture of the 5th metacarpal neck. Mean age 25 years. No mention of gender.</td>
<td>Casting with the MCP joint in flexion in a short-arm cast with volar outriggers with ring and small finger interphalangeal joints in extension, SAC-VOR (N=40) vs. casting with MCP joint in neutral extension and a cast with a 3-point mold about the fracture site, MCP-ext (N=41). All patients underwent a fracture reduction prior to cast placement. Cast was removed after 4 weeks. Assessments at 1 week, 4 weeks, and 3 months after the start of treatment.</td>
<td>Postreduction AP plane: SAC-VOR 5° vs. MCP-ext 14° (p&lt;0.05).</td>
<td>&quot;[W]e found that both methods of immobilization were equally effective in maintaining fracture reduction.&quot;</td>
<td>Data suggest comparable efficacy between (SAC VOR) and (MCP-ext) with a slight advantage to MCP-ext in terms of grip strength, patient tolerability and ROM.</td>
</tr>
<tr>
<td>Harding</td>
<td>2001</td>
<td>5.5</td>
<td>N = 73 (3 females, 62 males) Patients with minimally angulated (&lt;40°), closed fractures of the little finger metacarpal neck with no rotational deformity or associated injury. Mean age was 26.5 years</td>
<td>Molded metacarpal brace (N=37) vs. neighbor strapping for 5th metacarpal neck fracture (N=28) Follow up at 3 weeks.</td>
<td>Patients treated with brace complained of less pain (p = 0.001) and had slightly better range of finger movement (p = 0.03). More returned to work by 3 weeks (p = 0.007). None developed rotational or significant angular deformity.</td>
<td>&quot;The results of our study... showed a clear benefit over neighbor strapping for mean range of active range of motion of MCP joint, mean pain score, and return to work by 3 weeks.&quot;</td>
<td>There was no mention of control for co-interventions. For working populations this study suggests earlier return to work.</td>
</tr>
<tr>
<td>Kuokkanen</td>
<td>1999</td>
<td>5.5</td>
<td>N = 29 (26 males, 3 females) Patients treated for subcapital fractures of the fifth metacarpal bone. Mean age: 29 years.</td>
<td>Compression bandage for 1 week vs. splint immobilization (MCP 60° of flexion)</td>
<td>Angulation of fracture remained practically at same level compared with primary angulation in both groups. ROM of MCP (p = 0.02) and PIP (p = 0.01) joints higher in functional group at 4 weeks, but no difference at 3 months. Grip force was better in functional group at 4 weeks (p = 0.002).</td>
<td>&quot;We suggest that at least subcapital fractures of the fifth MC that are modestly and slightly angulated should be treated functionally, without reduction and splinting. Based on the present findings the correction achieved by closed reduction does not persist...&quot;</td>
<td>Small sample size. Patients in functional group had higher degree of pre-treatment angulation but still had equal or better functional outcomes in this population.</td>
</tr>
<tr>
<td>Braakman</td>
<td>1998</td>
<td>5.0</td>
<td>N = 50 (43 males, 5 females)patients</td>
<td>Ulnar gutter plaster cast vs. functional tape of 5th metacarpal fracture.</td>
<td>In both groups, fracture reduction partially lost at 1</td>
<td>The patients in the tape group showed a quicker and superior</td>
<td>Lack of randomization and allocation details.</td>
</tr>
<tr>
<td>Study</td>
<td>No mention of sponsorship or COI.</td>
<td>N= 40 (38 males, 2 females) with a fracture of the subcapital MC-V ≤ 3 days old and angulated ≤70º. Mean age 29 years.</td>
<td>Ulnar gutter plaster cast for 3 weeks followed by mobilization within pain limits (N=20) vs. 1 week of pressure bandage (N=20). Follow-up 6 and 12 weeks after fracture.</td>
<td>There were no significant differences between groups at 6 and 12 weeks follow-up.</td>
<td>functional recovery than those in the cast group. After 6 months, there were no significant differences between the groups with regard to functional and anatomical results or the number of patients with residual symptoms.</td>
<td>No blinding of assessor.</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>----------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Statius Muller 2003 RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>5.0</td>
<td>Compression glove worn on injured hand and early mobilization (N=21) vs. immobilization in plaster splint (N=21). Treatment lasted between 6-14 days after entry. All patients received hand exercises between 6-13 days after injury. Follow-up began at week 2, weekly intervals for 3 weeks.</td>
<td>Mean loss of total active motion (degrees): week 2 glove 56±26 vs. splint 84±33 (p=0.0036); week 3 glove 23±17 vs. splint 46±23 (p=0.0010); week 4 NS (p=0.15). Mean increase in circumference of PIP joint (mm): week 2 glove 2.2±2.8 vs. splint 4.5±3.2 (p=0.019); week 3 glove 0.5±2.5 vs. splint 2.1±2.8 (p=0.059); week 4 NS (p=0.27).</td>
<td>Mean loss of total active motion (degrees): week 2 glove 56±26 vs. splint 84±33 (p=0.0036); week 3 glove 23±17 vs. splint 46±23 (p=0.0010); week 4 NS (p=0.15). Mean increase in circumference of PIP joint (mm): week 2 glove 2.2±2.8 vs. splint 4.5±3.2 (p=0.019); week 3 glove 0.5±2.5 vs. splint 2.1±2.8 (p=0.059); week 4 NS (p=0.27).</td>
<td>Small sample size. Data suggest comparable efficacy between groups.</td>
<td></td>
</tr>
<tr>
<td>McMahon 1994 RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>4.5</td>
<td>N=42 with unilateral fresh closed stable fractures (displaced &lt;50% of width of shaft, angulated less than 40º and showed an angle of over 60º between plane of fracture and axis of shaft) of the shaft of single finger metacarpal Mean age plaster 27 years, compression glove 35 years. No mention of gender.</td>
<td>Compression glove worn on injured hand and early mobilization (N=21) vs. immobilization in plaster splint (N=21). Treatment lasted between 6-14 days after entry. All patients received hand exercises between 6-13 days after injury. Follow-up began at week 2, weekly intervals for 3 weeks.</td>
<td>Mean loss of total active motion (degrees): week 2 glove 56±26 vs. splint 84±33 (p=0.0036); week 3 glove 23±17 vs. splint 46±23 (p=0.0010); week 4 NS (p=0.15). Mean increase in circumference of PIP joint (mm): week 2 glove 2.2±2.8 vs. splint 4.5±3.2 (p=0.019); week 3 glove 0.5±2.5 vs. splint 2.1±2.8 (p=0.059); week 4 NS (p=0.27).</td>
<td>Mean loss of total active motion (degrees): week 2 glove 56±26 vs. splint 84±33 (p=0.0036); week 3 glove 23±17 vs. splint 46±23 (p=0.0010); week 4 NS (p=0.15). Mean increase in circumference of PIP joint (mm): week 2 glove 2.2±2.8 vs. splint 4.5±3.2 (p=0.019); week 3 glove 0.5±2.5 vs. splint 2.1±2.8 (p=0.059); week 4 NS (p=0.27).</td>
<td>Small sample size (N=42). Data suggest glove group experienced less pain and prevented loss of function and better range of motion during second and third weeks.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Sample Size</td>
<td>Design</td>
<td>Sponsorship/COI</td>
<td>Intervention</td>
<td>Primary Outcome Measures</td>
<td>Additional Notes</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>-------------</td>
<td>--------</td>
<td>----------------</td>
<td>-------------</td>
<td>--------------------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>Randall 1992</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>4.5</td>
<td>N=18 (13 males, 5 females) undergoing treatment of metacarpal fracture and hand has been immobilized for ≥2 weeks. Mean age 28.7 years.</td>
<td>Joint mobilization using traction and palmar/dorsal glide techniques (N=9) vs. control, no mobilization (N=9). Both groups received home exercises. Three appointments on alternate days over a 1 week period.</td>
<td>Increase in hand volume (cm³): week 2 glove 19±31 vs. splint 42±36 (p=0.029); week 3 NS (p=0.13); week 4 NS (p=0.69).</td>
<td>Small sample (18). Data suggest significant increase in metacarpal phalangeal joint motion after joint mobilization when compared to controls.</td>
</tr>
<tr>
<td>Konradsen 1990</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>4.0</td>
<td>N=100 with shaft or neck fracture of the 2nd-5th metacarpal bone. Median age plaster cast 22 years, functional cast 22.5 years. No mention of gender.</td>
<td>Immobilization by plaster cast as ulnar gutter cast for 5th metacarpal or as dorsal cast for 2nd-4th metacarpals (N=) vs. immobilization by functional cast, DeltaLite® allowing free ROM of wrist and digit joints (N=). Reduction was performed in all patients. Casts removed after 3 weeks. Assessments at 1 week, 3 weeks, and 3 months after injury.</td>
<td>Fracture angulation after cast removal (median degrees): subcapital – plaster cast 25 vs. functional cast 16 (p&lt;0.05); diaphyseal – plaster cast 14 vs. functional cast 5 (p&lt;0.01). Return to work in occupations where use of hands could be avoided (time in days): plaster cast 7 vs. functional cast 1 (p&lt;0.05).</td>
<td>Data suggest functional cast group returned to work in 1/3 the time compared to plantar cast group. Functional casting reduced volar angulation by 2/3 in metacarpal shaft fractures and 1/3 in metacarpal neck fractures as compared to plantar cast group.</td>
</tr>
<tr>
<td>Kim 2015</td>
<td>RCT</td>
<td>No mention of sponsorship. NO COI.</td>
<td>7.0</td>
<td>N = 46 (46 males, zero females) with displaced fifth metacarpal neck fractures with apex dorsal angulation &gt;30°. Mean age 29 years.</td>
<td>Antegrade intramedullary K-wire pinning (N =23) vs. percutaneous retrograde intramedullary K-wire pinning (N =23). All patients received an ulnar gutter short-arm splint post-surgery to be worn for 5 weeks.</td>
<td>Postoperative outcomes at 3 months: ROM antegrade 80 vs. retrograde 69 (p&lt;0.001); VAS points antegrade 2 vs. retrograde 4 (p&lt;0.001); grip strength % antegrade.</td>
<td>Data suggest antegrade intramedullary pinning had some clinical benefit to retrograde intramedullary pinning during recovery phase but these benefits are not present at 6 months.</td>
</tr>
</tbody>
</table>

**Fixation**

*Mean torque range of motion (TROM): treatment 73.6 vs. control 58.7 (no p-value reported).*

*The joint mobilization treatment given to the subjects in this study resulted in a significant gain in AROM and decrease in joint stiffness within a treatment session when compared to the control group.*

*Functionally treated patients returned to work faster than did patients in studies of nonimmobilization (Hunter and Cowen 1970, Arafa et al. 1986, Ford et al. 1989), perhaps because the short, but solid, bandage gave a feeling of security and provided pain relief.*

*Data suggest functional cast group returned to work in 1/3 the time compared to plantar cast group. Functional casting reduced volar angulation by 2/3 in metacarpal shaft fractures and 1/3 in metacarpal neck fractures as compared to plantar cast group.*
<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Description</th>
<th>Follow-up</th>
<th>Outcome Measures</th>
<th>Postoperative Findings</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Winter 2007 RCT</td>
<td>5.5</td>
<td>N = 36 male with fracture of neck of 5th metacarpal, recent and closed fracture. Mean age 31.4 years. Transverse pinning with 2 K-wires 1.5mm diameter (N = 18) vs intramedullary pinning with 3 K-wires 1mm diameter (N = 18). Both groups wore palmar splint for 1 week after procedure then physiotherapy 3x a week for 30 days. K-wires removed 6 weeks after surgery. Follow-up at days 8, 15, 30, 45, 60, and 90 after surgery.</td>
<td>81 vs. retrograde 71 (p&lt;0.001); DASH score, points, antegrade 4.3 vs. retrograde 10.3 (p&lt;0.001). Postoperative outcomes at 6 months: ROM (p=0.35); VAS (p=0.67); grip strength (p=0.41); DASH score (p=0.48).</td>
<td>postoperatively in terms of ROM, VAS, grip strength, and DASH score of the fifth metacarpophalangeal joint than percutaneous retrograde intramedullary pinning, but that the differences in clinical parameters are not sustained at 6 months postoperatively.</td>
<td>“This study suggests that intramedullary pinning is a particularly efficient procedure for treatment of the boxer’s fracture...” Small sample. Data suggest better functional outcomes with intramedullary pinning group unclear if patients were informed of surgical treatment.</td>
<td></td>
</tr>
<tr>
<td>Kruithaag 2009 RCT</td>
<td>5.0</td>
<td>N = 75 (6 male and 64 female) with unstable distal radius fractures (AO-type A3) suitable for non-bridging external fixation; &gt;10º of dorsal angulation and/or radial shortening of &gt;2mm vs. uninjured wrist. Mean age 62 years. H group: Hoffman compact II fixator (N = 37) vs D group: Dynawrist fixator (N = 38). All patients treated with closed reduction. Follow-up post-op, 6, 12, 24, and 52 weeks.</td>
<td>Median radial tilt (degrees): post-op Hoffman 8 volar vs. Dynawrist 2 volar (p = 0.002); at removal of fixators Hoffman 9 volar vs. Dynawrist 4 (p = 0.04). Mean loss of flexion (degrees): 6 weeks Hoffman 34 vs. Dynawrist 24 (p = 0.001).</td>
<td>“The Dynawrist bridging but dynamic fixator gives radiographic and functional outcome similar to that of the Hoffman II compact non-bridging fixator.” Data suggest comparable efficacy in both groups for radial tilt, inclination and radial length.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sletten 2015</td>
<td>5.5</td>
<td>N = 85 (61 male and 24 female) with little finger metacarpal neck fractures with ≥30° palmar angulation in the lateral view. At least 18 years.</td>
<td>Conservative treatment without reduction of the fracture (N = 43) vs Closed reduction and bouquet pinning (N = 42). Follow-up at 1 week, 6 weeks, 3 months, and 1 year.</td>
<td>Median operative time 30 minute. The palmar angulation was a median of 41° (range 30–58) in the conservative group at inclusion. In the operative group, palmar angulation was reduced from a median of 40° (range 30–59) – 17° (range – 9–31). At 1 year, The QuickDASH score was median 0 in both groups. No statistically significant or clinical relevant differences in QuickDASH scores at any time, but a worse QuickDASH Work score in the operative group at 6 weeks before pin removal, (25 versus 6 points, p = 0.07).</td>
<td>“There was a trend versus better satisfaction with hand appearance (p = 0.06), but longer sick leave (p &lt; 0.001) and more complications (p = 0.02) in the operative group.”</td>
<td>Data suggest comparable efficacy between conservative treatment vs. bouquet pinning of little finger metacarpal neck fractures for pain, finger ROM, grip strength, and quality of life. However, there was better patient satisfaction with hand appearance but longer sick leave in the surgical group.</td>
</tr>
</tbody>
</table>
5. **Recommendation: Routine X-rays in Follow-up of Fifth Metacarpal Neck Fractures**

Routine x-ray for follow-up of non-operative treatment of 5th metacarpal fractures is not recommended as it has little clinical impact on fracture management.

*Strength of Evidence – Not Recommended, Insufficient Evidence (I)*  
*Level of Confidence – Low*

**Rationale for Recommendation**
Routine radiographs in follow-up of non-operative treatment for 5th metacarpal neck fracture were not found to be of clinical utility(1259) except in only one case from two retrospective studies of 307 patients and 288 patients. Follow-up radiographs are indicated if physical examination suggests loss of reduction or instability within one week of the fracture.

**Shaft Metacarpal Fractures**
Shaft metacarpal fractures are usually transverse, oblique, spiral or comminuted. Determination of whether or not a fracture can be managed non-operatively is unclear.

1. **Recommendation: Non-operative Management of Metacarpal Shaft Fractures**

There is no recommendation for or against non-operative management of metacarpal shaft fractures.

*Strength of Evidence – No Recommendation, Insufficient Evidence (I)*  
*Level of Confidence – Low*

**Rational for Recommendation**
There are no quality studies and there are conflicting opinions regarding whether any angulation of the middle and index finger is acceptable,(84) versus whether up to 15° of dorsal angulation of the middle and index finger(89) can be tolerated. The ring finger is thought to tolerate 20°.(84) There is general agreement that rotational deformity is poorly tolerated. Thumb shaft fractures are rare, and those with less than 30° angulation can be managed with forearm-hand-based opponens splint for 3 to 4 weeks. Parameters of fifth digit fractures are discussed separately (see Boxer’s Fracture). Ultimately, decisions for non-operative versus surgical intervention balance acceptance of metacarpal shortening with risks accompanying surgical intervention.

Oblique fractures likely benefit from fixation (intra-medullary wires)(89) to prevent shortening. If adequate closed reduction is achieved and the fracture is stable, a 3-point brace (pressure points over the fracture apex and two counter-pressure points proximal and distal on the opposite side) can be used. Metacarpal shaft fractures that cannot be reduced, are unstable, or have multiple neighboring shaft fractures require fixation (pinning, wire, plate, lag screws).

2. **Recommendation: Surgical Management for Base Fractures of the Proximal Metacarpal**

Surgical management of base fractures of the proximal metacarpal is recommended as these fractures are rarely stable and require percutaneous pins or screws to maintain reduction.

*Indications – Extra-articular fractures with up to 15° of deformity in the 4th and 5th metacarpals, and only 5° in the 2nd and 3rd metacarpals can be managed with immobilization using a gutter splint holding the MCP in 70° flexion, wrist in neutral position, and allowing movement of the PIP and DIP joints.(84)*

*Strength of Evidence – Recommended, Insufficient Evidence (I)*  
*Level of Confidence – Moderate*

3. **Recommendation: Operative Fixation for Bennett’s Fracture and Rolando’s Fracture**

Operative fixation is recommended for Bennett’s and Rolando’s fractures as these fracture types are unstable.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*  
*Level of Confidence – Moderate*
4. **Recommendation: Surgical Management for Malrotated Phalangeal Fractures**
   Surgical management for malrotated phalangeal fractures is recommended as deformity and impairment may result.
   
   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – **Moderate**

**Rationale for Recommendations**
There are no quality studies for Bennett’s or Rolando’s fractures of the thumb. Bennett’s fracture of the thumb is a common metacarpal base fracture associated with dislocation, and requires operative fixation with one or two wires to maintain alignment of the shaft and joint surface, as does the base of the 5th metacarpal.(68) Rolando’s fracture is a comminuted intra-articular burst fracture at the thumb base requiring internal and external fixation to preserve metacarpal length and reapproximate articular fragments.(84)

**Evidence for the Use of Surgery**
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: operative fixation, middle, proximal phalangeal, metacarpal fractures, metacarpal, neck fractures, boxer's fracture, shaft metacarpal fractures, transverse, oblique, spiral, comminuted; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 69 articles in PubMed, 90 in Scopus, 0 in CINAHL, 18 in Cochrane Library, 175 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, Cochrane Library, Google Scholar, and from other sources. Zero articles were included.

**Follow-up Visits**
There are no quality studies on frequency or timing of return visits. X-rays for follow-up of all metacarpal fractures are reasonable; however, fractures at risk for displacement after reduction are particularly recommended to have repeat radiographic studies 7 to 10 days after injury to ensure no further displacement or malrotation has occurred – **Recommended, Insufficient Evidence (I), Level of Confidence - Moderate.** Motion and other hand exercises should be started at the earliest date the fracture becomes stable.

**Physical Methods/Rehabilitation**

1. **Recommendation: Ice, Compression, and Elevation for Acute Metacarpal Fractures**
   Ice, compression, and elevation are recommended for controlling edema related to acute metacarpal fractures.
   
   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – **Low**

2. **Recommendation: Early Mobilization for Acute Metacarpal Fractures**
   Early mobilization of acute metacarpal fracture (before 21 days) is recommended.
   
   Strength of Evidence – **Recommended, Insufficient Evidence (I)**
   
   Level of Confidence – **Low**

**Rationale for Recommendations**
There are no quality studies for physical methods of metacarpal fracture management. However, it is believed that controlling edema and early mobilization result in a more favorable outcome. Inflammation associated with traumatic injury, fracture hematoma and any resultant soft tissue damage from fracture displacement including adjacent tendons and ligaments results in edema and routine tissue repair processes. Immobilization in the presence of these inflammatory processes can result in adhesions to tendons, ligaments, capsules, or skin and subsequent stiffness and loss of function.(1260) Phalangeal fractures respond less favorably to immobilization than metacarpal fractures, with a predicted 84% return
of motion, compared to 96% return in metacarpal fractures. Immobilization continued longer than 4 weeks is reported to further reduce the return to motion to 66%. (1222)

There are no quality studies defining the efficacy or advantages of specific physical methods in reducing the effect of inflammation and immobilization. Control of edema after injury has been thought to be important in restoring function. (1261) Ice, compression, and elevation should be emphasized, with particular emphasis on hand elevation overnight. (1262)

Early mobilization to promote venous return via muscle contraction and thus reduce swelling and propensity towards complex regional pain syndrome (CRPS) is advocated for stable fractures. Early motion of simple metacarpal fractures initiated within 21 days of injury is likely to result in earlier recovery of motion and strength, and earlier return to work without adversely impacting fracture alignment. (1263) Tendon gliding range of motion exercises should be initiated as soon as possible based on the fracture immobilization method.

**Evidence for the Use of Joint Mobilization**

There are 3 moderate-quality RCTs incorporated into this analysis. (1308, 1309, 1327) (Kuokkanen 99; Statius Muller 03; Sletten 15)

**Ice:**
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Ice, Compression, Elevation, Metacarpal, Middle, Proximal, Phalangeal, boxer’s, Fractures, Bone; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 0 in Google Scholar and 0 in other sources. Zero articles met the inclusion criteria.

**Joint mobilization:**
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Joint mobilization, early ambulation, Middle and Proximal Phalangeal and Metacarpal Fractures (fifth metacarpal neck fractures, boxer’s fracture, shaft metacarpal fractures - transverse, oblique, spiral, comminuted ); controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 4 articles in PubMed, 56 in Scopus, 380 in CINAHL, 3 in Cochrane Library, and 3 in Google Scholar. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 1 from Cochrane Library and 1 from Google Scholar. Of the 4 articles considered for inclusion, 3 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kuokkanen 1999</td>
<td>No mention of sponsorship or COI.</td>
<td>5.5</td>
<td>N = 29 (26 males, 3 females) Patients treated for subcapital fractures of the fifth metacarpal bone. Mean age: 29 years.</td>
<td>Compression bandage for 1 week vs. splint immobilization (MCP 60° of flexion)</td>
<td>Angulation of fracture remained practically at same level compared with primary angulation in both groups. ROM of MCP (p = 0.02) and PIP (p = 0.01) joints higher in functional group at 4 weeks, but no difference at 3 months. Grip force was better in functional group at 4 weeks (p = 0.002).</td>
<td>“We suggest that at least subcapital fractures of the fifth MC that are modestly and slightly angulated should be treated functionally, without reduction and splinting. Based on the present findings the correction achieved by closed reduction does not persist...”</td>
<td>Small sample size. Patients in functional group had higher degree of pre-treatment angulation but still had equal or better functional outcomes in this population.</td>
</tr>
<tr>
<td>Statius Muller 2003</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>5.0</td>
<td>N= 40 (38 males, 2 females) with a fracture of the subcapital MC-V ≤ 3 days old and angulated ≤ 70°. Mean age 29 years.</td>
<td>Ulnar gutter plaster cast for 3 weeks followed by mobilization within pain limits (N=20) vs. 1 week of pressure bandage (N=20). Follow-up 6 and 12 weeks after fracture.</td>
<td>There were no significant differences between groups at 6 and 12 weeks follow-up.</td>
<td>“[A] pressure bandage for 1 week and immediate mobilization is a sufficient alternative treatment of a boxer’s fracture, if this is not angulated greater than 70° and not rotated.”</td>
</tr>
<tr>
<td>Sletten 2015</td>
<td>RCT</td>
<td>This work was supported by a grant from Sofies Mindes Ortopedi AS, Oslo, Norway. NO COI.</td>
<td>5.5</td>
<td>N = 85 patients with little finger metacarpal neck fractures with ≥ 30° palmar angulation in the lateral view. Mean age Conservative Group 29 (18–67) and Operative group 25 (18–68) Gender (M:F) Conservative group (39:4) Operative (39:3)</td>
<td>Conservative group, received an initial plaster-of-Paris applied for pain for one week, then buddy strapping was applied over the proximal phalanges of the little and ring fingers, and the patients started active exercises. Operative Group underwent closed reduction and internal fixation by antegrade, intramedullary bouquet pinning then The postoperative regime. was identical to the conservative regime N = 43</td>
<td>For conservative vs. operative: QuickDASH (0 vs. 0 (p=0.54)), VAS overall Satisfaction (97 vs 100 (p=0.17)), TAM (º) (261 vs 260 (p=0.68)), Grip strength (kg) (49 vs 49 (p=0.78)),</td>
<td>“After 1 year, there were no statistical differences between the groups in QuickDASH score, pain, satisfaction, finger range of motion, grip strength, or quality of life. There was a trend versus better satisfaction with hand appearance (p = 0.06), but longer sick leave (p &lt; 0.001) and more complications (p = 0.02) in the operative group.”</td>
</tr>
</tbody>
</table>
Distal Forearm Fractures
Special Studies and Diagnostic and Treatment Considerations

**X-RAYS**

Recommendation: X-ray for Suspected Distal Forearm Fractures

X-rays in the posterior-anterior and lateral views are recommended as a first-line study for suspected distal forearm fractures.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*
*Level of Confidence – High*

**Rationale for Recommendation**

There is no quality evidence for evaluation of x-ray studies for evaluation of suspected distal radial fractures. However, x-ray studies are standard of practice for suspected fracture. Therefore, as a first-line study, PA, lateral and if available oblique x-ray image views are recommended. Radiographic evaluation should provide the provider necessary information on location, configuration, displacement, subluxation, likelihood of stability, and concomitant potential of soft tissue injury. Contralateral wrist x-ray images should be considered as a comparison that may improve reliability of some radiographic measurements, particularly for a more accurate determination of stability and provide greater guidance on indication for treatment.

**Evidence for the Use of X-rays**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Distal Forearm Fracture, xray, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 22 articles in PubMed, 3 in Scopus, 24 in CINAHL, 0 Cochrane Library, and 11,100 from Google Scholar. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles were included.

**MRI**

Recommendation: MRI for Diagnosing Distal Forearm Fractures

MRI is recommended to diagnose suspected soft-tissue trauma after x-ray images confirm a complex displaced, unstable, or comminuted distal forearm fractures.

*Strength of Evidence – Recommended, Insufficient Evidence (I)*
*Level of Confidence – Low*

**Rationale for Recommendation**

There is no quality evidence that MRI is superior to radiographs for the initial detection of distal radial fractures and should not be generally used as a first line test. Upon confirmation of displaced, comminuted or unstable fracture, MRI may be an important diagnostic technique for the evaluation of suspected injuries of soft tissues related to distal radius fractures, such as to the flexor and extensor tendons or the median nerve. Other potential indications include identification of triangular fibrocartilage complex perforations, ruptures of carpal ligaments, and demonstration of contents of the carpal tunnel.(1264-1266)
**Evidence for the Use of MRI**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: magnetic resonance imaging, MRI, distal forearm fracture, distal forearm fractures, colles' fracture, colles fracture, colles fractures, dinner fork deformity, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 19 articles in PubMed, 117 in Scopus, 1 in CINAHL, 4 in Cochrane Library, and 640 from Google Scholar. Zero articles met the inclusion criteria.

**CT**

*Recommendation: CT for Diagnosis and Classification of Occult and Complex Distal Forearm Fractures*

CT is recommended for investigation of occult and complex distal forearm fractures to gain greater clarity of fracture displacement, articular involvement, and subluxation of the distal radioulnar joint.\(^{(1267)}\)

*Indication* – Negative x-rays with occult fracture strongly suspected.

*Strength of Evidence* – **Recommended, Insufficient Evidence (I)**

*Level of Confidence* – **Moderate**

**Rationale for Recommendation**

In contrast to MRI, CT should be considered when x-ray images are negative but on the basis of physical findings an occult fracture is strongly suspected. CT may also be useful for evaluation of complex comminuted fractures, providing superior depiction of distal radial articular surface involvement, fragment positioning, and diagnosis of subluxations of the distal radioulnar joint.\(^{(1267, 1268)}\) The value of CT has been demonstrated by Katz et al, who showed the use of CT scanning for evaluation of articular step off and gaping, comminution, and treatment influenced observers to change treatment plans developed from radiographs and resulted in increased interobserver reliability in the proposed management of these injuries.\(^{(1269)}\)

**Evidence for the Use of CT**

There are 3 quality studies incorporated into this analysis.\(^{(1339, 1342, 1343)}\) (Johnston 92; Harness 06; Avery 14)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: CT, CAT, computed tomography, distal, Forearm, radial, Radius fractures, bone Fractures, Colles' Fracture, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 302 articles in PubMed, 20 in Scopus, 3 in CINAHL, 16 Cochrane Library, and 20 from Google Scholar. We considered for inclusion 1 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 1 from other sources. Of the 3 articles considered for inclusion 3 diagnostic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score</th>
<th>Number</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>Type of CT</th>
<th>X-ray used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes assessed</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Copyright© 2016 Reed Group, Ltd.
Avery III 2014 Retrospective Study

| 5.5 | 17 sets of images | Wr ist | Dist al Ra dial Fr act ur e | GE Light Spe ed VCT | - | + | + | - | - | - | CT and traction radiographs had the about the same ability to identify fracture fragmen ts, except for the volar rim fragmen t. The volar rim was correctly identifie d 72% of the time on traction radiography compar ed to CT’s 60% (p<0.01) . CT correctly identifie d the radial column more often than traction radiographs (71.8% vs 65.8%, p=0.04) | “The information obtained from the traction radiographs compared with CT imaging showed little significant difference with regard to fracture fragment characteriz ation and led surgeons to consistent treatment recommend ations with both imaging modalities” | Small sample. Data suggest use of traction radiographs may be an alternativ e to CT imaging for diagnosi ng and assessin g distal radial fractures. |

No ment ion of how man y patie nts, mea n age, or gend er.
| Harness 2006 Retrospective | 7.5 | 30 | Wr ist Fr actur e | GE Adv antag e 3.1 W orkstat ion | + | - | + | + | - | - | + | - | Pertaining to a coronal fracture line, 3D CT imaging resulted in a sensitivity of 0.82, a specificity of 0.50, and accuracy of 0.77. 2D CT imaging resulted in values of 0.81, 0.56, and 0.77, respectively. When combined, the two images had slightly better results: a sensitivity of 0.87, a specificity of 0.56, and an accuracy of 0.82. | “Three-dimensional computed tomography improves both the reliability and the accuracy of radiographic characterization of articular fractures of the distal part of the radius and influences treatment decisions” | Dats suggest use of 3-D CT for analyzing complex intra-articular distal radius fractures. |
| Johnston 1992 | 5.5 | 22  | Wrist/Hand | Acute distal radial and/or carpal injury | GE 9800 scanner | + | - | + | - | - | - | - | Only 19 of the 22 patients had a radial distal fracture. 3 sets of plain film were interpreted as normal. However, a CT scan revealed that all three were fractures. CT scan enhanced the details of the fractures. In one case, a “innocent lip fracture” on plain film turned into an intra-articular compression of the scaphoid fossa on the CT. | “CT has an advantage over conventional tomography in lending itself to potential three-dimensional reconstruction.” | Data suggest CT visualizes more detail in evaluating acute distal radial fractures compared to plain radiographs. |
|-------------|-----|-----|------------|----------------------------------------|-----------------|---|---|---|---|---|---|---|---|---|---|---|
| Mean age = 31.5 | 13 men | 9 women |

**Follow-up Visits**

No quality evidence exists for specific follow-up care of distal radial injuries outside of identified recommendations listed in this section. Routine follow-up as with other fractures should be followed, with consideration of forearm girth changes with reduced swelling after the immediate injury period that may necessitate re-casting or immobilization device adjustments, and monitoring the potential for reduction failure with subsequent radiographic studies.
Medications

NSAIDS

Recommendation: NSAIDs for Acute Colles’ Fracture (Distal Forearm Fracture) Analgesia

The use of NSAIDs to control bone pain associated with Colles’ fracture is recommended as there does not appear to be any negative effect on bone fracture union or functional recovery.

Indications – Bone pain associated with Colles’ fracture.

Frequency/Duration – Scheduled dosage rather than as needed is generally preferable.

Indications for Discontinuation – Resolution of pain, lack of efficacy, development of adverse effects particularly gastrointestinal.

Strength of Evidence – Recommended, Evidence (C)

Level of Confidence – Moderate

Rationale for Recommendation

There are three moderate-quality studies that show NSAIDs are effective for pain relief of distal forearm fracture.(1270-1272) Flurbiprofen was more effective than placebo in conjunction with bier block manipulation and for post manipulation pain.(1272) Piroxicam was more effective than paracetamol,(1270) and diflunisal was equally effective as mefenamic acid. No changes in Gartland and Werley functional assessment scores(1272) or functional recovery in post menopausal women(1270) were found, indicating there is no significant benefit other than pain relief from the use of NSAIDS.

Evidence for the Use of NSAIDs for Distal Forearm Fractures

There are 4 moderate-quality RCTs or prospective studies incorporated into this analysis.(1344-1347) (Thomas 86)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: anti-inflammatory agents, non-steroidal, NSAIDS, non-steroidal anti-inflammatory, ibuprofen, acetaminophen, distal, forearm, radial, radius, fractures, bone fractures, Colles’ fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 13 articles in PubMed, 25 in Scopus, 0 in CINAHL, 18 in Cochrane Library, 5,993 in Google Scholar, and 3 from other sources. We considered for inclusion 4 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 4 articles considered for inclusion, 4 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Davis 1988</td>
<td>Prospective study</td>
<td>N = 100 (gender not specified) with Colles’ fractures. Average age for groups I and II; 55.7 and 64.1.</td>
<td>7.0</td>
<td>Group 1, 50mg flurbiprofen (f) (n = 53) vs. Group 2 or placebo (p) randomized after dividing into group 1 (displaced fracture requiring Bier’s block and manipulative reduction) or group 2, no reduction (n = 45).</td>
<td>Mean grip strength (mmHg) Group 1 f/p: Week 2: 59/53, Week 6: 92/93, 1 year: 192/189. Mean grip strength (mmHg) Group 2 f/p: Week 2: 88/82, Week 6: 112/149, 1 year: 195/207. One-year assessment results (percentages) Group 1 f/p: patients who needed physiotherapy 11(45)/7(35), patients with residual pain 10(40/9(45), patients with restricted activities 10(40)/7(35). 1-year assessment results</td>
<td>“[F]lurbiprofen provides significant pain relief and does not significantly delay union of Colles’ fractures.”</td>
<td>Data suggest efficacy without delaying union.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Sample Size</td>
<td>Description</td>
<td>Outcome Measures</td>
<td>Results</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>-------------</td>
<td>-------------</td>
<td>------------------</td>
<td>---------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adolphson 1993</td>
<td>6.0</td>
<td>N = 42 (42 female) Mean age and range 63 (52-79)</td>
<td>20mg a day per os piroxicam (Feldene®) for 8 weeks after initial 48 hours vs. 500mg paracetamol as rescue drug.</td>
<td>7% mean decrease in bone mineral content in radius after 8 weeks for piroxicam; 10% decrease in control (p = NS). Pain piroxicam/placebo 10 days: 2.1/3.1, 4 weeks 1.0/2.5, 8 weeks 1.0/0.9 (p &lt;0.05). Grip Strength piroxicam/placebo 10 days/4 weeks 10/6, 8 weeks 32/26 (p = NS).</td>
<td>&quot;The patients who received piroxicam had significantly less pain during plaster treatment, but there was no difference in the rate of functional recovery between the groups.&quot;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thomas 1986</td>
<td>5.0</td>
<td>N = 55 (21 males, 34 females) with fracture of distal end of radius, parallel to the wrist joint and with a tendency to dorsal displacement, treated in the normal way with no external skeletal fixation, fracture splinted in below-elbow plaster cast for between 4 to 6 weeks; mean age of both groups = 55 [no mean average listed for entire study population]</td>
<td>Normal treatment of fracture plus receiving three 50mg tablets of diclofenac, a prostaglandin inhibitor, a day for seven days (N = 29, Men = 10, Women = 19) vs Normal treatment of fracture plus receiving three 50mg placebo tablets a day for seven days (N = 26, Men = 11, Women = 15). Follow-up at two weeks after removal of cast</td>
<td>Comparison of loss of total range of movement between diclofenac vs. placebo groups – Student’s t test (0.05 &lt; P &lt; 0.1), Patients’ perception of pain between diclofenac vs. placebo groups – ratings from none (11 vs 6), improved (15 vs 17), no change (2 vs 2), worse (1 vs 1) – Chi squared (X² = 1.44, 0.05 &lt; P &lt; 0.1), Patients’ perception of stiffness between diclofenac vs. placebo groups – none (12 vs 3), improved (13 vs 20), no change (4 vs 3), worse (0 vs 0) – Chi squared (X² = 6.88, 0.05 &lt; P &lt; 0.1)</td>
<td>&quot;Both subjective and objective tests of recovery at 2 weeks after removal of splintage following fractures of the distal end of the radius showed that those patients treated with a prostaglandin inhibitor recovered better than those who received placebo. This form of treatment may prove most valuable in patients who might otherwise be slow to recover or in whom a rapid recovery is especially desirable.&quot;</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Initial Care

#### Non-Displaced Distal Radial Fracture

1. **Recommendation:** Immobilization Period of Three or Less Weeks (Early Mobilization) for Non-displaced or Minimally Displaced Distal Radius Fractures

   Immobilization of non-displaced or minimally displaced distal forearm fractures limited to 3 weeks is moderately recommended and has equivalent or superior functional outcomes than periods greater than 3 weeks for non-displaced or minimally displaced distal radius fracture.

   **Strength of Evidence** – Moderately Recommended, Evidence (B)
   
   **Level of Confidence** – Moderate

   **Rationale for Recommendation**

   Six moderate-quality studies (1273-1278) support limiting immobilization of non-displaced or minimally displaced non-articular fractures of the distal radius to a period of 3 weeks or less. The clinical definition of minimally displaced fractures, however, is not established by quality evidence, as the available literature lacks a consistent standardized fracture classification, such as Frykman’s or the AO classification systems. In general, the inclusion and exclusion criteria for entry into interventional studies reviewed in this Guideline may act as a defacto guideline, defining minimally displaced as fractures with less than 10° of dorsal angulation, less than 10° of radial angulation, and less than 2 to 3mm of radial shortening. (106, 1274, 1279-1281)

   In each study comparing immobilization of 3 or 5 weeks, patients demonstrated either improved functional measures such as pain scores (1274) wrist swelling, wrist and grip strength, and better subjective patient assessments with shorter immobilization periods, or no measurable differences between the groups indicating there is no advantage to longer immobilization periods. There were no differences in radiographic findings in any of the studies associated with duration of immobilization. (1273, 1275) There is no quality evidence supporting immobilization for periods greater than 3 weeks in these cases. Although there is one low-quality study that suggests equivalent functional results are achieved with fewer cases of complex regional pain syndrome (1 vs. 5), (106) there is insufficient evidence to support 1-week immobilization.

2. **Recommendation:** Use of Functional Brace or Splint over Traditional Casting for Non-displaced or Minimally Displaced Distal Radius Fractures

   The use of functional bracing or splinting that will allow mobilization of the radial-carpal joint while maintaining stabilization of the fracture is moderately recommended over traditional casting to immobilize the forearm and wrist for non-displaced or minimally displaced Colles’ fractures.

   **Strength of Evidence** – Moderately Recommended, Evidence (B)
   
   **Level of Confidence** – Moderate

   **Rationale for Recommendation**

   Copyright © 2016 Reed Group, Ltd.
There are multiple moderate-quality studies providing moderate evidence in support of functional bracing or splinting over traditional casting for non-displaced or minimally displaced fractures of the distal radius. (1274, 1275, 1282-1285) Functional bracing or splinting techniques described allow for mobilization at the radiocarpal joint. Various splinting techniques have been described, including the use of the lightweight removable splints, (1285) posterior splint with tubigrip, (1274) crépe bandage, (1275) elastic bandage, (1282) triple point loading brace with adjustable Velcro straps, (1284) and 3-point loading functional plaster brace. (1283)

As there are no direct comparisons between types of functional bracing, no specific recommendation can be made as to which if any technique is superior. The importance of early radiocarpal joint mobilization appears to be most important factor. Improved functional outcome through early mobilization may be a surrogate or confounder to the recommendation for shorter durations (3-week period) of immobilization, which essentially achieves the same objective of reducing immobilization of the radiocarpal joint. The literature is unclear if there might be an additive effect for functional bracing combined with immobilization of the fracture for 3 weeks or less, as functional bracing was compared to traditional casting of 4 to 6 weeks duration.

3. **Recommendation: Casting/Bracing Non-displaced or Minimally Displaced Colles’ Fractures in Pronation or Supination**
   
   There is no recommendation for or against casting/bracing the forearm and wrist in pronation or supination for non-displaced or minimally displaced Colles’ fractures.

   **Strength of Evidence – No Recommendation, Insufficient Evidence (I)**
   
   **Level of Confidence – Low**

   **Rationale for Recommendation**

   There are two moderate-quality studies on cast positioning of the forearm and hand, either supination or pronation, and functional outcomes. One study found no advantage to supination over conventional Colles’ casting; (1286) the other found forearm casting in pronation superior to above-elbow supination. (1287) As both techniques were last reported on more than 20 years ago, and with more recent evidence indicating that functional splinting is more effective casting, no recommendation is made regarding casting in pronation or supination in patients with non-displaced Colles’ fractures, although pronation is likely superior to supination if casting is attempted. Casting the forearm and wrist in pronation may provide benefit over casting in supination, although neither is recommended if functional bracing or splinting is an available treatment option.

**Evidence for Immobilization/Fixation for Non-displaced Colles’ Fracture**

There are 26 moderate-quality RCTs and 1 prospective study incorporated into this analysis. (106, 1239, 1348-1352, 1357-1361, 1363-1377) (Tumia 03; Bunger 84; Arora 11; Wik 09; Bong 06; Sarmiento 80; Gupta 91; Rosetzsky 82; Wahlstrom 82; Uzzaman 08; Ismatullah 12) There are 2 low-quality RCT in Appendix 1. (1375, 1378) (Gupta 11)

**Early Immobilization:**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Early Immobilization & Mobilization & Colles’ Fracture Or Distal Radial Fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 76 articles in PubMed, 30 in Scopus, 8 in CINAHL, 12,970 in Google Scholar, 18 in Cochrane Library, and 0 from other sources. We considered for inclusion 5 from PubMed, 5 from Scopus, 3 from CINAHL, 1 from Cochrane Library, 3 from Google Scholar, and 0 from other sources. Of the 17 articles considered for inclusion, 9 randomized trials and 8 systematic studies met the inclusion criteria.

**Functional Bracing:**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Functional Bracing & Casting, Distal Radius Fracture; splinting, splints, splintage, braces, bracing, functional, mobilization, rehabilitation, and recovery. We found and reviewed 76 articles in PubMed, 30 in Scopus, 8 in CINAHL, 12,970 in Google Scholar, 18 in Cochrane Library, and 0 from other sources. We considered for inclusion 5 from PubMed, 5 from Scopus, 3 from CINAHL, 1 from Cochrane Library, 3 from Google Scholar, and 0 from other sources. Of the 17 articles considered for inclusion, 9 randomized trials and 8 systematic studies met the inclusion criteria.
Radial Fractures or Colles’ Fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 7 articles in PubMed, 4 in Scopus, 1 in CINAHL, 5 in Cochrane Library, 11,230 in Google Scholar, and 0 from other sources. We considered for inclusion 4 from PubMed, 1 from Scopus, 1 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 0 from other sources. Of the 7 articles considered for inclusion, 6 randomized trials and 1 systematic studies met the inclusion criteria.

Casting:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Casting and Bracing and Colles’ Fractures Or distal Radial Fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 6 articles in PubMed, 35 in Scopus, 7 in CINAHL, 14 in Cochrane Library, 8830 in Google Scholar, and 0 from other sources. We considered for inclusion 3 from PubMed, 17 from Scopus, 1 from CINAHL, 0 from Cochrane Library, 4 from Google Scholar, and 0 from other sources. Of the 22 articles considered for inclusion, 18 randomized trials and 4 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moir 1995</td>
<td>RCT</td>
<td>6.5</td>
<td>N = 85 (70 females/9 males) individuals with distal Colles’ fracture; Median age. Group 1: 55 (22-86) Group 2: 60 (21-84)</td>
<td>Functional brace vs. control (dorsal plaster splint for 2 weeks followed by casting). Follow up at 10-14 days, 5-6 weeks, and 8, 13, and 26 weeks.</td>
<td>Functional score results: brace vs. control (lower score is better): 8 weeks 10 vs. 14 (p = 0.02); 13 weeks 4 vs. 11 (p = 0.003); 26 weeks 2 vs. 5 (p = 0.02). Grip strength as % of uninjured side: 8 weeks 50 vs. 35 (p = 0.0006); 26 weeks, 73 vs. 71 (p = 0.6). Analogue pain score (0-10): median: Splint removal 1 vs. 2 (p = 0.02); 8 weeks 1 vs. 2 (p = 0.048).</td>
<td>“The brace gave better functional results than conventional plaster treatment. The improved function was apparent up to 6 months after injury. Finger function and pinch strength were also better in the brace-treated patients. Anatomical results were similar in the two groups.”</td>
<td>The brace-treated fractures were initially less severely displaced than control fractures. “The improved functional results, particularly in terms of pinch and grip strength, are particularly important in the group of elderly patients who live alone.”</td>
</tr>
<tr>
<td>Stewart 1984</td>
<td>RCT</td>
<td>5.0</td>
<td>N = 243 (No mention of Gender) patients with fractures of the distal radius; No mention of Mean age.</td>
<td>Conventional Colles’ plaster vs. (Sarmiento) supinated cast-brace vs. below elbow cast-brace. Follow-Up at 6 weeks, and 3, 6 months.</td>
<td>Anatomical assessment excellent or good/total: plaster 45/93; supinated brace 43/70; short brace 43/72. Functional results mean score at 3 months/6 months: plaster 10.0/6.3; supinated brace 9.5/6.7; short brace 10.7/6.9. Incidence of carpal tunnel compression symptoms was 17% at 3 months and 12% at 6 months. No statistical significance between groups for incidence of symptoms.</td>
<td>“Early hand function and the supinated position advocated by Sarmiento were found to confer no anatomical or functional advantage; we could see no reason to change from the use of conventional plaster casts in the treatment of uncomplicated Colles’ fractures.”</td>
<td>Author suggests 4 indications for use of below-elbow cast brace: request by patient for complete freedom of movement of fingers and thumb; pre-existing finger stiffness or painful arthritis of carpometacarpal joint of thumb; the possibility that patient may develop Sudeck’s osteodystrophy; and to all direct access to the hand for dressings in patients with soft-tissue injuries.</td>
</tr>
<tr>
<td>Turnia 2003</td>
<td>RCT</td>
<td>5.0</td>
<td>N = 339 (31 male and 139 female) categorized into minimally displaced and displaced requiring manipulation groups. Mean age of 58.4 years.</td>
<td>Conventional Colles’ plaster cast (N = 163) vs Prefabricated functional brace or the Aberdeen Colles’ fracture brace (N = 166). Follow-up for 14 weeks.</td>
<td>Functional scores cast/brace non-manipulated group Week 8: 6.7/5.5; Week 24: 2.6/2.7 manipulated group Week 8: 11.4/10.6; Week 24: 5.4/5.8. Mean pain score cast/brace non-manipulated group 10 d: 2.2/2.4 p = 0.27; Week 24: 1.0/1.0 p = 0.96; manipulated group 10 d: 1.8/2.1 p = 0.19; Week 24: 0.5/0.5 p = 0.043.</td>
<td>“There was no significant difference in the functional outcome between the two treatment groups.”</td>
<td>Author comment on younger patients having better functional results not presented in body of study results. There appears to be no advantage to flexible brace over cast.</td>
</tr>
<tr>
<td>Study</td>
<td>Methodology</td>
<td>Design</td>
<td>N</td>
<td>Description</td>
<td>Follow-up</td>
<td>Study outcomes</td>
<td>Findings</td>
</tr>
<tr>
<td>---------------</td>
<td>-------------</td>
<td>--------</td>
<td>-------</td>
<td>-----------------------------------------------------------------------------</td>
<td>-----------</td>
<td>------------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Bürger 1984</td>
<td>RCT</td>
<td></td>
<td>4.5</td>
<td>N = 145 (20 male and 125 female) with Colles’ fracture. Age not given.</td>
<td>Follow-up</td>
<td>Functional bracing in supination or FUSU (N = 68) vs Dorsal Plaster Immobilization or DPI (N = 77).</td>
<td>&quot;Functional bracing in supination provided superior results in the treatment of particularly displaced intra-articular Colles’ fracture.&quot;</td>
</tr>
<tr>
<td>Abbaszadegan 1989</td>
<td>RCT</td>
<td></td>
<td>4.0</td>
<td>N = 80 (No mention of gender) &quot;un-displaced or minimally displaced Colles’ fractures&quot;; No mention of mean age.</td>
<td>Follow up</td>
<td>4 weeks in dorsal plaster cast vs. an elastic bandage. Follow up at 10-12 days. 1, 2, 3, and 6 months.</td>
<td>&quot;Elastic bandage treatment resulted in less pain, improved grip strength and better subjective satisfaction at one year. It did not result in increased fracture displacement when compared to conventional plaster splints. Functional treatment of the minimally displaced fractures.&quot;</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Methodology</td>
<td>N</td>
<td>Description</td>
<td>Results</td>
<td>Notes</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>-------------</td>
<td>---</td>
<td>-------------</td>
<td>---------</td>
<td>-------</td>
<td></td>
</tr>
</tbody>
</table>
| Christensen 1995 | RCT | No mention of sponsorship or COI. | 5.5 | Early Immobilization  
N = 33 patients with undisplaced fractures of the distal radius;  
Mean Age 3 wk group : 61 (29-78), 5 wk group: 64 (40-84).  
Immobilizing plaster splints at either 3 or 5 weeks for undisplaced fractures.  
Follow Up for 3 months and 9 months. | Differences in modified median Gartland and Werley scores at 3, 9 months insignificant (3 weeks/5 weeks): Score-3 months 2.0/3-0; Score: 9 months 1.0/1.0. | No difference in "radiological healing at 3 months or in the functional scores after 3 and 9 months." Early mobilization at 3 weeks appears to have no negative or positive impact on nondisplaced fractures. |
| Davis 1987 | RCT | No mention of sponsorship or COI. | 5.0 | Early Immobilization  
N = 55 (11 males/44 females) patients with slightly displaced fractures of the distal radius;  
Mean age, Group 1: 56.6 Group 2: 55.6  
After 2-week period of posterior splinting, patients randomized to tubigrip vs. below elbow cast for 3 additional weeks.  
Follow-up at 2, 5, and 7 weeks. | No significant difference of pain between groups. Gartland and Werley's functional assessment (excellent or good) total: Week 5 cast 11/25; Week 5 tubigrip (tg) 23/27 p <0.05; Wk 7 cast 22/25 p <0.01, p <0.05; Week 7 tg 25/27 p <0.01. Complications of treatment cast/tg: Fracture displaced 3/2; Physiotherapy needed 3/1. | "Unnecessary to splint slightly displaced fracture of the distal radial metaphysis for 4 weeks… a faster functional recovery will be obtained [if fractures are in an unrestrained tubigrip support] in a manner that had been shown to be acceptable to most patients." No blinding in this study. |
| Dias 1987 | RCT | No mention of sponsorship or COI. | 5.0 | Early Immobilization  
N = 187 (no mention of gender) patients with unilateral Colles' fractures that were older than 55;  
No mean age.  
Undisplaced fractures treated either with conventional 5 weeks cast (Group 1) or crêpe bandage (Group 2) and early mobilization. Displaced fractures were treated either with conventional 5 week cast (Group 3) or modified 5 week cast (Group 4).  
Follow-Up at weeks 1, 5, 9, and 13. | Early mobilization more resolution of wrist swelling first 5 weeks. At 9 and 13 weeks, wrist girth was similar. Deterioration rate of radiological deformity was similar in conventionally treated groups as with mobilization groups. Grip strength recovery expressed as a percentage of strength of contralateral hand much better in early mobilization groups. Undisplaced fractures Group 1/Group 2: Week 5 36.1/45.7 p <0.001; Week 9 51.7/63.5 p <0.005; Week 13 58.3/76.2 p <0.001. | "Early wrist movement hastened functional recovery and led to earlier resolution of wrist swelling. Discomfort was no greater than in patients who were treated conventionally. The bony deformity, which recurred irrespective of the method of treatment, was not adversely affected by early mobilization." This study includes weaknesses in randomization and baseline comparability. |
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Sponsorship or COI</th>
<th>Participants</th>
<th>Intervention</th>
<th>Follow-Up</th>
<th>Outcomes</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arora 2011</td>
<td>4.5</td>
<td>RCT</td>
<td>No sponsorship or COI</td>
<td>N = 73 (18 male and 55 female) with distal radial fracture; mean age 76.7 (65-89).</td>
<td>Group 1, operative group that underwent Open reduction and internal fixation (ORIF) 12 weeks after injury (N = 36) vs Group 2, immobilized in short arm cast for 5 weeks (N = 37).</td>
<td>Follow-up at baseline, 6, 12 weeks, 6 and 12 months.</td>
<td>Disabilities of the Arm, Shoulder and Hand Score (DASH) at 6 weeks group 1 vs 2; 18.8±17.9 vs 34.4±22.5 (p = 0.00). At 12 weeks; 13.3±14.8 vs 23.2±19.3 (p = 0.02).</td>
<td>Significantly more complications in operative group, 13 vs 5 (p&lt;0.05).</td>
</tr>
<tr>
<td>McAuliffe 1987</td>
<td>4.5</td>
<td>Prospective RCT</td>
<td>No mention of sponsorship or COI</td>
<td>N = 108 (All Women) who had a Colles’ fractures.</td>
<td>Plaster immobilization for 3 (Group A) or 5 (Group B) weeks.</td>
<td>Follow Up baseline, 3 months, and 1 year.</td>
<td>72% of Group A reported good or excellent results relating to pain, disability, and range of movement at 3 months while 66% of Group B did; after 1 year 85% of Group A had a good or excellent result and 77% did in Group B. Group A showed statistical significance for pronation after 1 year, less pain at time of plaster removal, 3 months and 1-year follow up as well as stronger grip strength after 1 year.</td>
<td>In this elderly population, mobilization after 3 weeks may lead to less short-term disability.</td>
</tr>
<tr>
<td>Millet 1995</td>
<td>4.0</td>
<td>RCT</td>
<td>No significant clinical differences found between the treatment groups.</td>
<td>N = 90 female with unilateral Colles’ fracture;</td>
<td>5 week below elbow plaster cast (N = 45)</td>
<td>All patients in early mobilization reported greater comfort after switching from</td>
<td>“Early mobilization produced less pain and a stronger grip. It did not lead to any greater loss of reduction of the fracture. However, there was no significant improvement in the final range of movement of the wrist.”</td>
<td>Data suggest at 12 months, ROM, pain level and PRWE and DASH scores equivalent. Patients in surgical group reported better grip strength throughout trial.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>N</td>
<td>Population</td>
<td>Intervention</td>
<td>Outcome Measures</td>
<td>Findings</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>---</td>
<td>------------</td>
<td>--------------</td>
<td>-----------------</td>
<td>---------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prospective Study</td>
<td>1998</td>
<td>4.0</td>
<td>N = 52</td>
<td>Plaster immobilization for 1 week vs. plaster immobilization for 3 weeks in minimally displaced fractures.</td>
<td>Functional Cooney score; 1 week (SD) vs. 3 weeks (SD): 6 weeks 61.6 (12.1) vs. 56.8 (19.7) 3 months 77.4 (13.8) vs. 71.5 (19.2); 1 year 86.8 (10.9) vs. 82.2 (18.6). One-week group Cooney score generally higher at every re-evaluation period than 3-week plaster group. Differences, however, not statistically significant.</td>
<td><em>No dislocations occurred. All patients experienced eventful healing with good or excellent results in 92% of cases. We believe, therefore, that only minimal immobilization is required in these fractures and that they should be mobilized as soon as comfort allows.</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stoffelen</td>
<td>2010</td>
<td>6.5</td>
<td>N = 60 (Predominately female) elderly Chinese people with dorsal angulated fracture of the distal radius;</td>
<td>Group 1 (N=30) Patients given plaster of Paris cast preceded by closed reduction. Vs Group 2 (N=30) Patients were treated with K-Wire Follow up at 2 weeks prior to injury and then 1, 2, 4, 6 weeks, 3, 6 months and 1 year after assessment of radiographs.</td>
<td>No statistically significant differences between the K-Wire treatment and the plaster of Paris group.</td>
<td><em>Although our study showed that the ‘tripod technique’ [K-Wire] is safe without significant complications, there is an Cochrane review of wiring for distal radial fractures… We do not provide a biomechanical rationale to explain our ‘tripod technique’ but we feel that it is a better construct to prevent collapse of the fracture/</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>O’Connor</td>
<td>2003</td>
<td>6.5</td>
<td>N = 66 (22 males/44 females) adult patients with minimally displaced radial fractures;</td>
<td>(N=32) Below-the-elbow plaster of Paris cast vs. (N=34) lightweight removable “Futuro” splint for minimally displaced Colles’ fractures;</td>
<td>No significant differences in pain scores. Cast satisfaction higher in splint group at weeks 1, 2, and 6. No difference in anatomical outcome. Functional scores and wrist range of motion</td>
<td>*A lightweight splint provides an acceptable, comfortable and economic alternative to plaster of Paris and allows faster restoration of function without an</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Casting/Bracing</em></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td><em>Patients in splint group were educated on rationale for splint use as authors found cultural bias toward the traditional cast.</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Mean Age, Group 1: 56.6 (16-81), Group 2: 57 (18-79)</td>
<td>Follow-up at 1, 2, 6, and 12 weeks.</td>
<td>were better at 6 weeks, but the differences disappeared at 12 weeks.</td>
<td>increased risk of malunion.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>--------------------------------------------------</td>
<td>------------------------------------</td>
<td>----------------------------------------------------------------</td>
<td>-----------------------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ledingham 1991</td>
<td>N = 60 (50 females/10 males); Mean Age, Group 1: 60.2, Group 2: 61.3.</td>
<td>(N=30) Plaster-of-Paris functional brace (brace) vs. (N=30) Standard below-elbow cast (control). Follow-Up at baseline, 24 hrs, 7-14 days, and 35-42 days.</td>
<td>Final radiological result; Brace vs. Control: Overall, brace group had better radiological results than control (lower score better) mean score 2.5 vs. 4.3 (p &lt;0.05). No significant differences between &lt;60-years-old brace and control or between brace under and over 60 years old. Significant difference in controls vs. under 60 years (12.7 vs. 4.4, p &lt;0.005). Functional grading results (Excellent + Good) using modified Gartland and Werley significant difference of brace vs. control at 12 weeks, but not at 26 weeks.</td>
<td>&quot;With the Plaster-of-Paris brace described in this paper, we have shown improved final radiological and early functional results compared to the standard below-elbow cast.&quot;</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Grafstein 2010</td>
<td>N = 101 (78 females/23 Males) with a displaced fracture of the distal radius requiring closed reduction.</td>
<td>Circumferential casting or CC (N = 40) vs Volar–dorsal splinting or VDS (N = 31) vs Modified sugar-tong splinting or MST (N = 30). Follow-up at 8 weeks and 6 months.</td>
<td>Median pain scores were not statistically different between the groups. 22 patients (22%, 95% CI: 13.9%-30.1%) had radiographic loss of reduction: VDS= 5 patients (16%, 95% CI 3.1%-28.9%), CC= 8 patients (20%, 95% CI: 7.6%-32.4%), and MST= 9 patients (30%, 95% CI: 13.6%-46.4%) (p = 0.17).</td>
<td>&quot;Rates of loss in anatomic position were not statistically significant among the 3 types of dressings used. However, there was a clinically important trend of increased loss of reduction with the use of MST splinting.&quot;</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Authors demonstrated in radiographic and functional grading that patients over 60 years old may benefit the most, although sample sizes were small.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Gender</th>
<th>Age</th>
<th>Fracture Type</th>
<th>Treatment</th>
<th>Follow up</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moroni 2004</td>
<td>RCT</td>
<td>40 (All female)</td>
<td>Osteoporotic patients who are 65 years of age or older; Mean Age &gt;65 years old</td>
<td>Group 1 (N=?) Patients who received plaster cast with closed reduction. Vs Group 2 (N=?) Received external fixation. Follow up at 2 weeks, 6 weeks, and 3 months.</td>
<td>Redisplacements Group 1 vs Group 2: 4 vs 0. Volar Angle at post op group 1 vs group 2: 8.6±5.8 vs 3.4±1.8. At 6 weeks: -1.9±9.4 vs 1.9±3.4 (p&lt;0.0005). Radials Angle at post op, group 1 vs 2: 20.6±4.9 vs 23.5±3.5. At 6 weeks, 17.1±6.3 vs 23.3±3.5 (p=0.008). Horesh Demerit Point Score at 3 months, Group 1 vs Group 1: 7.7±3.3 vs 6.6±3.4 (p&lt;0.006).</td>
<td>In conclusion, our study supports the use of external fixation in the treatment of osteoporotic wrist fractures. Both radiographic and clinical results were better in the external fixation group than in the plaster cast group.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cohen 1997</td>
<td>Prospective RCT</td>
<td>30 (22 females/8 male)</td>
<td>who had varying degrees of Radial distal fractures; Mean Age Group 1 56 (33-89) Group 2 58 (19-86)</td>
<td>Group 1 (N=10) Non displaced fractures, 5 fiberglass tape, 5 QuickCast tape. Vs Group 2 (N=10) (Displaced but stable after reduction fractures) 5 in QuickCast, 5 with fiberglass tape. Vs Group 3 (N=10) (Displaced fractures requiring Pin fixation) 5 with quick cast, 5 with fiberglass. Follow up at cast removal 5.5 weeks to 6.5 weeks</td>
<td>Number of cast Applications, Group 1, 2, and 3, Fiberglass vs QuickCast: Group 1; 2.2 vs 1.2, Group 2; 2.2 vs 1.0, Group 3; 3.0 vs 2.0. (p&lt;0.001). Problems with cast answer (1-10) Fiberglass vs Quickcast: 1.0±0.8 vs 0.5±0.4 (p&lt;0.001). Some cast complications within both groups, not significant.</td>
<td>In sum, a short-arm cast constructed of the QuickCast does save approximately one cast change in the treatment of distal radius fractures with no apparent effect on fracture healing. The QuickCast does, however, cost more in materials alone. This financial differential must be weighed against the labor saved of a single cast application with additional savings of time for the applicer and patient.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cohen 2001</td>
<td>RCT</td>
<td>200 (No mention of gender)</td>
<td>patients who sustained arm or leg injuries required cast support (N=29 individuals with Radial Distal fracture);</td>
<td>Group 1 (N=14) patients with Forced Rigidity Casting (FRC) Vs Group 2 (N=15) patients treated with Complete Plaster of Paris synthetic cast (Standard)</td>
<td>Increase in Ability FRC vs Standard: favored group 1 (p=0.0002). Satisfaction better in FRC group (p=0.0009).</td>
<td>Data suggest increased patient satisfaction with FRC vs. conventional plaster of Paris cast with comparable efficacy.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>Sample Size</td>
<td>Age Range</td>
<td>Fracture Type</td>
<td>Treatment</td>
<td>Follow-Up</td>
<td>Result</td>
</tr>
<tr>
<td>------------------</td>
<td>------</td>
<td>-----------</td>
<td>-------------</td>
<td>------------</td>
<td>---------------</td>
<td>-----------</td>
<td>-----------</td>
<td>--------</td>
</tr>
<tr>
<td>Wik 2009</td>
<td>4.0</td>
<td>RCT</td>
<td>72</td>
<td>Mean Age: 50</td>
<td>Colles'</td>
<td>Cast v. Splint</td>
<td>5 weeks</td>
<td>Reduction and a complete plaster cast (N=34) vs reduction and a dorsal plaster splint (N=38). Immobilization for 5 weeks with follow-up at 1 and 10 days and 5 weeks after reduction.</td>
</tr>
<tr>
<td>Bong 2006</td>
<td>4.0</td>
<td>Prospective RCT</td>
<td>85 (85 fractures, 26 male and 59 female)</td>
<td>Mean age: 64</td>
<td>Distal radial</td>
<td>Splint</td>
<td>7-10 days</td>
<td>Group 1 immobilized using short-arm radial gutter splint (N = 38) vs Group 2 immobilized with sugar tong splint. Radiographs taken in respective splint.</td>
</tr>
<tr>
<td>Sarmiento 1980</td>
<td>4.0</td>
<td>RCT</td>
<td>156</td>
<td>Median age: 49</td>
<td>Colles'</td>
<td>Cast / Splint</td>
<td></td>
<td>Bracing in either pronation, fractures were immobilized in a long-arm cast with the wrist at 20° of volar flexion and ulnar deviation; the elbow at 90° of flexion and the forearm in either pronation (N = 78) vs Supination the elbow at 90° of flexion and the forearm in supination</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>Sponsorship/COI</td>
<td>Participants</td>
<td>Intervention 1</td>
<td>Intervention 2</td>
<td>Follow-up</td>
<td>Results</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>------</td>
<td>--------</td>
<td>-----------------</td>
<td>--------------</td>
<td>----------------</td>
<td>----------------</td>
<td>-----------</td>
<td>---------</td>
</tr>
<tr>
<td>Gupta 1991 RCT</td>
<td>4.0</td>
<td>N = 204 (82 male and 122 female) with displaced Colles’ fractures. Mean age 46 years.</td>
<td>Plaster immobilization with either: Palmar flexion or PF (N = 60) vs Neutral or NP (N = 75) vs Dorsiflexion or DF wrist position (N = 69).</td>
<td>Follow-up for 15 weeks.</td>
<td>Functional results excellent or good/total: Type III PF 20/28; NP 26/34; DF 28/32 Type IV PF 10/17; NP 8/19; DF 15/17; Type V PF 9/15; NP 13/22; DF 16/20</td>
<td>“After manipulation of a Colles’ fracture, immobilization of the wrist in dorsiflexion would appear to provide better maintenance of reduction.”</td>
<td>Immobilization of wrist in palmar flexion had detrimental effect on hand function.</td>
<td></td>
</tr>
<tr>
<td>Rosetzsky 1982 RCT</td>
<td>4.0</td>
<td>N = 46 (15 male and 35 female) with Colles’ fractures of the forearm. Mean age was 45 years.</td>
<td>Polyurethane casts (N = unknown) vs Traditional plaster-of-Paris braces (N = unknown).</td>
<td>Follow-up at 6 weeks.</td>
<td>No significant difference for secondary adjustment of casts between groups, (p &gt; 0.90). No significant differences for failure of retaining fracture reduction, (p &gt; 0.50).</td>
<td>“Polyurethane braces are a good supplement to plaster-of-Paris bandage in such fractures and recommended in selected cases.”</td>
<td>Alternative to plaster of Paris in 1980s.</td>
<td></td>
</tr>
<tr>
<td>Wahlström 1982 RCT</td>
<td>4.0</td>
<td>N = 42 (all women) with extra articular fractures. Mean age 65 years.</td>
<td>Immobilization in pronation (N = 14) vs Supination (N = 12) vs Midway position (N = 16).</td>
<td>Follow-up at 10 days and 1-4 months after reduction.</td>
<td>Five fractures had to be re-reduced, one from pronation, one from midway and three from supination group. Patients with redislocation ≥10° number pronation 2/14, midway 6/12, and supination 8/16.</td>
<td>“The position of the forearm during immobilization is of importance for the degree of redislocation.”</td>
<td>Applicable to cast application rather than long-term functional results.</td>
<td></td>
</tr>
<tr>
<td>Uzzaman 2008 RCT</td>
<td>4.0</td>
<td>(N=40) (19 females/11 males) patients with displaced Colles fracture at the emergency of outpatient department</td>
<td>Closed reduction and two crossed percutaneous Kirschner wire fixation combined with plaster cast support (Arm A, N=20) v. conventional method-reduction by closed manipulation and maintained by plaster cast immobilization (Arm B,</td>
<td>Anatomic end result of Arm A was better than Arm B, p&lt;0.05. There was a significant Satisfactory result in Arm A compared Arm B, p&lt;0.05.</td>
<td>“Closed reduction and Percutaneous kirschner wire fixation combined with plaster cast immobilization is better method than the conventional plaster cast immobilization – in restoration of preinjury anatomical alignment</td>
<td>Data suggest percutaneous fixation group superior to cast alone group for maintaining radial length and angulation resulting in better function and also had less reported complications.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Methodology</td>
<td>Number of Patients</td>
<td>Description</td>
<td>Outcome</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>-------------</td>
<td>--------------------</td>
<td>-------------</td>
<td>---------</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ismatullah 2012</td>
<td>Prospective RCT</td>
<td>(N=30, 13 males/17 females) adult patients with a comminuted distal radius fracture; Mean Age 49.8 ±16.05</td>
<td>Group 1 (N=15) Treated with plaster casting. Vs. Group 2 (N=15) Treated with external fixation. Follow-Up at baseline and 12 weeks.</td>
<td>Green &amp; O’Brien Criteria rankings, Group 1 v 2, 12 weeks; Group 1: 4 were excellent, 3 were good, 4 were fair, 4 were poor. Group 2: 5 were excellent, 6 good, 2 fair, and 2 poor. “We recommend external fixation in comminuted fractures of the distal radius, which are potentially unstable fractures. It decreases the complications of re-displacement and shortening which may occur when these fractures are managed with closed reduction and casting.”</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Data suggest external fixation in comminuted distal radius fractures better than casting.
**Displaced Distal Radial Fracture**

Distal radial fractures with radiographic measurements of 10° or more of dorsal angulation, more than 2 mm of radial shortening or with any degree of radial shift require reduction to reduce the risk for deformity and disability. Closed reduction should result in no more than 5° of dorsal angulation and no more than 2 mm of radial shortening. Unstable fractures are defined as fractures with bone loss or bone involvement that will not allow for structural integrity without the use of internal or external fixation of the bone. Examples include fractures with dorsal comminution or radial lateral shift of more than 2 mm, have been proposed as limits for consideration of surgical intervention.(107, 108)

**CLOSED REDUCTION TECHNIQUE**

*Recommendation: Closed Reduction Technique for Displaced Distal Radial Fractures*

Manipulation and dynamic traction devices are recommended for closed reduction technique for displaced distal radial fractures as they have demonstrated equivalent ability to achieve initial reduction of injury.

- **Strength of Evidence** – Recommended, Evidence (C)
- **Level of Confidence** – Moderate

**Rationale for Recommendation**

For closed reduction, there is one high-quality and two moderate-quality studies comparing the effectiveness of manipulation and traction techniques for displaced Colles’ fracture. There was no difference in immediate reduction results using Chinese finger dynamic traction devices compared to manipulation under anesthesia.(1279, 1293, 1294) Long-term outcomes also showed no differences in post reduction failures, as both methods have 25 to 29% loss of reduction with casting. It is likely the loss of reduction is unrelated to reduction technique, and rather more related to immobilization technique. In a group of elderly patients, there were no differences in functional outcomes or deformity between those that underwent manipulation and casting versus those that were non-reduced and casted if the degree of displacement had less than 30° of dorsal angulation and 5 mm of radial shortening.(1295)

Despite non-superiority of reduction outcomes for manipulation or dynamic traction, one author in two papers reports lower rates of severe reduction pain and reduced long-term neurological deficits with dynamic traction (paresthesia, reduced 2-point discrimination) compared with manipulation under hematoma block.(1293, 1294) This author suggests the difference may have been related to the anesthetic technique rather than the reduction technique.

As noted earlier, the lack of a standard fracture classification system across each of these studies inhibits prognostic or treatment indications to be generalized. For these studies, Earnshaw used criteria of >10° of dorsal angulation, > 5 mm radial shortening, no marked comminution or displacement of articular surfaces. Kongsholm included mostly Frykman II, VI, VII, VIII fractures in the study, which may have included comminuted fractures, and those enrolled by Kelly included up to 5 mm of radial shortening and 30° of dorsal angulation.

**Evidence for the Use of Closed Reduction Technique for Distal Radial Fractures**

There are 4 moderate-quality RCTs incorporated into this analysis.(1279, 1293-1295)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: closed reduction technique, distal, forearm, radial, radius fractures, bone fractures, colles’ fracture, displaced; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 14 articles in PubMed, 24 in Scopus, 13 in CINAHL, 0 in Cochrane Library, 19930 in Google Scholar, and 0 from other sources. We considered for inclusion 4 from PubMed, 0 from Scopus, CINAHL, Cochrane Library, Google Scholar, and from other sources. Of the 4 articles considered for inclusion, 4 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Earnshaw 2002 RCT No sponsorship or COI.</td>
<td>7.5</td>
<td>N = 225 (172 female/53 male) displaced Colles-type fractures.</td>
<td>Closed reduction with either finger-trap (n = 112) vs. Manual manipulation (n = 111).</td>
<td>87% of fractures were successfully reduced. &quot;By five weeks, fifty-six (25%) of the 225 fractures had been treated with surgical intervention because of failed closed treatment and only sixty-five (29%) remained in a satisfactory position.&quot;</td>
<td>&quot;The two methods of fracture reduction did not differ with regard to the eventual position of the fracture or the rate of failure.&quot;</td>
<td>All reductions performed post Bier's block. Loss of reduction during the period of cast immobilization is common in this study.</td>
</tr>
<tr>
<td>Kongsholm J Orthop Trauma 1987</td>
<td>5.5</td>
<td>N = 116 with acute displaced Colles' fracture</td>
<td>(Group A) dynamic reduction device with no anesthesia vs. (Group B) 8-10ml of 1% lidocaine with plaster cast.</td>
<td>2/62 patients in Group A displayed symptoms and signs of neurological impairment at 5 weeks compared to 11/54 patients in Group B, p &lt;0.01. 1 year follow up resulted in figures of 4/62 and 8/54 with p &lt;0.05.</td>
<td>&quot;The dynamic reduction technique without local anesthesia results in a significantly lower frequency of neurological complication than manual reduction after injection of local anesthetic into the fracture hematoma.&quot;</td>
<td>The neurologic complications included subjective paresthesia, positive Tinel's sign or 2 point discrimination &gt;4mm. Authors note &quot;nerve damage&quot; was mild and in no case in either group did it lead to surgical neurolysis.</td>
</tr>
<tr>
<td>Kelly 1997 RCT</td>
<td>5.0</td>
<td>N = 30</td>
<td>Reduction of the fracture under Bier’s block then immobilized in dorsoradial plaster of Paris slab compared to plaster immobilization only in elderly population.</td>
<td>11/15 in Bier’s block group and 9/15 in immobilization only group considered that their wrist was of normal appearance or had only slight deformity visible. Functional outcome Bier’s block/immobilization: Gartland and Werley score 5.8/6.6. Grip strength % predicted 48.8±17%/55.8±19%.</td>
<td>&quot;There was no detectable difference between the groups in any of the outcome measures.&quot;</td>
<td>Study suggests reduction does not provide any benefit over risk of Bier’s block to the elderly population within the parameters of 30° of dorsal angulation and 5mm of radial shortening.</td>
</tr>
<tr>
<td>Kongsholm Injury 1987</td>
<td>4.0</td>
<td>N = 116</td>
<td>(Group A) dynamic bone alignment device compared without anesthesia to (Group B) traditional manual reduction using local infiltration anesthesia</td>
<td>No differences between the groups in &quot;no pain&quot; or &quot;slight pain.&quot; However, for severe pain Group B had 19 vs. 5 patients, p &lt;0.001.</td>
<td>&quot;Dynamic reduction without anesthesia seems to be a less painful method for the patients than traditional manual reduction under local anesthesia.&quot;</td>
<td>Study did not follow longitudinal results of reduction.</td>
</tr>
</tbody>
</table>
CASTING OR FUNCTIONAL BRACING

Unlike casting for non-displaced and minimally displaced distal radial fractures, there are few studies comparing casting technique and functional bracing for displaced distal radial fractures and most of the available work was conducted more than 20 years ago. There is no defined standard for casting technique and forearm positioning that provides significant advantage over any other technique for displaced distal forearm fractures.

**Recommendation: Use of Functional Brace or Splint over Traditional Casting for Displaced Distal Radial Fracture**

There is no recommendation for or against the use of a functional brace or splint that will allow mobilization of the hand while maintaining stabilization of the reduced displaced distal radial fracture.

**Strength of Evidence – No Recommendation, Insufficient Evidence (I)**

**Level of Confidence – Low**

**Rationale for Recommendation**

There are few studies that compare specific casting or immobilization techniques. Rather, bracing and casting has been studied in the greater context of allowing hand functionality (functional brace) compared with traditional Colles’ casting (elbow flexion, forearm pronation with ulnar deviation) as well as position of the wrist (palmar flexion, neutral, dorsiflexion) and forearm position (pronation, supination).(1277, 1296-1301) One moderate-quality study of 339 patients with non-specific displaced fractures showed no difference in casting versus functional bracing.(1300) Two moderate quality studies found bracing in the supine position may have advantages for intra-articular fractures,(1296, 1299) whereas bracing in pronation may provide advantage for extra-articular fracture.(1297) However, another moderate-quality study with 250 participants found no differences between hand and ulnar positioning.(1302) In several of these studies, the authors concluded results were related to displacement of original fracture and degree of successful reduction more than immobilization technique. Thus, there are insufficient data to recommend specific casting or immobilization techniques for displaced Colles’ fractures.

**Evidence for the Use of Casting/Functional Bracing for Displaced Forearm Fractures**

There are 10 moderate-quality RCTs or prospective studies incorporated into this analysis.(1277, 1292, 1296-1301, 1303, 1304) There is 1 low-quality RCT in Appendix 2.(1302)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: casting or functional bracing, displaced distal radial fracture, distal, forearm, radial, radius fractures, bone fractures, colles’ fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 30 articles in PubMed, 13 in Scopus, 1 in CINAHL, 41 in Cochrane Library, 3174 in Google Scholar, and 7 from other sources. We considered for inclusion 4 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 7 from other sources. Of the 11 articles considered for inclusion, 11 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/ Year Study Type (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Casting or Functional Bracing</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tumia 2003 RCT</td>
<td>5.0</td>
<td>N = 339 (31 male and 139 female) categorized into minimally displaced and displaced requiring manipulation groups. Mean age of 58.4 years.</td>
<td>Conventional Colles’ plaster cast (N = 163) vs Prefabricated functional brace or the Aberdeen Colles’ fracture brace (N = 166). Follow-up for 14 weeks.</td>
<td>Functional scores cast/brace non-manipulated group Week 8: 6.7/5.5; Week 24: 2.6/2.7 vs manipulated group Week 8: 11.4/10.6; Week 24: 5.4/5.8. Mean pain score cast/brace non-manipulated group 10 d: 2.2/2.4 p = 0.27; Week 24: 1.0/1.0 p = 0.96; manipulated group 10 d: 1.8/2.1 p = 0.19; Week 24: 0.5/0.5 p = 0.043.</td>
<td>“There was no significant difference in the functional outcome between the two treatment groups.”</td>
<td>Author comment on younger patients having better functional results not presented in body of study results. There appears to be no advantage to flexible brace over cast.</td>
</tr>
<tr>
<td><strong>Arora 2011 RCT</strong></td>
<td>4.5</td>
<td>N = 73 (18 male and 55 female) with distal radial fracture; mean age 76.7 (65-89).</td>
<td>Group 1, operative group that underwent Open reduction and internal fixation (ORIF) 12 weeks after injury (N = 36) vs Group 2, immobilized in short arm cast for 5 weeks (N = 37). Follow-up at baseline, 6, 12 weeks, 6 and 12 months.</td>
<td>Disabilities of the Arm, Shoulder and Hand Score (DASH) at 6 weeks group 1 vs 2; 18.8±17.9 vs 34.4±22.5 (p = 0.00). At 12 weeks; 13.3±14.8 vs 23.2±19.3 (p = 0.02). Patient-Rated Wrist Evaluation (PRWE) group 1 vs 2, at 6 weeks; 36.4±28.7 vs 64.9±29.0 (p = 0.00), at 12 weeks; 33.7±32.0 vs 54.4±31.8 (p = 0.01). Grip Strength (kg) group 1 vs 2, 6 weeks; 14.1±10.7±5.6 (p = 0.01). At 12 weeks; 15.7±6.2 vs 2.5±4.4 (p = 0.02). At 6 months; 19.8±7.4 vs. 16.1±5.6 (p=0.02). At 12 months; 22.2±6.3 vs 18.8±5.8 (p = 0.02). Significantly more complications in operative group, 13 vs 5 (p&lt;0.05).</td>
<td>“[H]owever, at twelve months after surgery, the active range of motion, the pain level, and the PRWE and the DASH scores were not different between the operative and nonoperative treatment groups.”</td>
<td>Data suggest at 12 months, ROM, pain level and PRWE and DASH scores equivalent. Patients in surgical group reported better grip strength throughout trial.</td>
</tr>
<tr>
<td>Bünger 1984 RCT</td>
<td>4.5</td>
<td>N = 145 (20 male and 125 female) with Colles’ fracture. Age not given.</td>
<td>Functional bracing in supination or FUSU (N = 68) vs Dorsal Plaster Immobilization orDPI (N = 77).</td>
<td>Primary treatment; DPI vs. FUSU: Anatomic end results (excellent/good)/total 65/72 vs. 59/64 (p &lt;0.05). Functional results at 6 months (excellent/good)/total 62/72 vs. 59/62 (p &lt;0.5)</td>
<td>“Functional bracing in supination provided superior results in the treatment of particularly displaced intra-articular Colles’ fracture.”</td>
<td>Suggests the functional benefit from FUSU is primarily secondary to decreased fracture redislocation.</td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Level</td>
<td>Participants</td>
<td>Intervention</td>
<td>Outcomes</td>
<td>Comments</td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>-------</td>
<td>--------------</td>
<td>--------------</td>
<td>----------</td>
<td>----------</td>
</tr>
</tbody>
</table>
| Wik 2009 | RCT | 4.0 | N = 72 females who sustained low-energy trauma and displaced Colles’ fractures initially suitable for closed reduction and immobilization in plaster cast. Age >50. | Reduction and a complete plaster cast (N = 34) vs Reduction and dorsal plaster splint (N = 38). Immobilization for 5 weeks with follow-up at 1 and 10 days and 5 weeks after reduction. Mean dorsal angulation 10 days after reduction: slightly better in the dorsal plaster splint group, (p = 0.04). Radial length at 5 weeks was better in the complete plaster group, (p = 0.02). | “[S]urgeons caring for such cases may choose the immobilization method for the first 10 days following reduction according to their individual preferences and those of the injured person.” |}
<p>| Bong 2006 | Prospective RCT | 4.0 | N = 85 (85 fractures, 26 male and 59 female) who were used had acquired a displaced distal radial fracture; mean age 64 (27-91). | Group 1 immobilized using short-arm radial gutter splint (N = 38) vs Group 2 immobilized with sugar tong splint. Follow-up 7-10 days after initial injury (N = 47). Radiographs taken in respective splint. No significant difference between loss of fracture reduction, volar tilt, radial height, radial inclination. Disabilities of the ARM, Shoulder, and Hand (DASH) scores, Group 1 vs group 1 at 1 week; 62±19 vs 70±15 (p=0.044). | &quot;Based on our study we recommend that surgeons consider using a short-arm radial gutter splint for the initial immobilization of displaced distal radius fractures.” | Sparse baseline comparability details. Data suggest both long and short arm splints are effective in maintaining the reduction of distal radius fractures but the short arm splint was preferred by patients. |
| Millet 1995 | Prospective Study | 4.0 | N = 90 female with unilateral Colles’ fracture. Mean age of 61 years. | 5 week below elbow plaster cast (N = 45) vs 3 week plaster cast with 2 week flexible cast. Displaced fractures in both groups were manipulated. (N = 45). Patients followed for 3 years. | All patients in early mobilization reported greater comfort after switching from plaster to flexible casting. Mean grip scores and joint mobilities higher at all time points with early mobilization, reaching levels of statistical significance at 6, (p &lt; 0.01) months for grip score and 3 months for joint mobility, (p = 0.04). | “Early mobilization is a satisfactory treatment option for Colles’ fracture and may, in fact, hasten functional recovery.” |
| Rosetzsky 1982 | RCT | 4.0 | N = 46 (15 male and 35 female) with Colles’ fractures of the forearm. Polyurethane casts (N = unknown) vs | No significant difference for secondary adjustment of casts between groups, (p &gt; 0.90). No significant differences for failure of “Polyurethane braces are a good supplement to plaster-of-Paris bandage in such | No significant clinical differences found between the treatment groups. | Alternative to plaster of Paris in 1980s. |</p>
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>Participants</th>
<th>Treatment</th>
<th>Follow-up</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sarmiento 1980 RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>N = 156 (50 male and 106 female) with Colles’ fractures. Mean age of 49 years.</td>
<td>Traditional plaster-of-Paris braces (N = unknown). Follow-up at 6 weeks.</td>
<td>retaining fracture reduction, (p &gt; 0.50). fractures and recommended in selected cases.”</td>
<td>“Treatment with functional bracing in supination position yielded 90% excellent or good functional results.”</td>
</tr>
<tr>
<td>Gupta 1991 RCT</td>
<td>No sponsorship. No mention of COI.</td>
<td>N = 204 (82 male and 122 female) with displaced Colles’ fractures. Mean age 46 years.</td>
<td>Plaster immobilization with either: Palmar flexion or PF (N = 60) vs Neutral or NP (N = 75) vs Dorsiflexion or DF wrist position (N = 69). Follow-up for 15 months.</td>
<td>Functional results excellent or good/total: Type III PF 20/28; NP 26/34; DF 28/32 Type IV PF 10/17; NP 8/19; DF 15/17; Type V PF 9/15; NP 13/22; DF 16/20</td>
<td>“After manipulation of a Colles’ fracture, immobilization of the wrist in dorsiflexion would appear to provide better maintenance of reduction.”</td>
</tr>
<tr>
<td>Wahlström 1982 RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>N = 42 with extra articular fractures. Mean age 65 years.</td>
<td>Immobilization in pronation (N = 14) vs Supination (N = 12) vs Midway position (N = 16). Follow-up at 10 days and 1-4 months after reduction.</td>
<td>Five fractures had to be re-reduced, one from pronation, one from midway and three from supination group. Patients with redislocation ≥10° number pronation 2/14, midway 6/12, and supination 8/16.</td>
<td>“The position of the forearm during immobilization is of importance for the degree of redislocation.”</td>
</tr>
</tbody>
</table>

Gupta 1991 RCT
No sponsorship. No mention of COI.

4.0
N = 204 (82 male and 122 female) with displaced Colles’ fractures. Mean age 46 years.

Plaster immobilization with either: Palmar flexion or PF (N = 60) vs Neutral or NP (N = 75) vs Dorsiflexion or DF wrist position (N = 69). Follow-up for 15 months.

Functional results excellent or good/total: Type III PF 20/28; NP 26/34; DF 28/32 Type IV PF 10/17; NP 8/19; DF 15/17; Type V PF 9/15; NP 13/22; DF 16/20

“After manipulation of a Colles’ fracture, immobilization of the wrist in dorsiflexion would appear to provide better maintenance of reduction.”

Immobilization of wrist in palmar flexion had detrimental effect on hand function.

Gupta 1991 RCT
No sponsorship. No mention of COI.

4.0
N = 204 (82 male and 122 female) with displaced Colles’ fractures. Mean age 46 years.

Plaster immobilization with either: Palmar flexion or PF (N = 60) vs Neutral or NP (N = 75) vs Dorsiflexion or DF wrist position (N = 69). Follow-up for 15 months.

Functional results excellent or good/total: Type III PF 20/28; NP 26/34; DF 28/32 Type IV PF 10/17; NP 8/19; DF 15/17; Type V PF 9/15; NP 13/22; DF 16/20

“After manipulation of a Colles’ fracture, immobilization of the wrist in dorsiflexion would appear to provide better maintenance of reduction.”

Immobilization of wrist in palmar flexion had detrimental effect on hand function.

Wahlström 1982 RCT
No mention of sponsorship or COI.

4.0
N = 42 with extra articular fractures. Mean age 65 years.

Immobilization in pronation (N = 14) vs Supination (N = 12) vs Midway position (N = 16). Follow-up at 10 days and 1-4 months after reduction.

Five fractures had to be re-reduced, one from pronation, one from midway and three from supination group. Patients with redislocation ≥10° number pronation 2/14, midway 6/12, and supination 8/16. “The position of the forearm during immobilization is of importance for the degree of redislocation.”

Applicable to cast application rather than long-term functional results.
REDUCTION ANALGESIA

1. **Recommendation: Bier Block Analgesia for Manipulation of Acute Displaced Distal Forearm Fractures**
   
   Bier block analgesia is moderately recommended as a first-line technique for manipulation of acute displaced distal forearm fractures.
   
   **Strength of Evidence – Moderately Recommended, Evidence (B)**
   
   **Level of Confidence – Moderate**

2. **Recommendation: Hematoma Block Analgesia for Manipulation of Acute Displaced Distal Forearm Fractures**
   
   Hematoma block analgesia is recommended for manipulation of acute displaced distal forearm fractures.
   
   **Strength of Evidence – Recommended, Evidence (C)**
   
   **Level of Confidence – Low**

3. **Recommendation: Dynamic Reduction for Acute Distal Forearm Fractures**
   
   Dynamic reduction is recommended as an alternative technique for distal forearm fractures as it may result in less reduction pain than hematoma block, and may have a lower neurologic complication rate than a hematoma block.
   
   **Strength of Evidence – Recommended, Evidence (C)**
   
   **Level of Confidence – Low**

**Rationale for Recommendations**

There are three moderate-quality studies that support the use of Bier block (intravenous local anesthetic) over hematoma (local infiltration) block for pain control during manipulation and reduction of displaced Colles’ fracture.(1305-1307) In addition, those manipulated under Bier block were found to have better anatomic outcomes,(1306, 1307) lower remanipulation rates,(1306) and better grip strength at 6 months.(1307) Interestingly, medical staff may prefer hematoma infiltration over bier block based on perception of ease of technique, analgesia quality, and risk avoidance despite patient satisfaction and preference for Bier block.(1305) There are no quality studies comparing Bier block with any of the other techniques. Thus, Bier block is recommended as a first-line technique for achieving adequate analgesia and for potentially improving chance for better anatomic and functional outcome.

Dynamic reduction which does not require anesthesia block may hold slight advantage over hematoma block from the patient’s perspective, although there is one study of moderate quality available on the topic, wherein less severe pain was reported during dynamic reduction than those receiving local infiltration.(1293) There was no difference in reduction quality, and no longitudinal results were reported. The same author reported in a different paper, likely of the same study group, that hematoma infiltration resulted in higher subjective paresthesia or mild deficit in 2-point discrimination at 5 weeks and 1 year compared with the dynamic traction group.(1294) Hematoma infiltration provided lower pain scores during reduction and quicker onset of analgesia than patients receiving IV pentazocine (Talwin®) and diazepam (Valium®).(1308) Finally, in one moderate-quality study, hematoma block showed no difference with cubital block, and both were judged to be substandard.(1309)

**Evidence for Reduction Analgesia for Displaced Distal Forearm Fractures**

There is 1 high-(1308) and 6 moderate-quality(1293, 1294, 1305-1307, 1309) RCTs incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: reduction analgesia, bier block, analgesia, hematoma block analgesia, dynamic reduction, distal, forearm, radial, radius fractures, bone fractures, Colles’ fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, random; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 5 articles
in PubMed, 11 in Scopus, 0 in CINAHL, 1 in Cochrane Library, 100 in Google Scholar, and 3 from other sources. We considered for inclusion 3 from PubMed, 1 from Scopus, 0 from CINAHL, Cochrane Library, and from Google Scholar, and 4 from other sources. Of the 8 articles considered for inclusion, 8 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Singh 1992 RCT</td>
<td>9.0</td>
<td>N = 66 (46 male and 20 female) with Colles’ fracture. Mean age groups A and B: 36±16 and 39±15.</td>
<td>Group A, received 30mg pentazocine with 5mg diazepam (N = 33) vs 20cc or 20cc of 1.5% Xylocaine (N = 33). Follow-up for 15 hours.</td>
<td>“The pain scores during reduction in the local anesthetic group were markedly lower (mean 2.7, median 1.8) than the scores in the sedation group (mean 8.2, median 8.7), p &lt; 0.001.”</td>
<td>“Hematoma block by local anesthetic is a safe and effective alternative to sedation in reduction of Colles fracture.”</td>
<td>Patients receiving local anesthesia had lower pain and quicker reductions than those receiving sedation.</td>
</tr>
<tr>
<td>Kendall 1997 RCT</td>
<td>5.5</td>
<td>N = 142 (17 male and 125 female) with Colles’ fracture. Mean age for Bier’s block and Haematoma groups: 65 and 6 years.</td>
<td>Bier’s block (N = 72) vs Haematoma block with either alkalinized non-alkalinized local anesthetic (N = 70). Follow-up not specified.</td>
<td>Mean pain scores Bier’s block/hematoma: administration of anesthetic 2.8/5.3 p &lt;0.001; manipulation of fracture 1.5/3.0 p &lt;0.01. Alkalinized vs. non-alkalinized hematoma block alkalinized/non-alkalinized: median pain score on administration 4.4/5.9 p = 0.08; median pain score on manipulation 3.5/3.0 p = NS. More remanipulations in hematoma block (17/70) than Bier’s block (4/72) (p = 0.003).</td>
<td>“Bier’s block is superior to hematoma block in terms of efficacy, radiological result, and remanipulation rate.”</td>
<td>Trend to decreased pain with alkalinized v non-alkalinized group but did not reach significance.</td>
</tr>
<tr>
<td>Kongsholm J Orthop Trauma 1987 RCT</td>
<td>5.5</td>
<td>N = 116 (6 male and 56 female) with acute displaced Colles’ fracture. Mean age 61.6 years.</td>
<td>Group A, dynamic reduction device with no anesthesia (N = 62) vs Group B, 8 to 10ml of 1% lidocaine with plaster cast (N = 54). Follow-up for 1 year.</td>
<td>2/62 in Group A displayed symptoms and signs of neurological impairment at 5 weeks vs. 11/54 in Group B, p &lt;0.01. 1 year follow up resulted in figures of 4/62 and 8/54 with p &lt;0.05.</td>
<td>“The dynamic reduction technique without local anesthesia results in a significantly lower frequency of neurological complication than manual reduction after injection of local anesthetic into the fracture hematoma.” Authors speculate “that the mechanism for such nerve damage is scarring and fibrosis in the vicinity of the nerves, which is secondary to elevated pressure in the tissues probably caused by the increased volume load due to the injection.”</td>
<td>Neurologic complications included subjective paresthesia, positive Tinel’s sign or 2 point discrimination &gt; 4mm. Authors note that “nerve damage was mild and in no case in either group did it lead to surgical neurolysis.”</td>
</tr>
<tr>
<td>Fathi 2015 RCT</td>
<td>5.0</td>
<td>N = 143 (76 male and 67 female) with distal radial fracture. Mean procedural sedation and analgesia or PSA group, received 0.05 mg/kg</td>
<td>Pain numeric rating scale before / 5 / 10 / and 15 minutes after treatment:</td>
<td>“Ultrasound guided haematoma block may be a safe and effective”</td>
<td>Data suggest comparable efficacy in both groups but</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>Sponsorship/COI</td>
<td>N</td>
<td>Characteristic</td>
<td>Intervention</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>--------</td>
<td>-----------------</td>
<td>---</td>
<td>---------------</td>
<td>--------------</td>
</tr>
<tr>
<td>No mention of sponsorship. No COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haasio 1990</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abbasza-degan Acta Orthop Scand 1990;61:348-9</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kongsholm Injury 1987</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Sponsorship/COI</th>
<th>N</th>
<th>Characteristic</th>
<th>Intervention</th>
<th>Outcome</th>
<th>Conclusion</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>No mention of sponsorship. No COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haasio 1990</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abbasza-degan Acta Orthop Scand 1990;61:348-9</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kongsholm Injury 1987</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cobb 1985 RCT</td>
<td>4.0</td>
<td>N = 100 with Colles' fractures. Aged over 15 years.</td>
<td>Bier's block, fracture was manipulated 10 minutes after injection (N = 44) vs Local infiltration, fracture was manipulated 10 minutes after injection (N = 56). Follow-up for 20 minutes.</td>
<td>“Pain scores during manipulation were higher for patients receiving local infiltration vs. bier block (mean 5.53/10 vs. 3.67/10, P, 0.003). No difference was noticed in the period of postoperative painlessness between the groups: Bier's block 3-7 (3-0) hours; local infiltration 4-0 (3-0) hours.”</td>
<td>Despite findings, author states “For patients with fresh Colles' fracture local anesthetic infiltration was more popular among accident service staff (table), giving satisfactory anesthesia, being simpler and quicker to perform, and avoiding risks of a large intravenous does of local anesthetic agent reaching the general circulation.”</td>
<td>Paper highlights difference in staff perception (thought local was better) vs. patients own perception (preferred Bier's).</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Physical Methods/Rehabilitation

ELECTROMAGNETIC FIELDS

Recommendation: Use of Low Frequency Electromagnetic Fields to Stimulate Bone Healing of Distal Radial Fractures

The use of extremely low frequency (1-1000 Hz) electromagnetic field therapy to stimulate bone healing in patients with non-displaced fractures is not recommended.

Strength of Evidence – Not Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
There is one moderate-quality study of extremely low frequency (ELF) electromagnetic field (EMF) therapy, which is hypothesized to stimulate bone healing as measured by scintigraphy. The study found early increased bone activity in the first two weeks vs. control, but the differences disappeared after Week 2. In a subset of patients with displaced fractures that were re-displaced during the study, EMF of ELF resulted in higher scintimetric scores; however, the clinical significance of this finding is unknown.

Evidence for the Use of Electromagnetic Fields for Distal Radial Fractures
There are 3 moderate-quality RCTs incorporated with this analysis.(1389-1391) (Cheing 05; Lazovic 12)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Electromagnetic field therapy, electromagnetic therapy, PEMFT, Pulsed electromagnetic field therapy, magnetic therapy, magnet therapy, distal, Forearm, radial, Radius Fractures, bone Fractures, Colles’ Fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 60 in Scopus, 0 in CINAHL, 14 in Cochrane Library, 100 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 2 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 1 from other sources. Of the 3 articles considered for inclusion, 3 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wahlström 1984</td>
<td>5.5</td>
<td>N = 30</td>
<td>Electromagnetic fields of extremely low frequency (N = 15) – Received treatment vs. Control (N = 15) – Did not receive treatment</td>
<td>Scintimetric exam treated group/ control group: Week 1: 23.9±6.4/ 18.5±6.5 p &lt;0.05; Week 4: 44.6±13.6/ 41.6±15.0.</td>
<td>&quot;The clinical relevance of the results is not known, but one interpretation of the data is that the stimulation with EMF of ELF improves (accelerates) the early phase of fracture healing. The data warrant further investigation of fresh fracture treatment with this method.&quot;</td>
<td>Magnitude of differences disappeared at 4 weeks, thus importance of results unclear.</td>
</tr>
</tbody>
</table>

No mention of sponsorship or COI.
<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Patient Population</th>
<th>Intervention</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cheing 2005RTC</td>
<td>83 patients diagnosed with stable distal radius fracture(s). Mean age = 63.1</td>
<td>Group A (N=23) Ice plus PEMF 30 min of ice plus PEMF vs Group B (N=22) Ice plus sham PEMF 30 min of ice plus sham PEMF Vs Group C (N=22) PEMF. No ice. PEMF alone Vs Group D, Control (N=16) Sham PEMF. No ice. Sham PEMF alone.</td>
<td>All treatment were done for 5 consecutive days Visual analogue scale pain scores, volumetric measurements and ROM were measured on days 1, 3, and 5. VAS: The VAS score on day 1 was ranging from low to medium. On day 3, there was no significant drops between the groups. But the sham PEMF with no ice had the least amount of reduction. On day 5, the score for Ice plus PEMF was significantly higher than the other three groups. Volumetric Measurement: Day one, baseline, measurements were comparable between the groups. On day 3, the sham PEMF and no ice group decreased less than the others. Day 5 revealed that Ice plus PEMF was better than the PEMF/no ice and sham PEMF/ no ice group. Also the ice/sham PEMF group was better than the shame PEMF/no ice group. ROM: Flexion improved significantly in the two PEMF (ice/no ice) group compared to the sham PEMF on day 3. Day 5 yielded similar results, but the differences was not significant. Pronation was the exact opposite. The difference between day 1 and 3 were not significant. But the difference between day 3 and 5 were.</td>
<td>Data suggest pain was significantly reduced via VAS as well as ulnar ROM deviation with the addition of PEMF to ice or PEMF alone compared to sham groups.</td>
</tr>
<tr>
<td>Lazovic 2012RTC</td>
<td>60 women who sustained unilateral extra-articular displaced stable DRF Mean age PEMF = 67.90 ± 5.56</td>
<td>PEMF Group (N=30) PEMF therapy 5 days a week for 2 weeks Vs. Control Group (N=30) No therapy. Follow up at 2-3 days after removal of cast.</td>
<td>PEMF yielded better mean results for edema, pain, and function scores compared to the control. However, only the edema score was significant (p=0.000). PEMF resulted in significant values, when comparing ROM to</td>
<td>Some baseline differences between groups which could cause PEMF group to show worse outcomes. PEMF group was older. Data suggest</td>
</tr>
</tbody>
</table>
Mean age
Control = 64.50 ± 6.02
60 women
0 men

the control, for flexion (p=0.003), extension (p=0.009) and supination (0.004). Other ROM values were high in the PEMF group except for radial deviation.

PEMF for distal radius fracture beneficial for increased ROM and decreased edema post cast removal.

EXERCISE
Exercise is not generally indicated acutely. Some patients have deficits after casting or surgery and require exercises and rehabilitation.

1. **Recommendation: Education after Cast Removal for Acute Colles’ Fracture**
   Referral of select patients needing education after cast removal for acute Colles’ fracture is recommended.
   
   *Strength of Evidence – Recommended, Insufficient Evidence (I)*
   
   *Level of Confidence – Low*

2. **Recommendation: Use of Physical or Occupational Therapy after Cast Removal for Acute Colles’ Fracture for Patients with Functional Deficits Unable to Return to Work**
   Referral of patients with functional deficits or those unable to return to work for physical or occupational therapy after cast removal for acute Colles’ fracture is recommended.
   
   *Strength of Evidence – Recommended, Insufficient Evidence (I)*
   
   *Level of Confidence – Moderate*

3. **Recommendation: Routine Referral for Physical or Occupational Therapy after Cast Removal for Colles’ Fracture for Patients Able to Return to Work**
   Routine referral for physical or occupational therapy after cast removal for Colles’ fracture of otherwise healthy patients who are able to return to work is not recommended.
   
   *Strength of Evidence – Not Recommended, Evidence (C)*
   
   *Level of Confidence – Moderate*

**Rationale for Recommendations**

There are two moderate-quality studies on the effects of physical or occupational therapy to hasten functional recovery once the cast is removed in non-surgical patients. One study, despite lack of blinding in the control group and small sample size, showed no added benefit to the addition of occupational therapy to home exercise instructions by the provider. This finding was consistent regardless of fracture angulation and functional scores.(1311) Conversely, in another study also weakened by small sample size and lack of blinding, physical therapy (passive joint mobilization) was found more effective than no therapy in increasing wrist extension and grip strength in the immediate cast removal period. However, there were no long-term measures to determine the duration of benefit in the intervention group.(1312) One low-quality study(1313) and one case series(1314) also found no functional benefit for physical therapy.

A few appointments for educational purposes for select patients are recommended. The numbers of appointments are dependent on the degree of debility, with one or 2 educational appointments appropriate for mildly affected patients. Patients with severe debility or those unable to return to work may necessitate 8 to 12 appointments that particularly emphasize progressive strengthening exercises. Additionally, while routine use may be of limited benefit, those patients who have muscle weakness or other debilities may also derive benefit from therapy including self-training exercises, particularly if unable to return to work. Therefore, occupational or physical therapy is recommended for select patients.

*Evidence for the Use of Physical or Occupational Therapy for Colles’ Fracture*
There are 8 moderate-quality RCTs incorporated into this analysis.(1392, 1393, 1396-1401) (Wakefield 00; Kay 00; Filipova 15; Valdes 05; Magnus 13; Kay 08) There are 2 low-quality RCTs and one other study (1355, 1394, 1395) in Appendix 2.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Education, Cast removal, Colles' Fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 64 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: physical therapy, occupational therapy distal, Forearm, radial, Radius Fractures, bone Fractures, Colles' Fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 4 articles in PubMed, 5 in Scopus, 2 in CINAHL, 1 in Cochrane Library, 79 in Google Scholar, and 1 from other sources. We considered for inclusion 4 from PubMed, 2 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 6 articles considered for inclusion, 4 randomized trials and 2 systematic studies met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Exercise; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 3 articles in PubMed, 21 in Scopus, 0 in CINAHL, 1 in Cochrane Library, 146 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 2 articles considered for inclusion, 2 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year of Publication</th>
<th>Study Type</th>
<th>Conflict of Interest (COI) Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wakefield 2000</td>
<td>RCT</td>
<td>6.5</td>
<td>N = 96 patients over the age of 55 with a distal radius fracture treated with immobilization in plaster; mean age 72 (55-90). Gender (M:F); 9:87</td>
<td>Group 1 which was taught and given home exercises by a physiotherapist in a fracture clinic and referred to a course of physiotherapy (N = 49) vs Group 2 which was instructed in home exercises only (N = 47).</td>
<td>Mean difference (95% CI) for group 1 vs. group 2 at 6 months. Flexion/Extension: 12.2 (5.4 to 19.2), (p =0.001).</td>
<td>“Home exercises are adequate rehabilitation after uncomplicated fracture of the distal radius, and routine referral for a course of physiotherapy should be discouraged.”</td>
<td>Data suggest home exercises for uncomplicated fractures are beneficial.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Effect Size</td>
<td>Sample Characteristics</td>
<td>Intervention</td>
<td>Outcome</td>
<td>Comments</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>-------------</td>
<td>------------------------</td>
<td>--------------</td>
<td>---------</td>
<td>----------</td>
<td></td>
</tr>
<tr>
<td>Christensen 2001</td>
<td>RCT</td>
<td>No mention of COI or Sponsorship</td>
<td>5.0</td>
<td>N = 30 with distal radius colles' type fracture; mean age 66 years; Gender (M:F) 3:27</td>
<td>Home exercise instructions for shoulder, elbow, wrist and fingers with and without occupational therapy.</td>
<td>No statistical significance between groups in dorsal angulation, radial angulation, axial radial length, or functional scores.</td>
<td>“For non-surgically treated patients with a distal radius fracture only instructions are necessary.”</td>
</tr>
<tr>
<td>Kay 2000</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>4.5</td>
<td>N = 39 patients with fractures involving the distal radius, and after removal of pins and/or cast; mean age for non-mobilisation group 51.6, mobilisation group 54.7; Gender (M:F) 12:27</td>
<td>Non-mobilisation group received advice and home exercises from a physiotherapist. (N = 20) vs Mobilisation group received advice, home exercises, and a six week course of passive mobilisation. (N = 19)</td>
<td>Mean difference (95% CI) for non-mobilisation group vs mobilisation group at initial, three weeks, six weeks: Flexion : ( -0.6 , -5.0, -1.3), (p = 0.02).</td>
<td>All data collected comparing extension, flexion, radial deviation, ulnar deviation, pronation, supination, web space, thumb motion scale, and grip strength were not statistically significant.</td>
</tr>
<tr>
<td>Valdes 2015</td>
<td>RCT</td>
<td>No sponsorship or COI</td>
<td>4.5</td>
<td>N = 50 patients with DRF and underwent volar plate fixation; Mean age Therapy group (28-81) Non therapy (23-92); Gender (M:F) 8:42</td>
<td>Therapy group received instruction from a standard pictorial home exercise program (HEP) and therapy with certified hand therapists. 2visits/wk for 16 visits N = 26 Non therapy group received standard pictorial HEP N = 24</td>
<td>No statistically significant differences between scores of PRWHE, wrist/forearm motion, pain or grip strength between groups.</td>
<td>“Supervised clinic-based therapy is equally beneficial for patients without complications. Clinic-based therapy may be preferable for patients with noteworthy complications after a distal radius fracture with volar plate fixation.”</td>
</tr>
<tr>
<td>Filipova 2015</td>
<td>RCT</td>
<td>No COI or sponsorship</td>
<td>4.0</td>
<td>N = 61 patients who were treated conservatively for distal radius fracture; mean age 60 ; Gender (M:F) 14:47</td>
<td>Group A received 9 PT sessions consisting of 20min galvanic baths, and 30 min individual kinesiotherapy. (N = 31)</td>
<td>Rehabilitation outcomes p values for a two-way (Time and Therapy) mixed ANOVA</td>
<td>“The combined therapy resulted in a statistically significant increase of grip strength in comparison with isolated physical therapy in the period of 12–16 weeks after the fracture. This</td>
</tr>
</tbody>
</table>

Data suggest no difference between groups.
<table>
<thead>
<tr>
<th>Study</th>
<th>Methodology</th>
<th>N</th>
<th>Description</th>
<th>Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Watt 2000</td>
<td>RCT</td>
<td>4.0</td>
<td>18 patients with Colles' Fractures; mean age 74.4, Non physiotherapy group 77.3; Gender (M:F) 1:17</td>
<td>Physiotherapy vs. non-physiotherapy following cast removal. Follow up at six weeks after cast removal</td>
<td>Clinical significant increase in wrist extension and grip strength after 6 weeks physiotherapy (passive joint mobilization). “Routine referral of Colles’ fracture patients to physiotherapy following cast removal is beneficial.”</td>
</tr>
<tr>
<td>Kay 2008</td>
<td>RCT</td>
<td>6.5</td>
<td>56 patients with DRF managed with pins and/or a cast; Mean age Experimental group 55, control group 55.8</td>
<td>Experimental group received a physiotherapist directed program of advice and exercise. (N = 28) vs. Control group who did not receive any physiotherapy intervention. Follow up at three and six weeks</td>
<td>No significant difference found between groups comparing wrist extension, ROM or strength. “An advice and exercise program provided some benefits over no intervention for adults following distal radius fracture.”</td>
</tr>
<tr>
<td>Magnus 2013</td>
<td>RCT</td>
<td>4.0</td>
<td>51 women with unilateral DRF; Mean Age Training group 63.3 ± 10 Control group 62.7 ± 10.2</td>
<td>Training group received strength training in non-fractured arm during casting and through follow up and standard clinical rehabilitation (N = 27) vs Control group, received standard</td>
<td>Fracture hand strength Training vs control at 12wks (17.3± 7.4 kg vs 11.8 ± 5.8kg (p &lt; 0.017) ) No significant differences in strength at 9, 12 or 26 wks. Fractured hand ROM training vs control group at 12 weeks (100.5 ± 10.0”</td>
</tr>
</tbody>
</table>
Gender (M:F) 0:51  
clinical rehabilitation (N = 24).  
Follow-up at week 1, 3, 6, 9, 12, 26  
19.2 vs 80.2 ± 18.7 ( p < 0.017 )
Not significant differences in ROM at 9, 12, 26 weeks  
No significant differences in patient rated wrist questionnaires at week 9 or 26.

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>strategies after unilateral injuries.</td>
</tr>
</tbody>
</table>

### Surgery

1. **Recommendation: Closed Reduction or External Fixation for Severely Displaced Extra-articular Fractures, Comminuted, or Displaced Intra-articular Fractures of the Distal Forearm**  
   Closed reduction or external fixation is moderately recommended for treatment of severely displaced extra-articular fractures, and for comminuted, displaced intra-articular fractures of the distal forearm.  
   
   *Strength of Evidence – Moderately Recommended, Evidence (B)*  
   *Level of Confidence – High*

2. **Recommendation: Cast Immobilization or External Fixation for Moderately Displaced Extra-articular Fractures, Non-comminuted or Non-displaced Intra-articular Fractures of the Distal Forearm**  
   Cast immobilization is moderately recommended for treatment of extra-articular fractures or distal forearm fractures that include moderately displaced extra-articular fractures, non-comminuted or non-displaced intra-articular fractures. External fixation is moderately recommended as a second option for fractures that fail reduction while immobilized.  
   
   *Strength of Evidence – Moderately Recommended, Evidence (B)*  
   *Level of Confidence – Moderate*

3. **Recommendation: Medullary Pinning As an Alternative to External Fixation**  
   Medullary pinning (k-wire or intramedullary fixation techniques) is recommended as an alternative to external fixation.  
   
   *Strength of Evidence – Recommended, Evidence (C)*  
   *Level of Confidence – Low*

4. **Recommendation: Bone Cement As an Alternative to External Fixation**  
   Remodellable bone cement (injected or open reduction) is recommended as an effective alternative to external fixation and casting.  
   
   *Strength of Evidence – Recommended, Evidence (C)*  
   *Level of Confidence – Low*

5. **Recommendation: Open Reduction and Internal Fixation Via Dorsal or Volar Plating**  
   Open reduction and internal fixation by either dorsal or volar plating is recommended if fracture remains unstable by other treatment methods. There is no clear evidence of a preferential approach.  
   
   *Strength of Evidence – Recommended, Insufficient Evidence (I)*  
   *Level of Confidence – Low*

   There is no recommendation for or against TFCC repair associated with distal radial fractures.  
   
   *Strength of Evidence – No Recommendation, Insufficient Evidence (I)*  
   *Level of Confidence – Low*

Rationale for Recommendations
Most comparative studies for surgical intervention of distal radial fractures includes external fixation using various named external fixators. In consideration of the topic, all studies using external fixators are considered as one particular treatment group, as few comparisons are made between individual types or brands of fixators. There are several moderate-quality studies available for this treatment. Overall, the available data is weakened by studies with small sample sizes, a lack of consistency in fracture types included in each study, and inconsistency in reported final outcomes measures (i.e., functional, radiographic) and duration of follow-up. Extra-articular fractures or distal forearm fractures that include non-displaced intra-articular fractures can be treated initially with external immobilization, using external fixation as a second option for fractures that fail reduction while immobilized. External fixation likely does not provide improved functional results in the elderly.

In consideration of support for external fixation, 5 moderate-quality studies that included either mixed fracture types or were limited to extra-articular, non comminuted and non-displaced intra-articular fracture types, generally showed equivocal or non-statistically significant positive trends in radiographic or functional outcomes when compared with casting.(1315-1319) Another moderate-quality study compared plaster cast to external fixation and found that patients with plaster casts showed significantly higher Maximum Voluntary Contraction than patients with primary external fixation on day immobilization device was removed until between 18 weeks and 1 year when the groups equalized.(1281) Therefore, there is evidence supporting non-operative treatment for these fractures using non-invasive immobilization techniques. However, for the more severe fractures, including comminuted extra-articular and displaced comminuted intra-articular types, there are 6 moderate-quality studies that support improved clinical outcomes from external fixation over casting.(1320-1325) Thus, there is strong evidence to support more invasive immobilization techniques for these more severe injuries. One exception to this may be for those over age 65, as there is one moderate-quality study that showed no difference in functional outcomes despite worse anatomical outcomes, suggesting the final anatomical outcome is less important in this age group.(1326) As there is only one study supporting this, there is insufficient evidence to make recommendation for or against greater emphasis on non-operative treatment in older populations.

Two moderate-quality comparison studies of external fixation with medullary pinning (percutaneous) resulted in mostly equivocal studies for radiographic and clinical outcomes measures,(1327, 1328) although both authors felt there were financial and post-operative care advantages with pinning over external fixation. One moderate-quality study, weakened by co-intervention differences, demonstrated improved outcomes with combined external fixation and additional k-wire fixation for displaced intra-articular fractures.(1323) If pinning is selected, there does not seem to be any difference in technique comparing Kapandji and Willinegger procedures,(1329) nor in the length of post-operative cast immobilization comparing 1 vs. 6 weeks.(1330) An alternative method for the treatment of distal radial fracture includes the use of bone cement. There are five moderate-quality studies found, although one author reported on the same study population in two different papers. The injection of remodellable bone cement, or the open reduction and use of remodellable bone cement, was shown to provide improved anatomic and functional outcomes compared to casting(1331, 1332) and external fixation(1333, 1334) and reducing immobilization time.(1335) There is only one moderate-quality study on the repair of triangular fibrocartilage complex (TFCC) with distal radial fractures.(1336) In a small sample size study limited to Frykman II and VI, closed reduction and casting had equivocal results to surgical repair. However, this study was published in 1989, prior to more recent anatomic studies and case series reports on TFCC. Therefore, no recommendation is made for TFCC repair based on insufficient evidence.

There is no quality evidence for specific internal fixation techniques in comparison to external fixation or other immobilization techniques. However, there is one moderate-quality study of two internal fixation techniques, which recommends against the use of pi-plates, which were more difficult to match properly to distal radius, and resulted in worse wrist flexion and extension outcomes than from 1/4 tube plates.(1337) Thus, with insufficient evidence for comparison, there are no recommendations for internal fixation techniques.
Evidence for Surgery for Displaced Distal Forearm Fractures

There are 39 moderate-quality RCTs or prospective studies incorporated into this analysis. (1356, 1367, 1402-1437) (Rozental 09; Foldhazy 10; Grewal 05; Grewal 11; Karantana 13; Kreder 05; Cassidy 03; Jeyam 02; Krishnan 03; Leung 08; Wei 09; Atroshi 06; Arora 11; Abramo 09; Egol 08)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: **Bone Cement** / Distal Forearm Fractures & Colles’ Fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 5 articles in PubMed, 12 in Scopus, 2 in CINAHL, 0 in Cochrane Library, and 6037 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 3 from Google Scholar, and 0 from other sources. Of the 3 articles considered for inclusion, 2 randomized trials and 1 systematic studies met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: **Cast Immobilization** / Distal Forearm Fractures & Colles’ Fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 5 in Scopus, 1 in CINAHL, and 2 in Cochrane Library, 6558 from Google Scholar, and 2 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 2 from Google Scholar, and 2 from other sources. Of the 5 articles considered for inclusion, 5 randomized trials and 0 systematic studies met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: **Closed Reduction** / Distal Forearm Fractures & Colles’ Fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 10 in Scopus, 2 in CINAHL, and 4 in Cochrane Library, 15380 from Google Scholar. We considered for inclusion 0 from PubMed, 3 from Scopus, 0 from CINAHL, 2 from Cochrane Library, 8 from Google Scholar, and 0 from other sources. Of the 13 articles considered for inclusion, 6 randomized trials and 1 systematic studies met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: **Medullary Pinning** / Distal Forearm Fractures & Colles’ Fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, and 0 in Cochrane Library, 2175 from Google Scholar, and 5 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 2 from other sources. Of the 2 articles considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: **Open Reduction** / Distal Forearm Fractures, Colles’ Fracture; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 5 articles in PubMed, 6 in Scopus, 2 in CINAHL, and 2 in Cochrane Library, 5425 from Google Scholar, and 10 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 1 from CINAHL, 1 from Cochrane Library, 3 from Google Scholar, and 3 from other sources. Of the 9 articles considered for inclusion, 7 randomized trials and 2 systematic studies met the inclusion criteria.
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Triangular Fibrocartilage Complex Repair (TFCC) / Distal Forearm Fractures & Colles’ Fractures; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 968 from Google Scholar, and 0 in other sources. Zero articles met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kreder 2006 RCT</td>
<td>7.5</td>
<td>N = 113 (74 female/39 male) skeletally mature with distal radius fractures. Age 16-75 years.</td>
<td>Closed reduction casting (n = 59) vs. Closed reduction and external fixation (n = 54). Follow-up for 2 years.</td>
<td>No statistically significant differences in functional, clinical, or radiographic outcomes found; 19 patients in external fixator group had additional percutaneous pin fixation; 5 patients initially randomized to cast group actually received external fixations within 3 weeks of surgery (within 2 weeks of initiating cast treatment) because their fractures displaced or acceptable closed reduction could not be achieved (n = 5; 8.5%).</td>
<td>“For distal radius fractures with metaphysial displacement but with a congruous joint, there exists a trend for better functional, clinical, and radiographic outcomes when treated by immediate external fixation and optional K-wire fixation.”</td>
<td>Author notes to achieve statistically significant results, a sample of n = 600 would be necessary. “…simply not enough patients or resources to definitively answer this functional question.”</td>
</tr>
<tr>
<td>McQueen 1996 RCT</td>
<td>6.0</td>
<td>N = 120 (13 males/107 females) patients with unstable fractures of distal radius; Mean age 63 (16-86).</td>
<td>Closed re-reduction with forearm cast (Group 1) vs. Open reduction and bone grafting (Group 2) vs. Closed re-reduction and application of Pennig external fixator (Group 3) vs. Closed re-reduction and application of Pennig external fixator, but at three weeks the ball joint was released to allow wrist movement (Group 4). Follow up at 6 weeks and one year.</td>
<td>Mean dorsal angulation correction better in open reduction and grafting group (Group 2) vs. control and external fixation groups at 6 weeks and 1 year. Groups 3 and 4 better than control, but no statistical difference between fixation and fixation with early mobilization. Mean mass grip strength as a percentage of normal side showed sequential improvement, but no statistical analyses done. Carpal malalignment had a significant association with diminished recovery of strength of mass grip (p = 0.02), chuck grip (p = 0.02) and key grip (p = 0.05) after 1 year. Similar association with recovery of range of rotation at 3 months (p = 0.005), 6 months (p = 0.002) and 1 year (p = 0.01). After 1 year radial shortening had a significant negative association with recovery of chuck (p = 0.005), key (p = 0.01) and pinch (p = 0.001) grip strengths.</td>
<td>“Functional results at 6 weeks, 3 and 6 months, and at one year showed no difference between any of the four groups despite anatomical disparity. The main influence on final outcome was carpal malalignment which had statistically significant negative effect on function.”</td>
<td>Despite differences in the final anatomical appearance of the distal radius, the incidence of carpal malalignment was similar in all groups. Authors state correction of palmar tilt is most important to reduce carpal malalignment. The four techniques in the study are equivocal in this study for improving palmar tilt.</td>
</tr>
<tr>
<td>Study</td>
<td>Score</td>
<td>Design</td>
<td>Patients</td>
<td>Intervention</td>
<td>Outcome</td>
<td>Conclusion</td>
</tr>
<tr>
<td>-------</td>
<td>-------</td>
<td>--------</td>
<td>----------</td>
<td>-------------</td>
<td>---------</td>
<td>------------</td>
</tr>
<tr>
<td>Jenkins 1987</td>
<td>6.0</td>
<td>RCT</td>
<td>N = 58 (no mention of gender) patients with a displaced Colles' fracture; no mention of mean age.</td>
<td>Forearm plaster vs. external fixator in patients with displaced Colles' fractures. Follow-up at 4, 8, and 16 weeks.</td>
<td>Mean loss of position significantly worse for plaster vs. fixator in dorsal angle 10.5° vs. 0.1° (p &lt;0.01), radial angle 6.5° vs. 0.7° (p &lt;0.01), radial length 3.7 vs. 0.3° (p &lt;0.01). Using a positional grading scale to rate changes between post-manipulation and union, 22 of 24 in plaster group had good or excellent post-manipulation, falling to 12 of 24 at union. In fixator group, no decrease, as all 30/32 with good or excellent post-manipulation patients remained at 30 of 32.</td>
<td>&quot;The external fixator proved more effective at holding the manipulated position, and the radiological loss of position during fracture union was minimal compared with that seen in patients treated in plaster.&quot;</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>External fixation is more effective than plaster in radiological scoring. There were no measurements of function in this study.</td>
</tr>
<tr>
<td>Abbaszadeh 1990 Acta Orthop Scand 1990;61:528-30</td>
<td>5.0</td>
<td>RCT</td>
<td>N = 47 (11 males/36 females) with severely displaced Colles' fractures types 3 and 4; Mean age 63 (22-75)</td>
<td>Prospective 1-year study of plaster cast or primary external fixation. Follow up at 4, 8, 12, 24 weeks, and 1 year.</td>
<td>Follow-up according to pain and subjective function: Pain cast/fixation (VAS 0-10); 8 weeks 3/2 (p = 0.04); 12 weeks 2/1 (p = 0.1); 24 weeks 2/0.5 (p = 0.009); 1 year 1/0 (p = 0.0002. Function cast/fixation (VAS 0-10); 8 weeks 5/7 (p = 0.1); 12 weeks 7/7 (p = 0.7); 24 weeks 8/9 (p = 0.1); 1 year 9/10 (p = 0.02). Functional outcome excellent or good/total: Plaster 12/19; Fixation 19/22.</td>
<td>&quot;Primary external fixation for severely malpositioned Colles' fractures might lead to a better radiographic and functional end result than conventional plaster-cast treatment.&quot;</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>5 fractures in plaster cast group redislocated after 11 days, and re-reduction and external fixation were required, with 3/5 reporting good or excellent results.</td>
</tr>
<tr>
<td>Merchan 1992</td>
<td>5.0</td>
<td>RCT</td>
<td>N = 70 with comminuted intra-articular fractures of the distal radius of types III to VIII;</td>
<td>Closed reduction and forearm plaster vs. application of a Clyburn dynamic external fixator.</td>
<td>&quot;Significant loss of position occurred in 27 (77%) of the plaster group at the 7-day examination… Patients stabilized with an external fixator had maintained their reduced position.&quot; In fixator group, 54.3% had excellent reduction, 34.3% good reduction, 8.7% fair reduction and 2.7 poor reduction compared to plaster group where 37.2% had excellent reduction, 17.2% had good reduction, 34.2% had fair reduction, and 11.4% had poor reduction. 4 occurrences of pin tract infection were found, however they were superficial and responded to treatment by cleansing and antibiotics. 3 encounters of pin loosening occurred. Reflex</td>
<td>&quot;It does appear that a good anatomic position combined with early rehabilitation of wrist function produces very favorable functional results in patients under 45 years of age.&quot;</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Study labeled as double blind; however, only an independent assessor could be blinded which was not well described.</td>
</tr>
<tr>
<td>Study</td>
<td>Rating</td>
<td>N =</td>
<td>Summary</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>-----</td>
<td>---------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stein 1990 RCT</td>
<td>5.0</td>
<td>126 with distal radius fracture; Mean age Group 1: 56 (20-89), Group 2: 54 (22-79).</td>
<td>Fixation with above-the-elbow cast immobilization vs. external fixation; Follow up at 1, 2, 4, and 6 weeks, then again at 6 months and 4 years. Patients categorized on 4 grade severity scale based on deformity, dorsal angulation, and shortening. Garland and Werley objective and subjective results showed 71 of 88 Grade I, II cast group had excellent or good results. In types II, IV fracture, external fixation had better scores 36 of 40 vs. 15 of 22 (p &lt;0.001).</td>
<td>Extraarticular fractures of the distal radius should be treated with cast immobilization. Comminuted intraarticular fractures of the distal radius should be treated with external fixation, which maintains accurate anatomic position until solid fracture healing is achieved.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pring 1988 RCT</td>
<td>4.5</td>
<td>75 (14 males/61 females) patients with Colles’ fractures; Mean age Group 1: 59.3, Group 2: 64.</td>
<td>Forearm cast alone vs. bipolar fixation of displaced fracture. Mean percentage grip cast/Bipolar: 7 weeks 28.5/21.6, 12 weeks 46.2/48.5, 6 months 63.8/67.6. “Nine fractures treated with plaster alone redisplaced and required manipulation. No patient initially treated with bipolar fixation required remanipulation. Functional results at 6 months did not reach statistical significance.”</td>
<td>A good final position (functional position) is desirable, even in the elderly; that bipolar fixation provides a method of achieving this, and that it is applicable to all but open fractures of the distal radius.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lagerström Scand J Rehabil Med 1999;31(1):55-62 RCT</td>
<td>4.5</td>
<td>33</td>
<td>Plaster cast (P-group) vs. external fixation using AO External Fixator® (E-group) vs. secondary fixator group (PE Group). Differences between uninjured and injured sides; P-group vs. E-group in Maximum Voluntary Contraction (MVC) (Newtons) (Higher difference is weaker); 6 weeks: 190.7*** SD = 49.0 vs. 206.7 *** SD = 77.5; 10 weeks: 126.4** SD = 48.8 vs. 155.6** SD = 59.6; 52 weeks: 32.6 SD = 38.1 vs. 34.2* SD = 35.0. *p&lt;0.05; **p&lt;0.01; ***p&lt;0.001.</td>
<td>For injured side patients with plaster casts showed significantly higher MVC (stronger) than patients with primary external fixation on day immobilization device removed until between 18 weeks and 1 year when groups equalized. Patients that failed casting and had external fixation had slower recovery trends.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jenkins 1988 RCT</td>
<td>4.0</td>
<td>106 (males/ females) who had sustained a Colles’ fracture sufficiently displaced to require</td>
<td>Forearm plaster vs. external fixator in patients (AO/ASIF minifixator) under age 60. Follow up at baseline, 1 week, 4 weeks, 3 months. Comparison of excellent and good outcomes/total: Subjective: external fixator 44/59, plaster 32/41. Objective external fixator 57/59, objective 40/41. Fixator group had much greater proportion of excellent than the plaster group. At 12 months, the plaster group had “The wrist’s immobilization does nothing to retard its early recovery. External fixation of these fractures is indicated solely for the purpose of improving long-term function by virtue of the improved anatomy that the treatment affords, and for study accounted for grip strength in dominant vs. non-dominant contralateral comparisons.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Copyright© 2016 Reed Group, Ltd.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Gender</th>
<th>Age</th>
<th>Treatment</th>
<th>Outcomes</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Howard 1989</td>
<td>RCT</td>
<td>4.0</td>
<td>N = 50 (No mention of Gender)</td>
<td>patients with severely displaced comminuted Colles’ Fractures; No mention of Mean Age.</td>
<td>Plaster with fracture manipulated under Bier’s block and supported by molded below-elbow plaster backslab vs. external fixation with 2 pairs self-tapping 2.0mm Hoffman pins inserted into radius, proximal to line of crossing of radial nerve.</td>
<td>Significantly reduced grip strength than externally fixated group (93.9% SD±9.4% vs. 84.1% ± 19.6 (p = 0.05).</td>
<td>for this reason methods of treatment that permit early wrist mobilization at the possible expense of the anatomical position are not justified.”</td>
</tr>
<tr>
<td>Young 2003</td>
<td>Prospective Study</td>
<td>4.0</td>
<td>N = 125 (28 males/97 females) with dorsally angulated fractures of the distal radius; Median Age Group 1: 54 (21-73). Group 2: 60 (24-75).</td>
<td>Group 1: primary bridging external fixation vs. Group 2: manipulation of the fracture with dorsal plaster slab converted to below-elbow plaster cast at 1 week</td>
<td>At 7-year follow-up, 17 died, 22 lost to follow-up, leaving 86. “There were no difference between groups for ranges of flexion, extension, pronation, supination and ulnar and radial deviation or grip strength.” Gartland and Werley scores similar with 34/36 of external fixation group and 47/49 of casting group reporting excellent or good scores. Residual wrist pain low with no differences between groups. Patients showing arthritic changes ext. fix n = 11/36, cast n = 9/49 not significant. Incidence of 14% reported for occurrence of radiological post-traumatic arthritis following intra-articular fractures.</td>
<td>“External fixation produces significantly better anatomical results than plaster in severely displaced comminuted Colles’ fractures and a significant improvement in function.”</td>
<td></td>
</tr>
<tr>
<td>Roumen 1991</td>
<td>RCT</td>
<td>4.0</td>
<td>N = 101 (8 males/93 females) with displaced Colles’ fracture; Mean Age 70.1</td>
<td>External fixator or conventional cast treatment (control) in patients that failed manipulation and Elderly patients with displaced Colles’ fractures treated with initial reduction and plaster backslab. At Week 1 and 2, patients with dorsal angulation &gt;10° or radial shortening &gt;5mm re-manipulated</td>
<td>“Radiographic result after distal radial fracture is significantly better if patients are treated by external fixation rather than by plaster immobilization. However, after 7 years, the outcome measures that the patient notices, such as range of movement and function, are no different between the two treatment methods.”</td>
<td>High dropout rate at long-term follow-up.</td>
<td></td>
</tr>
</tbody>
</table>

No correlation between anatomic and functional outcomes in elderly patients.
<table>
<thead>
<tr>
<th>Study Reference</th>
<th>Methodology</th>
<th>Design</th>
<th>Duration</th>
<th>Population</th>
<th>Intervention</th>
<th>Results</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allain 1999</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>7.0</td>
<td>N = 60 (45 female/15 male) with dorsally displaced extra-articular or non-comminuted intra-articular fractures of distal radius after trans-styloid K-wire fixation. Mean age 75 (21-83).</td>
<td>Postoperative immobilization for 1 (Group 1) or 6 (Group 2) weeks.</td>
<td>Patients followed at 1-year post-op. One reflex sympathetic dystrophy in Group 1, none in Group 2. Ulnar deviation statistically significant (p = 0.03) after early mobilization (mean difference between normal and impaired wrist). No significant differences in grip strength, (25 kg in Group 1 and 21 kg in Group 20, sick leave, functional discomfort, or outcome satisfaction.</td>
<td>“Addition of plaster cast immobilization of wrist after trans-styloid fixation with two K-wires, in Colles’ fractures may not be necessary if styloid fragment large enough to allow good K-wire fixation, as well as if fracture does not consist of more than 2 articular fragments.”</td>
</tr>
<tr>
<td>Strohm 2004</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>4.0</td>
<td>N = 100 (15 males/85 females) patients with Colles-type fracture of distal radius; Mean Age 65 (29-92)</td>
<td>Kirschner wire osteosynthesis via Kapandji procedure vs. Willenegger procedure.</td>
<td>Martini scores; Kapandji vs. Willenegger. Average 4 (range, 16-38 points) vs. 28 (range, 11-36 points) (p &lt;0.005). Difference in the modified Martini score between the treatment group was found for type-A2 (p = 1.004) and A3 (p = 0.007) fractures but not for type-C1 fractures (p = 0.6).</td>
<td>Conventional Kirschner wire fixation remains good method of osteosynthesis for treating displaced fractures of distal part of radius. “We found both the functional and radiographic outcomes of the Kapandji method to be significantly better than those of the Willenegger technique.”</td>
</tr>
<tr>
<td>Kapoor 2000</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>4.0</td>
<td>N = 90 (No mention of Gender) adult cases of acute displaced intra-articular fractures of the distal end of the forearm</td>
<td>Closed reduction and plaster immobilization vs. external fixation (Roger and Anderson type) vs. Open reduction and external fixation</td>
<td>Follow-up for 4 years. Final functional assessment (%); Plaster vs. Fixator vs. Open reduction: Good and excellent 43 vs. 80 vs. 63. Fair and poor 57 vs. 20 vs. 37. Average loss of arc with plaster 37° in comparison with 19° by external fixator. Average grip strength (in comparison with normal side) in “Displaced severely comminuted intra-articular fractures should be treated with an external fixator.”</td>
<td>Study intervention is different for fixation vs. internal fixation related to mobilization and physiotherapy. Follow-up times differ not clearly available.</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Age</td>
<td>Fracture Type</td>
<td>Fixation Method</td>
<td>Fixation Time</td>
<td>Outcome Measures</td>
<td>Observations</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>-----</td>
<td>----------------</td>
<td>------------------------</td>
<td>----------------------------------</td>
<td>---------------</td>
<td>--------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Ludvigsen 1997 RCT</td>
<td>6.0</td>
<td>N = 60 (7 males/53 females) with Colles’ Fracture type Older 3; Mean Age Group 1: 61 (31-80). Group 2: 58 (31-89).</td>
<td>External fixation vs. percutaneous pinning.</td>
<td>Patients immobilized for 6 weeks; outcome assessed after 6 months. Groups showed similar results with respect to radiographic parameters and function. All fractures healed and no difference in complication rate was observed.</td>
<td>Most unstable distal radial fractures, classified as Older’s type 3 and 4, can be treated with percutaneous pinning and a plaster cast, which is simpler and cheaper than external fixation.</td>
<td>With equivocal results, author justification for conclusion is based on other studies that loss of reduction may occur if external fixator is removed before 8 weeks, as radial shortening occurring during this time may result in loss of reduction.</td>
<td></td>
</tr>
<tr>
<td>Pritchett 1995 RCT</td>
<td>4.5</td>
<td>N = 100 (45 males/55 females) with distal radial fractures; Median Age Group 1: 65.3 (34-79). Group 2: 66.7 (37-83).</td>
<td>External fixation vs. medullary pinning.</td>
<td>Excellent or good outcome/total: external fixation 42/50, medullary pinning 48/50. Loss of ROM and grip strength slight and not significantly different between groups. Treatment outcomes of mean operating time, office visit numbers, use of more than 1 prescription drug, device removal, bathing and dressing problems, and other operation all favored medullary pinning.</td>
<td>“The two most important measures of outcome, patients complaints and cost, were significantly lower with pinning than with external fixation and we now believe that medullary fixation is the treatment of choice for these fractures.”</td>
<td>Non-clinical outcomes favor medullary pinning.</td>
<td></td>
</tr>
<tr>
<td>Hahnloser 1999 RCT</td>
<td>6.0</td>
<td>N = 46 (11 males/35 females) with unstable comminuted fracture of distal radius; Mean Age Group 1: 53.8 (17-77). Group 2: 58.2 (23-77).</td>
<td>Internal fixation via two 1/4 tube plates vs. [pi]-plate.</td>
<td>43% of [pi]-plates were too large and 19% could not be matched properly to distal radius. Range of wrist motion of the operative wrist expressed in percentage of the normal contralateral side: [pi]-plaster/tube plates Flexion 68 (±26 SD)/ 85 (±15); Extension 67 (±23)/ 86 (±12).</td>
<td>“With open reduction, cancellous bone grafting, and internal plate fixation in comminuted distal radial fractures, excellent results can be achieved. In our experience, we cannot recommend the [pi]-plate in its current shape and prefer to stabilize distal radius fractures and dorsal fragment dislocations with two 1/4 tube plates.”</td>
<td>Recommendation against pi-plate for internal fixation.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Method</td>
<td>N</td>
<td>Sex</td>
<td>Diagnosis</td>
<td>Comparison</td>
<td>Results</td>
<td>Notes</td>
</tr>
<tr>
<td>------------------------------</td>
<td>--------</td>
<td>---</td>
<td>------------</td>
<td>------------------------------</td>
<td>---------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Ekenstam 1989</td>
<td>RCT</td>
<td>5.0</td>
<td>41 males/31 females</td>
<td>Lidström Group IIIa-c or Frykman Groups II and VI</td>
<td>Triangular ligament was repaired after closed reduction (Group A) vs. closed manipulation and above-elbow cast (Group B).</td>
<td>Clinical examination results controls/group B/group A mean (SD): Strength 60(22)/58 (18)/59(25); Flexion 68(11)/58(11)/57(15). No difference for any part of clinical exam for 2 treatment methods.</td>
<td>“Repair of the ruptured triangular ligament in extraarticular fractures of the distal radius is not better than conventional treatment.” Dropout rate unclear. Randomization and baseline comparability not clear.</td>
</tr>
<tr>
<td>Schmalholz 1989</td>
<td>RCT</td>
<td>6.0</td>
<td>47 males/47 females</td>
<td>Frykman Types I and II</td>
<td>Bone cement (methylmethacrylate) (Group 1) vs. plaster cast (Group 2).</td>
<td>21/24 patients in Group 1 and 10/23 in Group 2 recovered full dorsiflexion; 8 in Group 1 and 1 in Group 2 regained full strength. Wrist appearance satisfactory for all in Group 1/none in Group 2 at 8 weeks. Group 1 function excellent in 6, good in 17, and fair in 1; Group 2 saw good in 2, fair in 12, and poor in 9. (p &lt;0.001).</td>
<td>“The operated on group were better with regard to all objectively measurable characteristics; all operated on fractures had healed radiographically, and the cement was surrounded by cortical bone.” Description of 2nd study sounds similar. Unclear if these 2 reports represent one trial with 3 arms split into 2 reports.</td>
</tr>
<tr>
<td>Schmalholz 1990</td>
<td>RCT</td>
<td>6.0</td>
<td>48 males/46 females</td>
<td>Redislocated Colles Fractures</td>
<td>Dorsal bone deficiency filled with bone cement (methylmethacrylate) (Group I) vs. external fixation (Group II).</td>
<td>Surgery on day 16 (median 16, range 14-18) in both groups. Group 1 (cement) had significant improvement in volar flexion, supination, pronation, and grip strength first 2-4 months post treatment. At 6 months all differences equalized. Group II, 24% had complications; none in Group I.</td>
<td>Final results equal in the 2 groups, but Group I improved earlier and had no complications. Open reduction and bone cement appears more effective than external fixation.</td>
</tr>
<tr>
<td>Sanchez-Sotelo 2000</td>
<td>RCT</td>
<td>5.0</td>
<td>110 males/97 females</td>
<td>Distal radius fractures</td>
<td>Remodellable bone cement (Norian SRS) and cast for 2 weeks vs. closed reduction and cast for 6 weeks.</td>
<td>Mean ranges of movement and mass grip strength as percentages of normal side. Norian SRS/Control: Extension 6 weeks 65.09±8.26/40.76±6.06 (p &lt;0.001); 1 year 95.7±3.2/90.1±3.4 (p &lt;0.01). Flexion 6 weeks 53.84±5.51/43.60±5.93 (p &lt;0.001); 1 year 86.2±3.41/77.8±4.2 (p</td>
<td>“The injection of a remodellable bone cement into the trabecular defect of fractures of the distal radius provides a better clinical and radiological result than conventional treatment.” Positive study for the use of remodellable bone cement over immobilization.</td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>N</td>
<td>Age</td>
<td>Fracture Type</td>
<td>Fixation Details</td>
<td>Outcome Measures</td>
<td>Findings</td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>---</td>
<td>-----</td>
<td>-------------</td>
<td>-----------------</td>
<td>-----------------</td>
<td>----------</td>
</tr>
<tr>
<td>Kopylov 2001</td>
<td>RCT</td>
<td>4.0</td>
<td>N = 23 (No mention of Gender) osteoporotic patients with distal radial fracture; Mean Age 68 in SRS group, 65 in external fixation.</td>
<td>External fixation vs. Norian SRS</td>
<td>Clinical findings reported in 1999 study. Stereometric findings are reported here. In all fractures there was a good correlation ($r^2 = 0.93, p = 0.0001$) between longitudinal radiostereometric analysis displacement from the first to last investigation.</td>
<td>Norian SRS/ Control: 3 months 45 (81.8%)/30 (54.5%); 12 months none 49 (89.1%)/38 (69.1%). Complication Norian SRS/ Control: Malunion 10 (18.2%)/23 (41.8%); Reflex sympathetic dystrophy 3 (5.4%)/4 (7.3%).</td>
<td></td>
</tr>
<tr>
<td>Kopylov 1999</td>
<td>RCT</td>
<td>4.0</td>
<td>N = 40 (36 men/4 women) with distal radial fractures; Mean age 67.5.</td>
<td>Stabilized with SRS injection and immobilized with cast for 2 weeks vs. externally fixed with Hoffman’s bar for 5 weeks</td>
<td>“SRS can be used in the treatment of unstable distal radial fractures. The more rapid recovery of grip strength and wrist mobility in the SRS group appears to be due to the shorter immobilization time.”</td>
<td>“The shorter immobilization time with SRS permitted earlier return of hand function. The question remains whether early mobilization by itself is enough to reach a good final result, even in the absence of fixation with SRS. That question is addressed in an ongoing study.”</td>
<td></td>
</tr>
<tr>
<td>Atroshi 2006</td>
<td>RCT</td>
<td>7.5</td>
<td>N=38 (7 males/31 females)</td>
<td>Group 1 (N=19) patients treated</td>
<td>No significantly different results in the mean DASH scores between both groups. No difference in</td>
<td>“The lack of a clear clinically relevant advantage”</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Data suggest similar efficacy between groups but non-</td>
<td></td>
</tr>
<tr>
<td>Wei 2009</td>
<td>Prospective Randomized trial</td>
<td>N=46 (13 males/33 females) patients with an unstable distal radial fracture; Mean Age Group 1: 58 ± 17.</td>
<td>7.0</td>
<td>Group 1 (N=22) patients treated with external fixation Vs Group 2 (N=12) patients treated with a radial column plate. Vs Group 3 (N=12) patients treated with a volar plate. Follow up at 10-14 days, 6 weeks, and 3, 6, and 12 months post-op.</td>
<td>Disabilities of the Arm, Shoulder and Hand (DASH) results, 6 weeks, group 3 vs group 1; 41 ± 23 vs 56 ± 19 (p=0.037). DASH 3 months group3 vs group 2 and vs group 1; 7 ± 5 vs 28 ± 17 (p=0.027), and 29 ± 18 (p=0.028). DASH at 1 year, group 3 vs 1 and 2; 4 ± 5 vs 18 ± 14 (p=0.025) and 18 ± 12 (p=0.056). Grip Strength (percentage value compared to uninjured side, at 6 months, group 1 vs 2; 75 ± 21 vs 53 ± 9 (p=0.042). Lateral pinch (% vs uninjured side); group 2 vs 3, at 3 months and 12 months. 66±14 vs 86±13 (p&lt;0.042), 73±8 vs 94±5 (p&lt;0.036). Range of motion; Extension, group 1 vs group 3 and 2 (degrees), 6 weeks; 10 vs 38 &amp; 32 (1 v 3 p=0.023), (1 v 2 p=0.032), respectively. Supination (degrees), group 1 vs 2 and 3, 6 weeks; 34 vs 57 and 55 (1 v 2 p=0.041), (1 vs 3 p=0.049) respectively. Radiographic Measurements; Radial inclination (deg), group 2 v 3, 6 weeks; 25.0±5.2 vs 21.1±7.0 (p=0.003). Radial Length (mm), group 1 vs 2,</td>
<td>In conclusion, this study provides new evidence supporting the trend toward fixation of distal radial fractures with locked volar plates.</td>
<td>Data suggest the use of a locked volar plate resulted in better patient reported outcomes at 3 months but at 6 months and 12 months, all 3 groups had good outcomes in terms of ROM, strength and radiographic alignment but the radial column plate group had significantly better radial inclination and length compared to other 2 groups.</td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Participants</td>
<td>Intervention</td>
<td>Follow-up</td>
<td>Summary</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>--------------</td>
<td>--------------</td>
<td>-----------</td>
<td>---------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Egol 2008</td>
<td>Prospective Randomized Trial</td>
<td>No sponsorship or COI.</td>
<td>N=88 (41 males/47 females) patients with a distal radius fracture that needed operative repair; Mean Age Group 1 49.9 (18-78). Group 2 52.2 (19-87).</td>
<td>Group 1 (N=38) patients that received external fixation and supplementary K-Wire fixation. Vs Group 2 (N=39) who were treated with volar plating.) Follow up at 2, 6 weeks, then at 3, 6, 12 months.</td>
<td>The mean DASH score in any of the intervals. For all parameters, as a percentage of the injured side, the range of movement was better in internally-fixed group; pronation (p&lt;0.001), supination (p=0.05), extension (p=0.05), radial deviation (p=0.002), reached statistical difference at 3 months. Similar complications.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rozental 2009</td>
<td>Prospective Randomized Trial</td>
<td>No sponsorship or COI.</td>
<td>N=45 () patients with an unstable fracture of the distal radius; Mean Age Group 1: 51 (19-77). Group 2: 52 (24-79).</td>
<td>Group 1 (N=23) patients treated with open reduction and internal fixation Vs Group 2 (N=22) patients treated with Closed reduction and percutaneous pinning. Follow up at 6, 9, 12 weeks, and 1 year.</td>
<td>Range of Motion Parameters (deg), group 1 vs 2, 6 weeks; Extension: 45±20 vs 16±13 (p&lt;0.01). Flexion: 50±12 vs 26±16 (p&lt;0.01). Supination: 79±21 vs 40±29 (p&lt;0.01). Pronation: 77±17 vs 63±26 (p=0.04). Ulnar Deviation: 27±10 vs 15±11 (p&lt;0.01). Radial Deviation: 15± vs 7±6 (p&lt;0.01). Grip Strength (% vs uninjured arm), group 1 vs 2, 6 weeks; 49.3±20.9 vs 25.6±30.1 (p&lt;0.01). Pinch Strength (% vs uninjured arm), group 1 vs 2, 6 weeks; 59.1±25.8 vs 38.8±27.0 (p&lt;0.01). DASH Score, group 1 vs 2, 6 weeks; 27±17 vs 53±28 (p&lt;0.01), 9 weeks; 17±17 vs 39±25 (p&lt;0.01), 12 weeks: 11±13 vs 26±23 (p=0.01). No significant difference between radiological outcome, return to</td>
<td>&quot;The present study confirms the hypothesis that volar plate fixation results in less functional disability in the first few months after treatment than does percutaneous pin fixation. At one year after the injury, we did not identify a difference between the treatment groups with regard to functional or radiographic outcomes.&quot;</td>
<td></td>
</tr>
</tbody>
</table>

Data suggest similar efficacy between groups but less re-operations were required in the external fixation group.
<table>
<thead>
<tr>
<th>Study</th>
<th>Patients</th>
<th>Group 1</th>
<th>Group 2</th>
<th>Findings</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grewal 2011</td>
<td>N=50 (12 males/38 females)</td>
<td>Group 1 (N=26)</td>
<td>Group 2 (N=24)</td>
<td>Group 1 scored 11 points lower on Patient-Rated Wrist Evaluation (PRWE) throughout whole study, except at 12 months (p=0.03). Group 1 vs 2 (specifically Volar locking plates) had significantly lower PRWE scores at baseline (p=0.03) and 6 weeks (p=0.06). No difference in radiological parameters, range of motion, grip strength, and complications.</td>
<td>&quot;[O]ur trial suggests that ORIF (Group 1) provides a short-term advantage over external fixation, but these differences do not persist over time. Our results are viewed with caution, given that the ORIF group also reported lower pain and disability at the initial preoperative assessment and because the trial had a small sample size.&quot;</td>
</tr>
<tr>
<td>Prospective RCT</td>
<td>Patients with fractures of the distal radius; Mean age Group 1: 58.0 ± 9.9. Group 2: 53.8 ± 11.7.</td>
<td>Patients with open reduction and internal fixation. Vs patients with external fixation procedures. Follow up at baseline, 6 weeks, and 3, 6, 12 months.</td>
<td></td>
<td></td>
<td>Data suggest ORIF group better than external fixation in short term but at 1 year, the results equalize among groups.</td>
</tr>
<tr>
<td>Krishnan 2003</td>
<td>N=60 (19 males/ 41 females)</td>
<td>Group 1 (N=30) pinned with a &quot;Delta&quot; frame and instructed to do wrist exercises Vs Group 2 (N=30) pinned in the &quot;Hoffman&quot; style and were not able to move wrist. Follow Up at 1, 6, 12, 26, 52 weeks.</td>
<td>No statistical difference between groups in extension, ulnar deviation, pronation and supination, grip strength, comparable complications in both groups except for rupture of extensor pollicis brevis tendon.. Flexion, Group 1 vs Group 2 at 6, 26, and 52 weeks median (range) in deg; 28 (10-60) vs 35 (10-90) (p=0.02), 45 (30-95) vs 55 (40-95) (p=0.008), 50 (25-100) vs 60 (45-100) (p=0.02). Radial deviation, group 1 vs Group 2, at 6 weeks; favored group 1 (p=0.002). Completing daily activities was better in group 2 than group 1 (p=0.034) at week 2.</td>
<td>In conclusion, this study demonstrated that the outcomes of patients with complex unstable intraarticular fractures of the distal radius are similar, regardless of whether they are treated with a static bridging external fixator or a dynamic non-bridging external fixator.</td>
<td>Data suggest comparable efficacy between groups.</td>
</tr>
<tr>
<td>Abramo 2009</td>
<td>N=50 (14 males/36 females)</td>
<td>Group 1 (N=25) who were treated with Open reduction and internal fixation Vs Group 2 (N=25) who were treated with closed</td>
<td>Grip strength (% vs uninjured arm), group 1 vs 2, 7 weeks; 47% vs 34% (p=0.01). Forearm rotation (deg), group 1 vs 2, 7 weeks; 129 vs 104 (p=0.006). Grip strength (% vs uninjured arm), group 1 vs 2, 1 year; 90 vs 78 (p=0.03). Forearm rotation (deg), group 1 vs 2, 7 weeks; 149 vs 136 (p=0.03). No difference in DASH scores between groups.</td>
<td><img src="" alt="Two methods compared will both give a good result with good DASH values, good grip strength, and good range of motion after a year. Overall, considering the subjective and objective results, the internal fixation group had better ROM, grip strength and fewer malunions than the external fixation group." /></td>
<td>At 1 year, data suggest internal fixation group had better ROM, grip strength and fewer malunions than external fixation group.</td>
</tr>
</tbody>
</table>
Hospital, the Swedish Medical Research Council, Alfred Osterlund foundation, the Great and Johan Kock Foundation, Maggie Stephens Foundation, Thure Carlsson Foundation, faculty of Medicine at Lund University.

Cassidy 2003
Randomized Prospective Trial

One of more of the authors received grants or outside funding from the Norian Corporation. Three authors were employees of Norian.

5.0

N=323 (51 males/272 females) patients who had sustained a displaced and/or unstable distal radial fracture; Mean Age Group 1: 63.5 ± 11. Group 2: 63.7 ± 12.

N=161 patients treated with Norian SRS cement and a closed reduction. (Group 1) Vs N=162 patients treated only with closed reduction and either external fixation or cast immobilization (Group 2).

Follow up evaluations were specified at 1, 2, 4, and 6-6 weeks, also at 3, 6, and 12 months.

Group 1 v Group 2 subjective pain rating difference; Group 1 lower at 2 and 4 weeks, (p=0.02, p=0.02, respectively). Group 1 required less pain medicine at 2 weeks (p=0.004). Group 1 vs Group 2 grip hand strength at 6-8 weeks, 18 lb vs 10 lb (p<0.0001). Group 1 at 6-8 weeks had better digital range of motion (p<0.01). Group 1 had significantly less swelling of forearm at 2 weeks, (p=0.0146), and various digits at 6-8 weeks. Jebsen dexterity test, Group 1 dominant hand fracture at 6-8 weeks took less time to pick up small objects (p=0.0023). Group 1 vs group 2, ulnar variance at 12 months. 2.0 vs 1.4 (p=0.02).

Complications largely due to loss of reduction, no significant difference in complications between groups.

"Our data suggest that Norian SRS cement provides adequate fixation for the majority of distal radial fractures to permit early wrist mobilization."

Data suggest Norian SRS cement is beneficial for most distal radial fractures and may allow faster recovery due to accelerated rehabilitation. The control group experienced a significantly higher number of post procedure infections.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Age</th>
<th>Intervention</th>
<th>Follow up</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Földhazy 2010</td>
<td>RCT</td>
<td>59 (6 males/53 females) with displaced fractures of the distal radius; Mean Age Group 1: 70 (62-81) Group 2: 73 (60-85).</td>
<td>Group 1 N=29 patients treated with open reduction and internal fixation with plaster casting. Vs Group 2 (N=22) Patients treated with closed reduction and external fixation. Follow up at 2, and 5 weeks, 2, 6, and 12 months.</td>
<td>No significant difference in Clinical outcomes, and complications. Slightly better dorsal extension and radial deviation in group 1 at final follow up (p=0.036 and p=0.043, respectively). Final dorsal angulation, group 2 vs 1, 1 year; 11±9 vs 20±14 (p=0.001).</td>
<td>“We believe that the results of this prospective, randomized and comparative study are that in 60–85 year old patients, with a displaced distal radial fracture after low energy trauma, no obvious clinical benefit could be demonstrated using closed reduction and external fixation as compared with closed reduction and plaster treatment.”</td>
<td></td>
</tr>
<tr>
<td>Kreder 2005</td>
<td>RCT</td>
<td>179 (109 males/70 females) skeletally mature patients with displaced intra-articular fractures of the distal radius; Mean Age Group 1: 40 (20-78). Group 2: 39 (20-81)</td>
<td>Group 1 (N=88) patients treated with Closed reduction and K-Wire Fixation. Vs Group 2 (N=91) Patients treated with open reduction and internal fixation. Follow up at 6 weeks, 12 and 24 months.</td>
<td>Patients in group 1 had better function overall, scoring a mean of 6 points (95% CI: 4.1-33.0) in Musculoskeletal Functional Assessment (MFA). Pain scores were better overall for group 1 (p=0.052) NS. MFA, group 1 vs 2, 6 months; 15.1 vs 37.9 Difference: -12.8 (95% CI: -23.7 - -1.9). Grip Strength, group 1 vs 2, improved throughout study by 10.1 lb (p=0.05).</td>
<td>“We recommend that open reduction be preceded by an attempt at minimally invasive percutaneous reduction. If an acceptable reduction is achieved then open reduction is unnecessary and function will be superior.”</td>
<td></td>
</tr>
<tr>
<td>Arora 2011</td>
<td>Prospective Randomized Trial</td>
<td>73 (18 males/55 females) with distal radial fracture that were unstable; Mean Age 76.7 (65-89).</td>
<td>Group 1 (N=36) individuals who were treated with open surgery and fixed with K-Wire, volar locking plate, or DVR. Vs No significant differences in clinical parameters. Significantly more complications in the operative treatment group (p&lt;0.05). DASH scores, 6 weeks group 1 vs 2; 18.8±17.9 vs 34.4±22.5 (p&lt;0.001). Patient-Rated Wrist Evaluation (PRWE) scores at 6 weeks; group 1 vs 2; 36.4±28.7 vs 64.9±29.0 (p&lt;0.001). DASH “Volar fixed-angle plate systems have made plate osteosynthesis popular for elderly individuals with osteoporotic bones. However, at twelve months after surgery, the active range of motion, the pain level, and the PRWE and the DASH scores suggest at 12 months, ROM, pain level and PRWE and DASH scores equivalent. Patients in surgical group reported better grip strength throughout trial.</td>
<td>Data suggest comparable efficacy between groups although primary external fixation group showed a positive radiographical effect. However, one third of the external fixation group had a complication.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>Participants</td>
<td>Interventions</td>
<td>Follow-up</td>
<td>Results</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>--------</td>
<td>--------------</td>
<td>--------------</td>
<td>-----------</td>
<td>---------</td>
</tr>
<tr>
<td>Karantana 2013</td>
<td>RCT</td>
<td>No sponsorship or COI</td>
<td>N=130 (no mention of gender) patients with a distal radial fracture; Mean Age not listed (18-73)</td>
<td>Group 1 (N=64) patients treated with open reduction and volar plating. Vs Group 2 (N=66) patients treated with closed reduction and external fixation.</td>
<td>Follow-Up at 6, and 12 weeks, also at 1 year.</td>
<td>Patient evaluation measure (PEM), group 1 vs 2, 6 weeks; 34±13 vs 45±12 (p&lt;0.001). Quick Dash, group 1 vs 2, 6 weeks; 41±21 vs 52±20 (p=0.002). Grip Strength (% vs uninjured arm), group 1 vs 2, 6 weeks; 40±23 vs 10±12 (p&lt;0.001). 12 weeks; 65±26 vs 45±22 (p=0.002). 1 year; 95±22 vs 84±19 (p=0.005). Range of Motion, group 1 vs 2, 6 weeks; Extension (deg): 57±22 vs 17±30 (p&lt;0.001). Flexion (deg): 59±18 vs 47±22 (p=0.001). Pronation (deg): 80±17 vs 65±28 (p=0.001). Supination (deg): 73±23 vs 37±26 (p&lt;0.001). More complications within group 2; (p=0.047).</td>
</tr>
<tr>
<td>Leung 2008</td>
<td>RCT</td>
<td>In support of their research for or preparation of this work, one or more of the authors received, in any one year, outside funding or</td>
<td>N= 137 (85 males/52 females) with an acute intra-articular of distal radial fracture; Mean Age 44.</td>
<td>Group 1 (N=74) fractures that were treated using external fixation and percutaneous pinning. Vs Group 2 (N=70) fractures that were stabilized with plates.</td>
<td>Follow-Up 6, 12, and 24 months.</td>
<td>Gartland and Werley point system results, group 1 vs group 2 at 24 months; 39% excellent, 55% good, 6% fair, 0% poor, vs 67% excellent, 30% good, 3% fair, 0% poor (p=0.04). Arthritis grade, group 1 vs group 2, at 24 months; 24% grade-0, 65% grade-1, 15% grade-2. Vs 44% grade-0, 52% grade-1, 4% grade-2 arthritis (p=0.001).</td>
</tr>
</tbody>
</table>

Data suggest comparable efficacy at 3 months and 1 year post procedure. The volar locking plate group did demonstrate some increased grip strength as well as anatomical improvement but these results were not significant.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Sex Distribution</th>
<th>Fracture Type</th>
<th>Mean Age Group 1</th>
<th>Mean Age Group 2</th>
<th>Treatment Group 1</th>
<th>Treatment Group 2</th>
<th>Follow-Up</th>
<th>Results</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grewal 2005</td>
<td>Prospective Randomized Trail</td>
<td>62</td>
<td>33 males/29 females</td>
<td>AO type C intra-articular distal radius fractures</td>
<td>45 ± 2.7</td>
<td>46 ± 2.7</td>
<td>Open reduction and internal fixation</td>
<td>Mini open reduction with percutaneous K-Wire fixation</td>
<td>2, 4, 6, 10-12 weeks, 6 months, and 1, 2 years</td>
<td>Complication Rate, Group 1 vs 2; 72.4% vs 24.2% (p=0.004). Grip Strength (% vs uninjured arm), group 1 vs 2, 86% vs 97% (p=0.019). Range of motion not significantly different. Radiographic outcomes not statistically different. Pain scores (DASH), group 1 vs 2, at 1 year; 22.1 vs 10.0 (p=0.02). after hardware taken out in some of the group 1 patients pain scores equalized at 2 years.</td>
<td>“Although dorsal Pi plates still may have a role in treating intra-articular distal radius fractures we have shown that mini open reduction with percutaneous K-wire and external fixation is a technique that provides a safe and effective alternative to open reduction and dorsal Pi plating when treating comminuted intra-articular distal radius fractures.” Data suggest comparable efficacy between groups with the dorsal plate groups having greater numbers of complications.</td>
</tr>
<tr>
<td>Jeyam 2002</td>
<td>RCT</td>
<td>21</td>
<td>0 males/21 females</td>
<td>Distal radial fractures type 1 and 2</td>
<td>74</td>
<td>71</td>
<td>K-wire using intrafocal technique, then castred for 4 weeks. (Group 1) vs (N=9) fracture site was cleaned and injected with Orthofix Bone source bone cement (Group 2).</td>
<td>Follow-Up at 1 day, and 1, 23, 6, 12, and 26 weeks.</td>
<td></td>
<td>Group 2, 1 week all three radiological parameters had deteriorated. Group 1 Vs 2, dorsal angle at 1 week, -7 (-19-6) and 6 (-5-15) (p&lt;0.05) remained significant throughout the entire study. Radial angle worse in group 2, not significant. Group 1 vs Group 2, Grip strength at 6 months: 11 (6-17) vs 8 (4-10) (p&lt;0.03).</td>
<td>“The results of this small study clearly indicate that hydroxyapatite cement (Bonesource) does not provide adequate fracture stability when used alone.” Data suggest that at 12 and 26 weeks, the hydroxyapatite group performed worse on grip strength, palmar flexion and dorsal flexion. There were no outcome measures where this group performed better.</td>
</tr>
</tbody>
</table>
Ganglion Cyst

Special Studies and Diagnostic and Treatment Considerations

There are no quality randomized trials for diagnostic testing in the evaluation of ganglia of the upper extremity. Generally, diagnosis is based on physical examination findings. Diagnosis is usually confirmed upon aspiration of mucinous fluid from the mass.

X-RAYS

1. Recommendation: Routine X-rays for Diagnosis of Wrist Ganglia
   
   X-ray to diagnose dorsal or volar wrist ganglia in select patients is recommended.

   Indications – Ganglia, especially occurring in the context of trauma where fracture may be present.
   
   Frequency/Duration – Obtaining x-rays once is generally sufficient.

   Strength of Evidence – Recommended, Insufficient Evidence (I)
   
   Level of Confidence – Low

2. Recommendation: Routine Use of X-rays for Evaluation of Dorsal or Volar Wrist Ganglia
   
   The routine use of x-ray to evaluate dorsal or volar wrist ganglia is not recommended.

   Strength of Evidence – Not Recommended, Insufficient Evidence (I)
   
   Level of Confidence – Moderate

Rationale for Recommendations

Patients develop ganglia for numerous reasons, ranging from trauma to arthritis to idiopathic. The threshold for obtaining x-rays should be low. Patients incurring ganglia due to trauma or other inciting events that may result in other traumatic sequelae such as fractures, dislocations, and sprains, should have x-rays. Patients incurring ganglia through non-traumatic means are candidates for initial management without x-rays. Some practitioners advocate the use of x-rays for routine evaluation of all patients with dorsal or volar wrist ganglia. However, there is no supporting evidence for this practice. In a prospective case series of 103 patients with volar and dorsal ganglia, three view wrist radiographs were obtained and a retrospective review of medical records completed. Findings on x-ray altered the course of management in 1 case (1%). (1338)

Evidence for the Use of X-rays

There is 1 low-quality study included in Appendix 2. (Sakamoto 13)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ganglion, Cyst, Cysts, Xray, X-ray, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 371 articles in PubMed, 298 in Scopus, 2 in CINAHL, 0 Cochrane Library, and 3240 from Google Scholar. We considered for inclusion 1 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 3911 articles considered for inclusion, 1 met the inclusion criteria.

MRI

Recommendation: MRI for Evaluation of Wrist Pain with Suspected Occult Dorsal or Volar Wrist Ganglia

There is no recommendation for or against the use of MRI for the evaluation of wrist pain with suspected occult dorsal or volar wrist ganglia as it may be of limited benefit in deciding on the course of treatment.

   Strength of Evidence – No Recommendation, Insufficient Evidence (I)
   
   Level of Confidence – Low

Rationale for Recommendation

In a small study of 20 patients with suspected occult ganglia, an MRI was obtained prior to surgical exploration and excision of the cyst. Comparison of MRI diagnosis with intra-operative findings and histological evaluation of the excised specimen resulted in MRI scanning sensitivity of 83%, specificity of
50%, and a positive predictive value of 94%.(292) The findings suggest in the absence of palpable mass, with no history of trauma or other conditions such as arthritis, an MRI may be beneficial for the diagnosis of occult symptomatic ganglia. However, in light of the results reported by Lowden,(114) which found nearly half of the asymptomatic population have an occult ganglia, the accuracy of these findings for screening purposes are questionable and the utility of a positive result may be of less clinical consequence. MRI may be useful in distinguishing synovitis from ganglion, which may be useful in determining the course of treatment.(1339) MRI is reasonable for patients who have had persistence of pain consistent with a ganglion lasting at least 3 weeks without trending towards improvement.

Evidence for the Use of MRI
There are 4 moderate-quality studies incorporated into this analysis.(1440-1443) (Anderson 06; Goldsmith 08; Vo 95; Cardinal 94)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: MRI, Magnetic resonance imaging, Ganglion Cyst, Wrist, hand, Ganglion, ganglia, dorsal, volar, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 19 articles in PubMed, 2037 in Scopus, 1 in CINAHL, 8 Cochrane Library, and 40 from Google Scholar. We considered for inclusion 0 from PubMed, 3 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 1 from other sources. Of the 4 articles considered for inclusion 4 diagnostic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Years</th>
<th>Score</th>
<th>Number</th>
<th>Study Type</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>Type of MRI used</th>
<th>Type of CT used</th>
<th>T1 weighted</th>
<th>T2 weighted</th>
<th>Myelography</th>
<th>X-ray</th>
<th>More than one rater</th>
<th>Long term follow-up</th>
<th>Clinical outcomes assessed</th>
<th>Surgery Performed</th>
<th>More than one rater</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anderson 2006</td>
<td>6.0</td>
<td>34 patients</td>
<td>Retrospective</td>
<td>Wrist</td>
<td>Dorsal occult ganglion cyst</td>
<td>1.5-T superconducting magnet</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>More than one rater</td>
<td>Negative</td>
<td>35 abnormalities were diagnosed with MRI: 25 ganglia, 16 dorsal occult ganglia and 6 synovitis. Surgery confirmed MRI diagnoses with an overall agreement of 71% (95% CI, 0.38-0.76). Sensitivity to ganglia was 89% (95% CI 56%-99%) to dorsal occult ganglia cysts was 94% (95% CI 70%-100%).</td>
<td></td>
</tr>
<tr>
<td>Goldsmith 2008</td>
<td>5.5</td>
<td>20 patients</td>
<td>Retrospective</td>
<td>Wrist</td>
<td>Occult dorsal ganglion cyst</td>
<td>Siemens’ 1.5 T imager</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>More than one rater</td>
<td>Positive</td>
<td>MRI found 16 of 20 wrist had an occult ganglion. Surgery was performed on all 20 patients, identifying 18 occult ganglion.</td>
<td>MRI scans provide relatively good reliability in establishing the diagnosis of an occult</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20 wrists</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Data suggest MRI is a good technique for visualizing occult dorsal wrist ganglia.</td>
</tr>
</tbody>
</table>
the 20 wrists had histologic features of a ganglion cyst. The 4 negative MRI were positive and 3 of the 18 positive in surgery were negative. MRI at the time of surgery provided a sensitivity of 83%, a specificity of 50% and a PPV of 94%. However, when evaluated with histologic findings, the sensitivity was 80%, specificity was 20%, the PPV was 75% and the accuracy was 65%. dorsal wrist ganglion
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Patients</th>
<th>Age or Gender</th>
<th>Imaging Details</th>
<th>Positive Patients</th>
<th>MRI Positive</th>
<th>Surgery</th>
<th>Histology</th>
<th>PPV</th>
<th>Notes</th>
</tr>
</thead>
</table>
| Vo 1995 Retrospective | 4.0  | 14       | No mention of age or gender | Chronic Dorsal Wrist pain of unknown etiology | 1.5-Tesla General Electric Signa | - + - - - + + + - | 10 of 14 were positive for occult dorsal wrist ganglion on the MRI. 7 of the 10 MRI positive patients underwent surgery after nonoperative treatment failed and was confirmed as positive through histologic examination. One of the positive patient developed a palpable ganglion. The two other positives were not confirmed. The PPV is 100%.
|                       |      |          |               |                 |                   |              |         |            |     |       |
| Cardinal 1994         | 4.0  | 14 wrists in 13 patients | Mean age = 30, 9 women, 4 men | Occult dorsal carpal ganglion | 1.5-T Imager, Signa Advantage | - + - - - + + + - | US identified 11 dorsal carpal ganglion cyst while MRI identified 9. One patient that was positive on US | MRI imaging and US are equally effective in the detection of occult dorsal carpal ganglion. Small sample. Data suggest comparable efficacy between MRI and US for detection. |
**ULTRASOUND**

**Recommendation:** Ultrasound for Evaluation of Chronic Wrist Pain with Suspected Occult Dorsal or Volar Wrist Ganglia

There is no recommendation for or against the use of ultrasound for the evaluation of chronic wrist pain with suspected occult dorsal or volar wrist ganglia. It may be beneficial in select cases in deciding on the course of treatment.

*Strength of Evidence – No Recommendation, Insufficient Evidence (I)*

*Level of Confidence – Low*

**Rationale for Recommendation**

In a small study of 57 patients with non-traumatic wrist pain and no palpable mass, ultrasound was used to determine the presence of ganglia at the wrist – 33 patients (58%) were found to have a ganglia of which 20 were treated with excision or aspiration and improvement of symptoms after the intervention. As MRI has demonstrated the prevalence of ganglia in asymptomatic study volunteers to be nearly 50%, (114) there is likely a high probability of finding ganglia on ultrasound as well. Thus, a positive finding of ganglion by ultrasound is of unknown clinical significance, particularly in that the study did not provide long-term follow-up for all of the patients that were found to have a ganglion cyst. If ultrasound is utilized, it would appear to be reasonable among patients who have had persistence of pain lasting at least 3 weeks without trending towards improvement.

**Evidence for the Use of Ultrasound**

There is 1 moderate-quality study incorporated into this analysis. (1444) (Osterwalder 97)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: ultrasonography, ultrasound, sonography, ganglion cysts, ganglion, ganglia, dorsal, volar, hand, wrist, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 43 articles in PubMed, 94 in Scopus, 0 in CINAHL, 7 in Cochrane Library, and 2,190 from Google Scholar. We considered for inclusion 1 from PubMed, 0 from...
Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 1 article considered for inclusion 1 diagnostic study met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>Type of Ultrasound</th>
<th>CT used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Osterwaler, 1997 Diagnostic</td>
<td>6.0</td>
<td>Wrist</td>
<td>suspected occult wrist ganglion who complained of wrist pain and palpation findings were inconclusive</td>
<td>For first three years - Aloka model SSD-6202S, for last two years - Hitachi model EUB-55S, both models used 7.5-MHz linear transducer and spacer</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>Out of the 168 patients examined by ultrasound 68 were diagnosed with a cyst and 85 were diagnosed with absence of a cyst. In 15 patients the diagnosis was not clear enough to get a definitive answer. Ultrasound sensitivity, specificity, accuracy, positive predictive value and negative predictive values plus the 95% confidence intervals were the following: 88% (73-96%), 85% (64-95%), 87% (76-94%), 90% (75-97%), 83% (62-94%)</td>
</tr>
</tbody>
</table>

**Initial Care**

1. Recommendation: Non-operative Management (No Treatment) for Acute Asymptomatic Wrist and Hand Ganglia

The use of non-operative management (no treatment) for acute asymptomatic wrist and hand ganglia is recommended as first-line management as the natural history for spontaneous resolution is more than 50%, and in recognition of the high recurrence rate of most other treatment strategies.

**Strength of Evidence** – Recommended, Insufficient Evidence (I)

**Level of Confidence** – Low

Rationale for Recommendation
There are many observational studies describing the natural history for ganglia to resolve without any treatment over time. More than 50% are likely to resolve within months to years. A recently published 6-year follow-up, reported a 58% spontaneous resolution rate in patients that received no intervention.(1340) Thus, in the asymptomatic patient, it is reasonable to provide patients reassurance that the mass is benign, and that the natural course is for most to resolve without treatment, making waiting a reasonable trial. However, patients may wish to have an intervention for cosmetic relief, and have reported higher satisfaction despite the higher risk of surgical or interventional complications.(1340)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: non operative management, no treatment, ganglion cyst, wrist, hand, ganglion, ganglia, dorsal, volar; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 56 articles in PubMed, 30 in Scopus, 0 in CINAHL, 3 in Cochrane Library, 12596 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar, and 0 articles from other sources. Zero articles met the inclusion criteria.

2. **Recommendation: Aspiration (without Other Intervention) for Acute Cosmetic and Ganglia Related Pain**

Aspiration (without other intervention) of the cystic fluid is recommended as it may result in immediate relief of acute cosmetic and ganglia related pain.

*Duration* – One aspiration is recommended.(1341) However, a long-term course of aspiration is usually of no benefit in terms of resolution. There is no recommendation on how many times aspiration should be attempted before advancing to other intervention. Variants of simple aspiration include steroid injection, splinting, multiple punctures, hyaluronidase, and sclerosing agents, reviewed below.

*Strength of Evidence* – **Recommended, Evidence (C)**

*Level of Confidence* – **Low**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: aspiration; ganglion cyst, wrist, hand, ganglion, ganglia, dorsal, volar; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 11 articles in PubMed, 29 in Scopus, 0 in CINAHL, 5 in Cochrane Library, 8,180 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 1 from Cochrane Library, 0 Google Scholar, and 1 from other sources. Of the 3 articles considered for inclusion, 2 randomized trial and 1 systematic studies met the inclusion criteria.

3. **Recommendation: Aspiration with Steroids**

There is no recommendation for or against the addition of steroids with aspiration.

*Strength of Evidence* – **No Recommendation, Insufficient Evidence (I)**

*Level of Confidence* – **Low**

**Rationale for Recommendations**

Aspiration with instillation of steroids is the most common treatment for upper extremity ganglia. Recurrence rates range from 14 to 83%. There are no quality studies that compare simple aspiration with the addition of steroids; thus, no quality evidence to address whether this results in potential benefits. However, a review of cohorts has shown an average recurrence rate of 51% for aspiration alone, and a recurrence rate of 52% with aspiration and steroids.(118) As the cystic structure has been shown histologically and with electron microscopy to have no synovial lining, but rather a network of collagenous fiber layers, there is little theoretical reason to believe that steroid agents would result in reducing
inflammation, as there is theoretically no tissue in the cyst to be inflamed. There is no recommendation for or against steroids when aspiration is used for immediate relief.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ganglion Cyst (wrist ganglia, dorsal or volar wrist ganglia), Aspiration with steroids; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 4 articles in PubMed, 15 in Scopus, zero in CINAHL, zero in Cochrane Library, 498 in Google Scholar, and zero from other sources. We considered for inclusion 3 from PubMed, zero from Scopus, zero from CINAHL, zero from Cochrane Library, zero from Google Scholar, and zero from other sources. Of the 3 articles considered for inclusion, 3 randomized trials and zero systematic studies met the inclusion criteria.

4. **Recommendation: Aspiration and Multiple Punctures of Cyst Wall**
   The technique of multiple punctures of the cyst wall is not recommended as it does not provide improved benefit over simple aspiration.

   **Strength of Evidence** – Not Recommended, Insufficient Evidence (I)
   **Level of Confidence** – Low

**Rationale for Recommendation**
There is one quality study comparing simple aspiration with multiple wall punctures,(1342) which did not show any significant difference in efficacy. A review of non-RCT studies comparing aspiration with multiple punctures showed an average of 64% recurrence rate, which is worse than aspiration alone.(118) Thus, there is no added benefit to making multiple punctures in the cystic wall, and may result in additional skin trauma and higher risk of infection, making this intervention not recommended.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Aspiration and multiple punctures of cyst wall, Ganglion Cyst (wrist ganglia, dorsal or volar wrist ganglia); controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed zero articles in PubMed, 2 in Scopus, zero in CINAHL, zero in Cochrane Library, 155 in Google Scholar, and zero from other sources. Zero articles met the inclusion criteria.

5. **Recommendation: Splinting after Aspiration for Acute or Subacute Dorsal or Volar Wrist Ganglia**
   There is no recommendation for or against the use of splinting after aspiration for the treatment of acute or subacute dorsal or volar wrist ganglia as splinting may have uncertain efficacy and may lead to prolonged joint stiffness.

   **Strength of Evidence** – No Recommendation, Insufficient Evidence (I)
   **Level of Confidence** – Low

**Rationale for Recommendation**
There are no quality studies comparing immobilization as an adjunct treatment. In a prospective series, immobilization after aspiration was not found to be of any significant benefit compared those without immobilization in a 1-year prospective study of volar, dorsal and digital ganglia.(1343) However, in an earlier study including multiple punctures, immobilization had a positive effect for successful outcomes.(1344) These conflicting results, in the absence of quality experimental data, preclude making recommendation for or against this intervention.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: aspiration, splint, splints, splinting, ganglion cyst, wrist, hand, ganglion, ganglia, dorsal, volar; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We
found and reviewed 0 articles in PubMed, 2 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 1,294 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

6. **Recommendation: Hyaluronidase Instillation after Aspiration**

   **There is no recommendation for or against the instillation of hyaluronidase into the cystic structure after aspiration.**

   **Strength of Evidence – No Recommendation, Insufficient Evidence (I)**
   
   **Level of Confidence – Low**

   **Rationale for Recommendation**

   One moderate-quality study compared the standard therapy of aspiration and steroids with the addition of hyaluronidase to the mixture. (1345) Although the study showed a positive effect on the patient reporting for excellent results, it was not statistically significant for good and excellent combined between the two groups. Thus, there is insufficient evidence for recommendation for or against this intervention.

   A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: aspiration, hyaluronoglucosaminidase, hyaluronidase, Ganglion Cyst, Wrist, hand, Ganglion, ganglia, dorsal, volar; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 2 in Scopus, 0 in CINAHL, 1 in Cochrane Library, 376 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 0 from other sources. Of the 1 article considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

7. **Recommendation: Aspiration and Sclerosing Agents**

   **Sclerosing agents such as phenol and hypertonic saline, which when instilled are intended to result in scarring and closure of the cystic potential space, are not recommended.**

   **Strength of Evidence – Not Recommended, Insufficient Evidence (I)**
   
   **Level of Confidence – Low**

   **Rationale for Recommendation**

   As the cystic structure as described histologically and with electron microscopy have determined there is no synovial lining, rather a network of collagenous layers, there is little theoretical reason to believe that sclerosing agents would result in inciting an inflammatory reaction. In one small prospective study of 29 patients in Africa, 2cc of hypertonic saline injected into the cyst structure after aspiration was reported to result in only one recurrence after a 2-year follow-up. (1346) A small study of 10 patients treated with phenol injection was reported with good results. (1347) From anatomic studies, it has been shown that the cystic structure is connected to the synovial space in some cysts, so that there is some theoretical risk that instilling sclerosing agents will directly enter into a joint with resultant poor consequences. Thus, these therapies are only reported in small studies with higher risk of causing harm, and are not recommended.

   A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: aspiration and sclerosing agents, phenol and hypertonic saline, ganglion cyst, wrist, hand, ganglion, ganglia, dorsal, volar; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, Scopus, CINAHL, Cochrane Library, 346 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, Scopus, CINAHL, Cochrane Library, Google Scholar, and 0 articles from other sources. Zero articles met the inclusion criteria.
8. **Recommendation: Surgical Excision for Subacute or Chronic Upper Extremity Ganglia**

   Surgical intervention is recommended for treatment of subacute or chronic upper extremity ganglia after a trial of non-operative management.

   *Strength of Evidence – Recommended, Evidence (C)*
   
   *Level of Confidence – Moderate*

   **Rationale for Recommendation**

   Surgical intervention is the most effective treatment method for upper extremity ganglia despite the significant recurrence rates and higher risk of complications.(115, 1341, 1348) As most upper extremity ganglia are asymptomatic, consideration of surgical risks and a trial of non-operative management are prudent before performing a surgical procedure for cosmetic reasons. One moderate-quality study exists comparing the recurrence rates of surgery to aspiration with steroids.(115) With a sample size of 28 dorsal ganglia, the success rate at 6 months was significantly higher with surgery (82% vs. 38%, p <0.05). The generalizability of the study is limited because of the small sample and the exclusion of other ganglia types. The success of surgery reported in non-randomized prospective case series suggest an overall recurrence rate between 5% and 40%. More recent surgical techniques that include comprehensive dissection and excision of the cyst, pedicle, and a cuff of the adjacent joint capsule are believed to have better results.(118)

   A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Surgical Excision, Ganglion Cysts, Ganglion, Ganglia, Dorsal, Volar, Hand, Wrist; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, random; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 11 in Scopus, 1 in CINAHL, 5 in Cochrane Library, 20 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 1 from Cochrane Library, 0 Google Scholar, and 0 from other sources. Of the 2 articles considered for inclusion, 2 randomized trials and 0 systematic studies met the inclusion criteria.

9. **Recommendation: Arthroscopic versus Open Excision**

   There is no general indication for one surgical technique (arthroscopic or open excision) over another for all cases and both are recommended. There may be advantages of arthroscopic procedures for ganglia originating in the radiocarpal joints, whereas open excision may have advantages in ganglia originating in midcarpal joints, although both have the same success rate.

   *Strength of Evidence – Recommended, Evidence (C)*
   
   *Level of Confidence – Low*

   **Rationale for Recommendation**

   There are two moderate-quality studies comparing open excision to arthroscopic resection of wrist ganglia. In both studies, rates of recurrence were low and not significantly different,(1349, 1350) thus showing no clear advantage for either technique. However, when comparing outcomes results for lost time, complications and functionality, arthroscopic excision of radiocarpal ganglia had faster recovery time and fewer complications than open excision, whereas open excision had better recovery and fewer complications than arthroscopic excision for midcarpal ganglia.(1350) However, these conclusions are weakened by small sample size and lack of statistical analyses to make recommendation for or against difficult. In a non-randomized trial,(116) the effectiveness of arthroscopic excision of dorsal ganglia in a 2-year follow-up study demonstrated 5% recurrence, although failure with arthroscopy was treated with open excision.

*Evidence for Aspiration and Surgery for Ganglion Cysts*

There are 67 moderate-quality RCTs incorporated into this analysis.(115, 1446, 1447, 1450, 1456-1459) (; Jagers Op Akkerhuis 02) There are 2 low-quality RCTs in Appendix 2. (1453, 1460) (Balazs 15, Varley 97)
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Arthroscopy, Arthroscopic, Open Excision, Surgery, Ganglion Cysts, Ganglion, Ganglia, Dorsal, Volar, Hand, Wrist; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 2 in Scopus, 1 in CINAHL, 1 in Cochrane Library, 20 in Google Scholar, and 2 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 1 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 2 from other sources. Of the 2 articles considered for inclusion, 0 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Aspirations and Multiple Puncture group</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stephen 1999 RCT No mention of sponsorship or COI.</td>
<td>4.0</td>
<td>N = 119 (male to female ratio 1:3.1) with ganglia. Age not given.</td>
<td>Simple aspiration (N = 65) vs Aspiration and multiple wall punctures (N = 54)</td>
<td>Follow-up for 1 year.</td>
<td>“16 of 51 ganglia (31%) treated by aspiration alone resolved and did not recur in contrast to 9 of 41 ganglia (22%) in the multiple puncture group.”</td>
<td>“The study has demonstrated that multiple puncture of the ganglion wall does not improve the results of simple ganglion aspiration.” Lack of study detail. No randomization or allocation details. Drop-out 23% at 1-year follow-up.</td>
</tr>
<tr>
<td><strong>Aspiration and Steroid Alone (prior use of Hyaluronidase)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Paul 1997 RCT No mention of sponsorship or COI.</td>
<td>4.0</td>
<td>N = 70 (29 male and 41 female) with ganglia of the wrist or hand. Mean age given.</td>
<td>Group 1, local anesthetic of 0.5% lignocaine plus 0.5 ml of ganglion contents were aspirated v via a 16 gauge needle (N = 35) vs Group 2, treated by conventional technique of aspiration under local anesthetic and immediate injection of 40 mg of Depomedrone (N = 35).</td>
<td>2 year follow-up.</td>
<td>Patients reporting excellent results significantly higher in hyaluronidase group (49% vs. 20%, p = 0.0051). However, good and excellent ratings combined showed trend for hyaluronidase (89% vs. 57%) but not significant, (p = 0.072).</td>
<td>“The cure rate with the combined use of hyaluronidase and methylprednisolone was 89% compared to 57% when treated by aspiration and instillation of methylprednisolone alone.” Lack of study details. 100% follow-up achieved at 2 years. Treatment may be beneficial for viscous cystic fluid that is too viscous for aspiration.</td>
</tr>
<tr>
<td><strong>Aspiration and Surgical Excision and Steroid Injection</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Limpaphayom 2004 RCT No mention of sponsorship or COI.</td>
<td>6.5</td>
<td>N = 28 (24 female and 4 male). Age surgery/aspiration: 29.9±9.79/32.0±13.08.</td>
<td>Surgery, 5 cc of 1% Xylocaine (N = 11) vs Aspiration, steroids, and immobilization (N = 13).</td>
<td>Follow-up for 6 months.</td>
<td>At 6 month follow-up, the success rate was 81.8% by surgical excision and 38.5% by aspiration,(p = 0.047).</td>
<td>“Result of treatment can be varied but by this RCT, surgery was shown to obtain a superior result in terms of success rate than aspiration, methylprednisolone acetate injection plus wrist immobilization.” Single trial of aspiration. Lack of blinding. Only included dorsal wrist ganglia.</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Description</td>
<td>Baseline Follow-up</td>
<td>Results</td>
<td>Notes</td>
<td></td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-------</td>
<td>-----------------------------------------------------------------------------</td>
<td>----------------------</td>
<td>---------------------------------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Latif 2014 RCT</td>
<td>4.0</td>
<td>N = 173 (36 male and 147 female) with ganglia within wrist, ankle and knee.</td>
<td>Group 1 who opted for aspiration and injection treatment (N = 143) vs Group 2 who opted for surgical treatment (N = 44). Follow-up baseline and 6 months.</td>
<td>Group 1 vs group 2 success at third week of injection: 82 (57%) vs 41 (33%). Success rate at 6 months (116 (81%) vs 0 (0%). Failure rate within group 1 vs group 2: 27 (19%) vs 3 (7%) (p &lt;0.028).</td>
<td>In symptomatic ganglia, surgical excision is a better treatment option as the failure rate is less compared to triamcinolone acetonide injection after aspiration.</td>
<td></td>
</tr>
<tr>
<td>Jagers Op Akkerhuis 2002 RCT</td>
<td>4.5</td>
<td>N = 89 (27 males, 62 females). Mean age is 39.5 years.</td>
<td>Hyaluronidase + Aspiration (N = 43) vs Surgical Excision (N = 46). Follow-up 1 year.</td>
<td>Hyaluronidase treatment resulted in recurrence in 33 of the 43 patients (77%). Recurrences after surgery were found in 11 of the 46 (24%) patients: six within 3 months and five between 3 months and 1 year.</td>
<td>Surgical excision is preferable to aspiration after hyaluronidase, assuming that the aim of treatment is resolution of the ganglion. However hyaluronidase and aspiration has a 23% success rate and can be used for those patients who prefer not to undergo surgery.</td>
<td>Data suggest surgical excision best treatment for symptomatic ganglia vs. injection-aspiration. At 6 months, injection-aspiration success rate 81.0% vs. surgical excision 93.0%. Failure rates significant at 19.0% for injection-aspiration group and 7.0% for surgical excision group.</td>
</tr>
<tr>
<td>Rocchi 2008 RCT</td>
<td>4.5</td>
<td>N = 51 (17 male and 24 female) with dorsal wrist ganglions. Mean age 29.8 years.</td>
<td>Arthroscopic resection (N = 41) vs Open excision of volar ganglion cyst (N = 10). Follow-up for 47.8 months.</td>
<td>Comparisons by radiocarpal ganglia (RCG) and midcarpal ganglia (MCG) locations. For open resection of RCG, mean functional recovery time 13 days with mean time lost from work 21 days, 15/20 reporting good results at 24 months and 3 bad results. Arthroscopic RCG 18/20 good results with 9 days recovery time and 9 days lost time. MCG subgroup, 5/5 good results with open excision with functional recovery time 10 days, lost time 17 days; 1/5 in arthroscopic group treated successfully.</td>
<td>Comparing our two groups, we noted rather better results with arthroscopy in the treatment of radiocarpal ganglia, and better results for open operation in the treatment of midcarpal ganglia.</td>
<td>No statistical analyses presented.</td>
</tr>
<tr>
<td>Kang 2008 RCT</td>
<td>4.0</td>
<td>N = 72 with ganglion recurrence or wrist pain. Mean age for the open</td>
<td>Arthroscopic technique consisted of 2 stab incisions at the standard 3-4 and 4-5 portal sites (N = 41)</td>
<td>At 4-8 weeks, 1/41 in arthroscopic group vs. 0/31 in open excision group had recurrence (p = 0.381). 17% in arthroscopic group reported success.</td>
<td>The results of our study suggest that the technique of arthroscopic surgery does not achieve superior rates of ganglion recurrence.</td>
<td>Lack of study details. High attrition rate at 12 month follow-up. No blinding.</td>
</tr>
</tbody>
</table>
No sponsorship. No mention of COI.

| group was 36 years and for the arthroscopic group 34 years. vs Open excision of dorsal ganglion cyst (N = 31). | Follow-up of 12 months. | residual pain vs. 10% (p = 0.369). At 1 year, no significant difference in pain or recurrence. |
Follow-up Visits
Because of the natural course of spontaneous resolution and recurrence, follow-up should be dictated by the course of treatment selected by the patient and physician.

Prescription Medications
No prescription medications are shown to be effective for treatment of upper extremity ganglia. By inference from other musculoskeletal conditions, NSAIDs may be of benefit as an analgesic for ganglia associated wrist pain, although there is no evidence of their efficacy.

Evidence for the Use of Medications
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: anti-inflammatory agents, non-steroidal, NSAIDS, non-steroidal anti-inflammatory, ibuprofen, acetaminophen; ganglion cyst, wrist, hand, ganglion, ganglia, dorsal, volar; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 3 in Scopus, 0 in CINAHL, 8 in Cochrane Library, 7,710 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Physical Methods/Rehabilitation
EXERCISE
Exercise is not generally indicated acutely. For those with residual deficits, particularly post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

Evidence for the Use of Exercise
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: exercise, exercising, physical activity; ganglion cyst, wrist, hand, ganglion, ganglia, dorsal, volar; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 1 articles in PubMed, 5 in Scopus, 0 in CINAHL, 9 in Cochrane Library, 15,300 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Hand Arm Vibration Syndrome (HAVS)
Special Studies and Diagnostic and Treatment Considerations
Currently there is no “gold standard” for the diagnosis and staging of hand-arm vibration syndrome (HAVS). Most authorities have adopted the Stockholm workshop scale(1351, 1352) which is subjective and relies on patient recall.(1353) This subjective system presents problems with reliability, particularly from patients pursuing compensation claims, which has been demonstrated in at least one study of persons reporting HAVS-related Raynaud’s phenomenon and submitting photographs of their hands during an active episode for review. Approximately 50% of the study population that reported to have captured their episode did not have supporting photographic evidence for what they were reporting.(1354)

In the pursuit of objective testing, there are a number of reported physical methods that attempt to provide measurable physiologic changes to support the diagnosis of HAVS. For measurement of vascular changes, the cold provocation test (CPT) has long been a cited maneuver. CPT is conducted by
immersing the hands in water at 10° C-15° C for 10 minutes, and comparing skin temperature recovery at 5 and 10 minutes with baseline prior to the cold water bath. The observer also looks for signs of blanching or white finger. There are several variations of this technique, which include adding more sophisticated temperature measurement instruments for measurement of finger skin temperature (FST) changes, or thermographic studies such as with infrared and dynamic infrared imaging. Finger systolic blood pressure (FSBP) measurement has also been described. Each of these tests attempts to reproduce or measure vascular changes associated with cooling.(1355-1359) Neurological testing has also been described through various methods. Most include measurement of sensory and motor functions, rather than nerve conduction or EMG studies. These tests include vibrotactile threshold tests, thermal aesthesiometry, grip strength, and dexterity testing.

Recommendation: Cold Provocation Test, Cold Stress Thermography (Finger Skin Temperature, Infrared, Dynamic Infrared, Laser Doppler Imaging), Finger Systolic Blood Pressure, Vibrotactile Threshold Testing, Thermal Aesthesiometry, or Nerve Conduction Velocity Studies to Diagnose Hand Arm Vibration Syndrome

There is no recommendation for or against the use of a cold provocation test, cold stress thermography (finger skin temperature, infrared, dynamic infrared, laser Doppler imaging), finger systolic blood pressure, vibrotactile threshold testing, thermal aesthesiometry, or nerve conduction velocity studies to diagnose hand arm vibration syndrome (HAVS).

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendations
Despite the widespread acceptance of physiologic testing, there are no quality RCTs comparing the utility of diagnostic methods for HAVS. Furthermore, there is poor correlation of these various physiological tests with the Stockholm workshop scales,(1360) and a general inability of these tests to reliably differentiate HAVS from controls.(1361, 1362)

A recent review of the literature concluded that there does not appear to be any single test with satisfactory diagnostic capability in diagnosing HAVS (white finger), but supports the use of cold provocation testing (CPT) as reasonable.(1363) However, a large scale review of cold provocation testing in over 40,000 UK miners being evaluated for compensation claims found only slight correlation of self-reported clinical severity and CPT results, concluding that CPT should not be used for evaluating the vascular component of HAVS.(1364) There remains no established standard for CPT methodology, which makes interpretation and comparisons difficult. While the test is relatively benign and inexpensive, the results are of unknown diagnostic utility.

There is little information available supporting the utility of thermographic imaging. Most of the reports are of small populations. The most recent study (21 patients) concluded that none of the available methods is sufficient for arterial constriction testing, but may be useful in follow-up testing of individuals.(1365) A similar story exists for finger systolic blood pressure monitoring as a diagnostic test. A recent prospective study measuring the changes in finger systolic blood pressure (FSBP) after segmental local cooling for vibration-induced white finger in vibration exposed vs. non-exposed populations showed a significant decrease in FSBP in the exposed group with reported HAVS vs. non-exposed as well as the exposed with no history of HAVS. The sensitivity and specificity of the FSBP test with a cut-off value of 75% of normal at 23 +/- 1 degrees C, were 65.2 and 87.5%, respectively, and at 21 +/- 1 degrees C, they were 73.9 and 82.5%, respectively.(1366) However, the study used self-report of HAVS and included retired (no longer exposed) persons in the exposed with HAVS group.

Testing for neurological deficits may be slightly more beneficial than vascular testing for confirming the severity of nerve damage associated with HAVS, although they are not definitive in objectively identifying HAVS. In a follow-up report of UK miners being evaluated for HAVS claims, 57,000 persons evaluated with vibrotactile threshold testing and thermal aesthesiometry showed some evidence that these tests are reliable indicators of underlying neurological damage.(1367)
Thus, there is insufficient evidence for making evidence based recommendations on the utility of each of the various tests currently available for the vascular and neurological components of HAVS. Administering a combination of these tests may improve the diagnostic utility when considered in context of the medical history and occupational exposures. Nerve conduction studies may also be indicated to rule out other associated or concomitant upper extremity disorders, although are not likely of useful benefit for diagnosis of HAVS. In addition to neurovascular physiologic testing, there are limited reports of serologic testing for HAVS.

**Evidence for the Use of Diagnostic Testing**

There are 3 moderate-quality studies incorporated into this analysis.\(^{1471, 1478, 1479}\) (Coughlin 01a; Coughlin 01b; Poole 04) There are 4 low-quality studies in Appendix 2.\(^{1480-1483}\) (Lindsell 99; Kurozawa 91; Bogadi-Sare 94; Lawson 97)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Hand-Arm Vibration Syndrome, Vibration white finger, dead finger, white fingers, hand-transmitted vibration, hand-arm vibration, traumatic vasospastic disease, Cold provocation, cold stress thermography, finger systolic blood pressure, vibrotactile threshold testing, thermal aesthesiometry, never conduction velocity, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 0 articles in PubMed, 2 in Scopus, 0 in CINAHL, 16 Cochrane Library, and 120 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 4 from Google Scholar, and 5 from other sources. Of the 9 articles considered for inclusion 7 diagnostic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score</th>
<th>N</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>Type of Thermography</th>
<th>CT used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes assessed</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coughlin 2001</td>
<td>Same as OCC MED Case Control</td>
<td>5</td>
<td>31 subjects in two groups. Group A: 10 healthy volunteers. 5 men, 5 women. Median age of 35. Group B: 21 patients. 20 men, 1 woman. Median age of 45</td>
<td>Hand</td>
<td>HAVS with RP</td>
<td>Cold Provocation</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>After cold provocation, the finger temperature and time for the finger temperature to return to pre-cooling levels were able to distinguish the HAVS group and the normal group. The sensitivity of CPT was low after cooling, but “CPT has a good sensitivity, specificity, positive predictive value and negative predictive value; it strongly supports a diagnosis of digital vasospasm.”</td>
<td>Data suggest CPT test has good sensitivity and specificity and supports a diagnosis of digital vasospasm.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Author/Year</td>
<td>Score</td>
<td>Study Design</td>
<td>Population/Case Definition</td>
<td>Investigative Test</td>
<td>Gold Standard / Comparative Test</td>
<td>Results</td>
<td>Conclusion</td>
<td>Comments</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------</td>
<td>-------</td>
<td>--------------</td>
<td>-----------------------------</td>
<td>--------------------</td>
<td>----------------------------------</td>
<td>---------</td>
<td>------------</td>
<td>----------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Poole 2004</td>
<td>6.0</td>
<td>Case Control</td>
<td>N = 46 24 Males with HAVS VS 22 Males without HAVS (Control) Mean age = 46</td>
<td>Measuring FSBP after cold provocation at 30, 15 and 10°C FST measurement following immersion of hands in 15°C water for 5 min</td>
<td>FSBP on the middle finger yielded a sensitivity of 60%, specificity of 84.1%, PPV of 71.5%, and a NPV of 75.9%. Compared to FSBP, FST had results of 68%, 71%, 61%, and 77%, respectively.</td>
<td>“Based on our data, the FSBP may also have limited use in confirming a positive diagnosis of vibration-induced vascular problems.”</td>
<td>Data suggest FSBP is of limited value as a diagnostic test for HAVS although it may have value in ruling out and/or confirm the vascular component of HAVS.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coughlin 2001 OCC MED</td>
<td>5.5</td>
<td>Case Control</td>
<td>N = 50 participants Two-Point discrimination</td>
<td>Depth sense perception</td>
<td>When testing using DSP, there was no significant difference in reach up to 95% 3 min after rewarming. The accuracy of the test was also the greatest towards the last stages of rewarming. The specificity and PPV were high during precooling stages and remained relatively high during the rewarming stages. NPV was low during the precooling stage and became high (&gt;90) during the rewarming stages.</td>
<td>“The increased sensitivity of the TPD disc would”</td>
<td>Data suggests the 2 point disc providers increased sensitivity for the assessment of</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
20 with HAVS VS 15 Sedentary worker VS 15 manual workers

No mention of mean age.

the right hand of all three groups. The left hand was significantly poorer in the HAVS group than the two others. DSP has a sensitivity of 41, specificity of 94, PPV of 82 and NPV of 70. When testing with TPD, both hands were significantly poorer in the HAVS group than the two other groups. TPD has a sensitivity of 46, specificity of 94, PPV of 84, and NPV of 72.

suggest that it should be used in preference to the DSP disc for the assessment of sensorineural dysfunction in patients with HAVS.

HAVS vs. the depth sense disc.

1. **Recommendation: Serologic Tests (Thrombomodulin, Soluble Intracellular Adhesion Molecule 1 [s1-CAM 1]) to Diagnose Hand Arm Vibration Syndrome**

   Serologic tests – thrombomodulin, soluble intracellular adhesion molecule 1 (s1-CAM 1) – are not recommended to diagnose HAVS.

   **Strength of Evidence** – **Not Recommended, Insufficient Evidence (I)**  
   **Level of Confidence** – **Low**

2. **Recommendation: Testing for Connective Tissue Disorders to Diagnose Hand Arm Vibration Syndrome**

   There is no recommendation for or against the use of testing for connective tissue disorders to diagnose HAVS.

   **Strength of Evidence** – **No Recommendation, Insufficient Evidence (I)**  
   **Level of Confidence** – **Low**

**Rationale for Recommendations**

There are no quality randomized studies on the utility of serologic testing or connective tissue disorders testing for HAVS. There does not appear to be any serologic tests that currently provide objective evidence or staging of HAVS. Objective serum tests, such as levels of soluble thrombomodulin (sTM) and soluble intercellular adhesion molecule-1 (sICAM-1), may provide some utility in the future as they have been shown to be statistically different in exposed groups with HAVS symptoms, but the usefulness is hampered currently by the lack of clear reference ranges,(1353) as each of the measurements for both comparison groups were still in the range considered normal. Testing for other causes of Raynaud’s phenomenon, particularly connective tissue disorders such as scleroderma and systemic lupus erythematosus may be beneficial when occupational exposure histories are not consistent with clinical presentation and the threshold for such testing should be low.

**Evidence for the Use of Serologic Testing or Connective Tissue Disorders Testing**
There is 1 moderate-quality study incorporated into this analysis. (Kanazuka 96) There is 1 low quality study in Appendix 2. (Kennedy 99)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Hand-Arm Vibration Syndrome, Vibration white finger, dead finger, white fingers, hand-transmitted vibration, hand-arm vibration, traumatic vasospastic disease, Cold provocation, cold stress thermography, finger systolic blood pressure, vibrotactile threshold testing, thermal aesthesiometry, never conduction velocity, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 0 articles in PubMed, 0 in Scopus, 4 in CINAHL, 9 Cochrane Library, and 150 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 from Google Scholar, and 2 from other sources. Of the 3 articles considered for inclusion 3 diagnostic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Study Design</th>
<th>Population/ Case Definition</th>
<th>Investigative Test</th>
<th>Gold Standard / Comparative Test</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kanazuka 1996</td>
<td>4.0</td>
<td>Case Control</td>
<td>N=175 Males 100 Patients with HAVS (Mean age = 63.0±6.3) Vs 25 Patients with collagen disease (Mean age = 43.5±16.8) Vs 50 Healthy patients (Mean age = 56.8±7.8)</td>
<td>TM one-step sandwich enzyme immunoassay</td>
<td>Not mentioned</td>
<td>Patients with HAVS had a significantly higher level of plasma TM (3.32±1.11 ng/mL) than the normal control (2.49±1.05 ng/mL, p&lt;0.0001). There was no significant difference between the HAVS group and the collagen disease group (3.65±2.02 ng/mL, p&lt;0.01).</td>
<td>“[W]e suggest that endothelial injury is present in vibration syndrome, the degree of endothelial injury in vibration syndrome equals that in collagen disease, and the endothelial injury in chain-saw operators is greater than that in rock-drill operators.”</td>
<td>Data suggest endothelial injury exists in patients with VWF as well as collagen disease.</td>
</tr>
</tbody>
</table>

**Initial Care**

There are no quality randomized clinical studies for the treatment of physiologic manifestations associated with HAVS. The most prudent form of treatment is to first remove or reduce the exposure to vibration, particularly in the earlier stages of symptom presentation. Smoking has been identified as a
risk factor for HAVS.\(^{1368}\) By inference, smoking cessation is a frequent recommendation to patients with HAVS. The effects of smoking on HAVS, if any, are thought to be a result of chronic platelet function inhibition,\(^{1369}\) effects on the microvasculature and that of nicotine on smooth muscle function. However, there is no quality evidence that smoking cessation will affect the course. As a risk factor, smoking cessation is recommended.

Other common advice based on the proposed pathophysiology of vasospasm includes avoidance of beta-blockers, sympathetic stimulants including caffeine, decongestants, amphetamines and even cocaine as they may act as potential triggers. Further, maintenance of hand and body temperature in cold environments may help avoid or reduce the risk of symptoms.

**Follow-up Visits**

There are no recommendations in the literature for the frequency or duration of follow-up visits. A few follow-up appointments to evaluate results of non-invasive measures, exposure avoidance, and serologic testing are indicated.

**Prescription Medications**

There are no quality studies of medications that prevent or improve symptoms related to HAVS. As the vascular component of HAVS mimics other causes of Raynaud’s phenomenon, calcium channel antagonists, which have positive benefit for many with non-vibration related cases, are often prescribed for HAVS.

**CALCIUM CHANNEL BLOCKERS**

*Recommendation: Calcium Channel Blockers for Advanced Subacute or Chronic Hand Arm Vibration Syndrome*

*Use of calcium channel blockers (nifedipine) for treatment of vascular symptoms similar to Raynaud’s phenomenon is recommended for advanced subacute or chronic HAVS.*

*Indications* – Patients with HAVS. Generally used in patients with sufficient symptoms such that removal from exposure is insufficient for management.

*Frequency/Dose* – Per manufacturer’s recommendations; generally initiated with low dose. Blood pressure should be monitored and may require lower doses, especially among those without higher blood pressures or among those with adverse effects.

*Discontinuation* – Resolution, intolerance, adverse effects.

---

**Strength of Evidence** – Recommended, Insufficient Evidence (I)

**Level of Confidence** – Low

**Rationale for Recommendation**

There is no quality evidence for the use of calcium channel blockers in HAVS population. It is a commonly accepted treatment for Raynaud’s phenomenon associated with connective tissue diseases with moderate benefit. A review of all calcium channel antagonist trials for non-HAVS Raynaud’s is beyond the scope of this text. Rather, as this medication is already frequently used for advanced HAVS, and with the lack of other treatments available, it may be considered a treatment for symptomatic patients once exposure sources are reduced or eliminated and other personal health measures have started.

**Evidence for the Use of Calcium Channel Blockers**

There are no quality studies incorporated into this analysis.

---

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: calcium channel blockers, hand arm vibration syndrome, vibration white finger, dead finger, white fingers, hand-transmitted vibration, hand-arm vibration, traumatic vasospastic disease; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization,
randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 19 articles in PubMed, 0 in Scopus, CINAHL, and Cochrane Library, 152 from Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, CINAHL, Cochrane Library, Google Scholar, and from other sources. Of the 1 articles considered for inclusion, 0 randomized trials and 1 systematic studies/background met the inclusion criteria.

**Physical Methods/Rehabilitation**

**EXERCISE**

Exercise is not generally indicated.

**Evidence for the Use of Exercise**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: exercise, exercising, physical activity, Hand-Arm Vibration Syndrome, vibration white finger, dead finger, white fingers, hand-transmitted vibration, hand-arm vibration, traumatic vasospastic disease; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 5 articles in PubMed, 2 in Scopus, 0 in CINAHL, 14 in Cochrane Library, 1,158 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Laceration Management**

**Special Studies and Diagnostic and Treatment Considerations**

There are no quality studies on diagnostic testing for the evaluation of wounds with lacerations. However, among the minority of wounds of sufficient severity, the use of imaging to rule out traumatic injury to bone or other structures is generally considered effective and well established. Yet, detection of retained soft tissue foreign bodies remains a clinical dilemma, with one study reporting up to 38% of foreign bodies in hand wounds going undetected by the initial provider, resulting in the second-leading cause of lawsuits in emergency medicine. Furthermore, if nerve injury is detected or suspected, then EDS may be indicated 2 to 3 weeks post-injury. An immediate EDS is not recommended as Walerian degeneration will not have been completed until at least 2 weeks post-injury, making earlier studies falsely normal.

**X-RAYS**

**Recommendation: X-rays for Evaluation of Lacerations with Suspected Fracture or Foreign Body**

X-rays are recommended for the evaluation of traumatic injury resulting in skin lacerations to rule out fracture or if a radiopaque foreign body is suspected.

- **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
- **Level of Confidence** – **Moderate**

**Evidence for the Use of X-ray**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Laceration management, x-ray, xray, radiography, lacerations with suspected fracture, foreign bodies, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 24 articles in PubMed, 20 in Scopus, 0 in CINAHL, 0 in Cochrane Library, and 1,880 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.
ULTRASOUND
Recommendation: Ultrasound for Evaluation of Suspected Superficial Foreign Bodies

Ultrasound is recommended for evaluating suspected radiolucent materials or as an alternative test when radiopaque foreign body is suspected but not detected on x-ray images.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Evidence for the Use of Ultrasound
There are 4 quality studies incorporated into this analysis. (1489-1492) (Soubeyrand 08; Tahmasebi 14; Wu 12; Fornage 86)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ultrasound, Laceration Management, Suspected superficial foreign bodies, ultrasonography, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 122 articles in PubMed, 62 in Scopus, 0 in CINAHL, 0 in Cochrane Library, and 8,560 from Google Scholar. We considered for inclusion 0 from PubMed, 2 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 2 from Google Scholar, and 0 from other sources. Of the 5 articles considered for inclusion 4 diagnostic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Study Type</th>
<th>Author/Year</th>
<th>Score</th>
<th>N</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>Type of Ultrasound</th>
<th>CT used</th>
<th>MRI used</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes</th>
<th>Long term follow-up</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diagnostic</td>
<td>Soubyrand 2008</td>
<td>7.5</td>
<td>30 injuries in 26 patients (19 males, 7 females) Mean age: 34 years</td>
<td>Hand and Wrist</td>
<td>Laceration Management /Lesion</td>
<td>Doppler Ultrasound</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>More than one rater</td>
<td>-</td>
<td>72 hours</td>
<td>There were 20 injuries of the finger and 10 of the palm. The right side was involved in 17 of 30 injuries (57%) and the dominant hand was involved in 11 of 30 injuries (37%). Injury at home occurred in 18 cases and at work in 10 cases. Two patients were injured on the street. Penetrating object was glass in 17 injuries, knife in 7 injuries, metallic object in 2, human teeth in 1, and a stone in 1. A complete US examination was performed in all 30 cases, despite moderate pain in two cases. Of 98 examined tendons, 81 appeared intact and 17 were damaged. Of 81 examined nerves, 63 appeared intact and 18 were damaged. Of 75 examined arteries, 61 appeared intact and 14 were damaged. The lesion path was visualized in 22 of the 30 injuries. In five injuries, the path did not extend beyond the fascial layer (superficial injury), and in two injuries, the path ended in the muscle. Foreign bodies were visualized in five injuries.</td>
<td>&quot;In conclusion, US proved highly effective in detecting tendon and arterial lesions. The results were less reliable regarding nerve damage. US may be effective in identifying hand lesions that require surgical repair and in selecting patients who can be treated without surgical exploration, provided they undergo a second physical examination 72 hours after the injury. Further studies in larger numbers of patients are needed to evaluate this possibility.&quot;</td>
<td>Data suggest US is effective in the detection of volar injuries without tendon or arterial lesions but not as good for detection of nerve lesions.</td>
</tr>
<tr>
<td>N=51 patients (41 males, 10 females)</td>
<td>Mean age: 24.95±1 3.4 years</td>
<td>Laceration Management USG</td>
<td>Predominant chief complaints of the patients were: foreign body sensation in 24, discharging wound in 15, and pain in 12 cases. Ten cases had a history of surgical exploration without the use of USG examination, which had no foreign body detected. On USG scan, 100% of the foreign bodies were echogenic. USG revealed a foreign body in 50 patients. All patients underwent surgical exploration or USG-guided removal. Forty-six patients had a foreign body removed. One patient had a negative USG exam and surgical exploring revealed a 7-mm thorn. USG was falsely positive in three cases with failed surgical manipulation due to the presence of air bubbles and scar tissue, as well in as one case with calcified granuloma. Foreign bodies were thorn, wood, glass, and plastic. The sites of the foreign bodies were foot, hand, leg, arm, forearm, ankle, wrist, knee, and thigh. Sizes of foreign body varied from 4-51 mm and in 50% of cases, the size of the foreign body was greater than 13 mm.</td>
<td>Predominant chief complaints of the patients were: foreign body sensation in 24, discharging wound in 15, and pain in 12 cases. Ten cases had a history of surgical exploration without the use of USG examination, which had no foreign body detected. On USG scan, 100% of the foreign bodies were echogenic. USG revealed a foreign body in 50 patients. All patients underwent surgical exploration or USG-guided removal. Forty-six patients had a foreign body removed. One patient had a negative USG exam and surgical exploring revealed a 7-mm thorn. USG was falsely positive in three cases with failed surgical manipulation due to the presence of air bubbles and scar tissue, as well in as one case with calcified granuloma. Foreign bodies were thorn, wood, glass, and plastic. The sites of the foreign bodies were foot, hand, leg, arm, forearm, ankle, wrist, knee, and thigh. Sizes of foreign body varied from 4-51 mm and in 50% of cases, the size of the foreign body was greater than 13 mm.</td>
<td>Data suggest US can detect radiolucent-soft-tissue foreign bodies that radiographs can not.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Thirty-four patients were enrolled in this study. There were 6 finger injuries, 11 hand injuries, 6 forearm injuries, 6 arm injuries, and 5 lower extremity injuries. Based on MRI or direct wound exploration, 4 patients had partial tendon injuries, 9 patients had complete tendon injury, and 21 patients had no evidence of tendon injury noted. Bedside ultrasound was able to accurately diagnose the extent of tendon injury in 33 of the 34 total cases. In comparison, physical examination accurately diagnosed 29 of the 34 total cases. On average, time to diagnosis and disposition based on bedside ultrasound findings was 46.3 minutes. In contrast, overall time to wound exploration, MRI, or consultation was 138.6 minutes.

“Bedside ultrasound is more sensitive and specific than physical examination alone for detecting tendon lacerations and takes less time to perform than traditional wound exploration techniques or MRI. Data obtained from bedside ultrasonography can be used to improve diagnostic accuracy and expedite patient care.”

No mention of gender or mean age. Data suggest bedside US increases the sensitivity/specificty in detection of tendon injuries when compared to physical exam alone.
Eight foreign bodies were found at surgery; glass in 4 cases, metal in 3 cases, and vegetable material in 1 case. All foreign bodies were visualized as hyperechoic on sonograms. An acoustic shadow was present in 2 cases only (glass fragments). A hyperechoic comet-tail artifact secondary to reverberations inside the dense echogenic foreign body was visualized in 3 cases. In 7 cases a surrounding hypoechoic mass ranged from 1.2-3 cm in diameter correlated well with inflammatory changes found at surgery. Seven of the eight foreign bodies were glass or metallic fragments and were radiopaque with sizes of 0.1-1 cm. In 1 case a vegetable fragment responsible for a cyst could not be seen on the radiograph, but was demonstrated on sonograms.

CT Recommendation: CT for Evaluation of Suspected Superficial Foreign Bodies

There is no recommendation for or against the use of CT for suspected superficial foreign bodies. CT is not routinely recommended, but may be indicated for the evaluation of suspected radiolucent materials and as an alternative test when radiopaque foreign body is suspected but is not detected on x-ray images or ultrasound.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendations

Most work-related lacerations presenting to clinics are too superficial to involve the bone or joints. However, if the injury mechanism or location of injury suggests a possibility of fracture, x-rays are indicated (see specific fracture sections for further recommendations). There are no quality studies of imaging techniques for the evaluation of suspected foreign bodies. If a foreign body is suspected, additional diagnostic testing should be considered dependent on the suspected foreign body type. For suspected radiopaque substances such as metals or glass, traditional x-ray reliably detects the foreign body 80 to 95% of the time. However, x-ray images do not reliably detect radiolucent foreign bodies such as wood, plastic, or vegetative material.

Ultrasound is increasingly being utilized for the evaluation of suspected radiolucent foreign bodies, although there are no quality studies available. There are several case series and cadaver studies.
providing reports of high sensitivity, although there are also a small number of false positives related to tendons or other artifacts. CT has reported high sensitivity for radiopaque substances, and moderate sensitivity for radiolucent materials. Because of increased costs, higher radiation exposure, with intermediate sensitivity, CT may be best used when a foreign body is suspected but not detected by x-rays or ultrasound. MRI is not indicated for evaluation of metallic foreign bodies in particular.

Evidence for the Use of CT
There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Laceration, Foreign, CT, CAT, Computerized Tomography, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 60 articles in PubMed, 12 in Scopus, 0 in CINAHL, 63 Cochrane Library, and 4680 from Google Scholar. Zero articles met the inclusion criteria.

Treatment
As previously stated, optimal results are accomplished by preventing infection through thorough wound cleansing, approximating wound edges with appropriate closure techniques, and providing a clean, moist environment to accelerate wound healing. If nerve injury is detected or suspected then appropriate surgical consultation should be considered.

WOUND PREPARATION
Infection is one of the primary factors that interfere with wound healing. Contamination of the wound from inoculation of skin flora or environmental bacteria, foreign bodies such as gravel, vegetation, dirt, and other industrial related compounds can act as a nidus for wound infection. Adequate wound anesthesia may be required for wound preparation. Anesthetic technique is reviewed later in this section.

1. Recommendation: Wound Cleansing, Irrigation, and Debridement
   Meticulous wound preparation after appropriate anesthesia using saline irrigation or copious amounts of running tap water, scrubbing, and debridement of devitalized tissue is recommended.
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – High

2. Recommendation: Wound Irrigation with Sterile Saline or Tap Water
   The use of either sterile saline or tap water is recommended for an irrigating solution.
   Strength of Evidence – Recommended, Evidence (C)
   Level of Confidence – High

3. Recommendation: Sterile or Clean Glove Use During Wound Cleaning
   The use of either sterile or clean gloves during wound cleaning is recommended.
   Strength of Evidence – Recommended, Evidence (C)
   Level of Confidence – High

Rationale for Recommendations
Wounds become infected when they contain more than 10⁵ bacteria per gram of tissue.(1376) As there is no test to determine the immediate bacterial load of a particular laceration, it seems prudent that all wounds should undergo some form of cleansing to decrease the amount of soil or presence of small foreign bodies to reduce the inoculation of bacteria and prevent infection. There are no quality studies comparing infection rates in wounds that are irrigated vs. non-irrigated or cleansed. However, it is widely accepted that irrigation and cleansing are best practice. Therefore, although there is a lack of quality evidence, wound cleansing, irrigation, and debridement are recommended. Optimal irrigating solutions and techniques are more controversial.
There is moderate evidence that irrigation solution does not make a significant difference in infection rates of routine laceration management. A high-quality study comparing tap water to sterile saline in a pediatric population showed no difference in infection rates at 48 hours.\(^{(1377)}\) Another moderate-quality study of 715 lacerations randomized to irrigation under regular tap water vs. sterile saline using pressure syringe irrigation also found no significant difference in infection rates.\(^{(1376)}\) Patients enrolled in tap water irrigation were instructed to wash their wound under regular tap water (U.S. location) for a minimum of 2 minutes. However, the power of these studies to detect differences in infection rates may be too low to conclude inferiority. On balance, there appears to be no difference in infection rates between the common practice of using sterile saline for routine laceration repair or using regular tap water for uncomplicated extremity lacerations and either wound irrigation with sterile saline or tap water is recommended. There is no quality evidence supporting the use of concentrated povidine-iodine solution instillation into the wound, although a low-quality study suggests some benefit in reduced infection rates.\(^{(1378)}\) Another low-quality study found no difference in infection rates between normal saline, povidine, and Shur Clens®.\(^{(1379)}\) There is some concern that concentrated povidine-iodine, hydrogen peroxide, and detergents may cause tissue toxicity.\(^{(195)}\)

There are no quality studies on irrigation pressures. High-pressure irrigation may result in increased trauma.\(^{(195)}\) Optimal pressures of 5 to 8 psi generated by large syringe and 16- to 19-gauge needle have been recommended.\(^{(195)}\) One moderate-quality study compared a commercial pressurized canister irrigation system with a standard syringe and 20-gauge catheter at maximal plunger force using saline and benzalkonium chloride.\(^{(1380)}\) The study had weaknesses but found no difference in infection rates or soft tissue trauma between the groups. The only advantage was that irrigation times were shorter (3.9 versus 7.3 minutes) using the canister.

For lacerations that involve skin areas where significant hair may hamper closure efforts, removal by clipping rather than shaving is commonly suggested to reduce potential sources of contamination resultant from disturbing bacteria on hair shafts, although there is no evidence to support this method in routine laceration repair. Debridement of devitalized tissue through surgical excision and scrubbing may also reduce the risk of infection. Generally, sterile technique has been recommended. However, there is one large moderate-quality study of 816 lacerations that showed no difference in infection rates in repair using sterile gloves versus non-sterile clean gloves,\(^{(1381)}\) thus either is recommended.

**Evidence for Wound Preparation**

There is 1 high-\(^{(1377)}\) and 3 moderate-quality\(^{(1376, 1380, 1381)}\) RCTs incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: wound preparation, wound cleansing, irrigation, debridement, wound healing, laceration, wound, cuts, management, repair, care, upper extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 4 articles in PubMed, 0 in Scopus, 15 in CINAHL, 5 in Cochrane Library, 8321 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 5 from Google Scholar, and 0 from other sources. Of the 4 articles considered for inclusion, 4 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bansal 2002</td>
<td>RCT</td>
<td>9.0</td>
<td>N = 46 (17 female and 28 male) with simple lacerations. Age range 2-15.</td>
<td>Wound irrigation by high pressure (25-40 PSI) syringe using tap water (N = 21) vs Normal sterile saline (N = 24). Follow-up for 48 hours.</td>
<td>Post irrigation culture positive in 11/21 (52%) for tap water, 7/24 for sterile saline (29%) p = 0.20. No difference in infection rates at 48 hours.</td>
<td>“Our study suggests that tap water may serve as a cost-saving alternative to normal saline for irrigating simple lacerations before repair.”</td>
<td>Hand lacerations were excluded. Pediatric population.</td>
</tr>
<tr>
<td>Moscati 2007</td>
<td>RCT</td>
<td>7.5</td>
<td>N = 715 with acute simple lacerations requiring sutures or staples. Age and gender not specified.</td>
<td>Tap water irrigation at sink (N = 300) vs High pressure sterile saline (N = 334). Follow-up for 48 hours.</td>
<td>11/374 in saline group developed infection (3.3%) vs. 12/339 (4.0%) with no significant difference between the groups.</td>
<td>“Compared with sterile saline, tap water for wound irrigation is more cost-effective and appears to be equally safe and efficacious.”</td>
<td>Sixty percent of enrolled lacerations were of upper extremity. Baseline comparability of common variables not presented. Author estimates total US savings $65.6 million by using tap water irrigation vs. current practice.</td>
</tr>
<tr>
<td>Chisholm 1992</td>
<td>RCT</td>
<td>5.0</td>
<td>N = 542 (male to female ratio 1.8:1 and 2.7:1 in Canister group) with lacerations requiring closure. Mean age for Syringe and Canister groups: 24.9 and 23.8 years.</td>
<td>220mL canister of sterile NS with 0.006% benzalkonium chloride (N = unknown) vs NS irrigation using 30-mL syringe, 20-gauge IV catheter tip 1 in. above skin edge, depress syringe plunger with maximal force (N = unknown). Follow-up</td>
<td>Face and hands most frequently lacerated. Mean irrigation time for pressurized canister group (281) 3.9 vs. 7.3 minutes for syringe irrigation group (254) (p &lt;0.0001). Wound complications occurred in 8/221 (3.6%) in syringe irrigation group and 12/245 (5.0%) in pressurized canister group, (p = 0.50).</td>
<td>“There was no significant difference in infection rates between the two groups. The pressurized canister group’s wounds were cleansed in almost half the time of those in the syringe group.”</td>
<td>Lack of control for dressing type, use of topical antibiotics. Final wound observations made by multiple observers including patient self-report office based practitioners, and ED practitioners.</td>
</tr>
</tbody>
</table>

### Wound Irrigation: Syringe Irrigation vs Pressurized Canister

Sterile vs Nonsterile Gloves for Uncomplicated Lacerations
### WOUND ANESTHESIA

Wound anesthesia is commonly obtained after completing a sensory examination through local infiltration, digital nerve block or topical application of anesthetic preparations. Anesthetic technique is most commonly performed based on wound location and the preference of the treating health care professional.

1. **Recommendation: Local Infiltration plus Topical Anesthetic or Digital Block for Finger Laceration Repair**

   Adequate anesthesia by either topical anesthetic plus local infiltration or digital block is moderately recommended for finger laceration repair. There is no recommendation of one technique over the other. For distal finger lacerations, digital block may be substantially less painful than local infiltration performed without topical anesthetic. If the operator and patient preference is digital block, the various techniques are described and evaluated in the management of phalangeal fracture section in this guideline.

   Strength of Evidence – Moderately Recommended, Evidence (B)
   Level of Confidence – Moderate

2. **Recommendation: Local Infiltration for Extremity Wound Repair**

   Instillation of local anesthetic for extremity wounds after sensory testing is recommended as the first-line technique for most laceration repairs unless the size or complexity would require potentially toxic doses of local anesthetic. Local anesthetic with epinephrine (except digits) is recommended.

   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Low

3. **Recommendation: Topical Anesthetics for Lacerations**

   The use of topical anesthetics, Tetracaine-Adrenaline-Cocaine (TAC) and EMLA, are recommended as an alternative to local infiltration for lacerations of the extremities (excluding digits) or as pre-treatment to reduce pain related to needle infiltration. However, these anesthetics have longer times to onset of effective anesthesia.

   Strength of Evidence – Recommended, Evidence (C)
   Level of Confidence – Low

**Rationale for Recommendations**

There are numerous quality studies of different anesthetic techniques for management of laceration repairs of the finger. There is one high-quality and one moderate-quality study comparing local infiltration to digital block for finger lacerations. However, in the high-quality study, both received topical anesthesia which may have otherwise confounded the results. The authors found no difference in pain of providing...
anesthesia or quality of anesthesia between the two techniques.

Digital anesthesia was preferred by providers and patients for both the application and quality of anesthesia in a moderate quality study, although it was uncertain if the comparison groups had similar baseline pain. Although there may be a modest advantage to digital anesthesia, there is not enough evidence to support one technique over the other, and both are recommended based on operator and patient preference.

There is one quality study that compared topical anesthetics with placebo, and that trial demonstrated efficacy, although it is a remote study utilizing Tetracaine-Adrenaline-Cocaine (TAC) and topical lidocaine. However, there are many trials comparing different topical agents. Topical anesthetics are applied to provide analgesia for subsequent local infiltration, or to provide anesthesia for wound repair. Topical anesthetics used for laceration repair without local infiltration are best used in highly vascularized regions, although they have also been used successfully in the extremity. In the past, TAC has been used effectively, but concerns of toxicity have resulted in the development and use of non-cocaine containing products. Lidocaine-Adrenaline-Tetracaine (LAT, LET) and EMLA are now the primary topical anesthetics used in the United States. LAT has been shown to be more effective than TAC in one high-quality study for topical anesthesia and as effective in another high-quality study for topical pre-treatment for infiltration. EMLA was also shown to be more effective for topical anesthesia than TAC in a moderate quality study.

There is one high-quality study comparing EMLA and LAT for topical anesthesia that demonstrated equal efficacy, with a slight advantage to LAT in the time to achieving anesthesia. Thus, there is sufficient evidence to support the use of LAT and EMLA for pretreatment and for primary anesthesia in select wounds in adult populations.

Although local infiltration is the most common technique, there are no quality studies of local anesthetic infiltration versus placebo. Nor are there any quality studies comparing topical anesthetics to local infiltration or nerve blocks. As local infiltration is the gold standard for most wound repair, and the failure of topical anesthetics is treated by local infiltration or nerve block in complicated wounds, there is no recommendation for the use of topical anesthetics over local infiltration.

There is one high-quality study comparing lidocaine solutions with buffering, the addition of epinephrine, and the use of diphenhydramine as an alternative for upper extremity wounds. Lidocaine with epinephrine with or without buffering was preferred by patients over diphenhydramine or buffered solutions without epinephrine. This result contradicts with common anecdote of using buffered solutions to reduce injection pain.

**Evidence for Wound Anesthesia**

There are 5 high-(1382, 1387, 1388, 1390, 1391) and 5 moderate-quality(1383-1386, 1389) RCTs incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: anesthesia, wound healing, laceration, wound, cuts, management, repair, care, upper extremity, local infiltration plus topical anesthetic; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 76 articles in PubMed, 39 in Scopus, 3 in CINAHL, 3 in Cochrane Library, 4524 in Google Scholar, and 5 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 5 Google Scholar, and 5 from other sources. Of the 10 articles considered for inclusion, 10 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chale 2006</td>
<td>9.0</td>
<td>N = 55 (16 female and 39 male) with traumatic lacerations of 1 finger. Age 40.1 (19.3) digital group: 36.3 (14.0) topical group.</td>
<td>Digital block 1 to 2 mL of lidocaine 1% was injected on both sides of the finger (N = 28) vs Local anesthesia 1 to 2 mL of lidocaine 1% was injected (N = 27). Both had topical anesthetics as co-intervention. 15 minute topical application</td>
<td>Wound outcomes: digital vs. local anesthesia: Time until onset of anesthesia in minutes: 7.7 vs. 1.9 p = 0.001. Mean pain of needle insertion in mm: 29.4 vs. 28.1 p = 0.87. Mean pain of anesthetic infiltration in mm 24.9 vs. 22.6, (p = 0.72).</td>
<td>“Digital and local anesthesia of finger lacerations with prior application of LET to all wounds results in similar pain of needle insertion, anesthetic infiltration, and pain of suturing.”</td>
<td>Application of LET to all wounds makes comparison of digital to local needle injection pain difficult in the absence of LET, which is most cases in the U.S.</td>
</tr>
<tr>
<td>Robson 1990</td>
<td>5.0</td>
<td>N = 60 (gender not specified) with lacerations of the digits. Age over 16 years.</td>
<td>Digital block 1 ml of anesthetic was applied (N = 28) vs Local anesthesia 2% plain lignocaine (N = 32). Follow-up unclear.</td>
<td>Assessment by patient and operator for pain related to application of anesthesia and suturing significantly better for digital block compared with local infiltration, (p &lt; 0.01).</td>
<td>“[D]igital block should be considered as the method of choice in all cases of digital lacerations requiring local anesthesia for their repair.”</td>
<td>No baseline comparison data was presented.</td>
</tr>
<tr>
<td>Ernst 1996</td>
<td>10.0</td>
<td>N = 200 (50 female and 130 male) with simple lacerations not involving vascular compromise infection. 18 years of age.</td>
<td>Group A, buffered 1% lidocaine (N = 45) vs Group B, buffered 1% lidocaine with epinephrine (N = 46) vs Group C, 1% lidocaine with epinephrine (N = 47)</td>
<td>“Buffered lidocaine (A) and buffered lidocaine with epinephrine (B) were significantly less painful to inject than was diphenhydramine with epinephrine (D) (p &lt; 0.01 for both the physicians and the patients). Lidocaine with epinephrine (C) was not statistically different from A, B, or D (p &lt; 0.05). For suturing (anesthesia effectiveness), the patients and the physicians</td>
<td>“Although we found buffered lidocaine solutions less painful to inject in this four-agent comparison study, we were unable to detect a statistically significant difference.”</td>
<td></td>
</tr>
</tbody>
</table>

**Digital vs. Local Infiltration**

**Injectable Agents**
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Study Design</th>
<th>Sponsorship/CoI</th>
<th>Patient Characteristics</th>
<th>Intervention</th>
<th>Findings</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ernst 1995</td>
<td>RCT</td>
<td>Sponsored by grant from Louisiana State University Emergency Medicine Residency Grant Fund. No mention of COI.</td>
<td>9.5</td>
<td>N = 95 (23 female and 76 male) with linear lacerations of face or scalp. Mean age LAT/TAC group: 33±11 / 34±13.</td>
<td>LAT or lidocaine – adrenaline-tetracaine (N = 48) vs TAC or tetracaine – adrenaline – cocaine (N = 47). Follow-up for unclear.</td>
<td>LAT found to have fewer painful sutures than TAC (p = 0.036). For physician ratings, difference between LAT vs. TAC groups showing that LAT more effective than TAC during suturing, (p = 0.093). Patient ratings however showed no significant difference in pain scores.</td>
<td>“We found that patients had smaller percentages of sutures causing pain in the LAT group than in the TAC group.”</td>
</tr>
<tr>
<td>Singer 2001</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>9.5</td>
<td>N = 60 (14 female and 44 male) with pretreating lacerations prior to lidocaine injection. Mean age 8.5 years.</td>
<td>EMLA cream (N = 31) vs LET or cream for pretreating lacerations prior to lidocaine injection (N = 29). Anesthetic application times range from 15 to 135 minutes, not other follow up specified.</td>
<td>“51/54 wounds received supplemental injection of lidocaine and were similar in the both groups (92% for LET vs 97% for EMLA, p = 0.47). Wounds treated with LET were more frequently anesthetic to a stick with a 27-gauge needle than wounds treated with EMLA (73% vs 40%, respectively, p = 0.01)... no difference in the median pain of supplemental lidocaine injection between the two groups.”</td>
<td>“[P]retreatment of uncomplicated lacerations ... with LET or EMLA cream results in a similar reduction in the pain of subsequent injection of lidocaine.”</td>
</tr>
<tr>
<td>Schilling 1995</td>
<td>RCT</td>
<td>Sponsored by FA Bean Education and</td>
<td>8.5</td>
<td>N = 171 (51 female and100 male) with uncomplicate d laceration on face or scalp. Mean age TAC/LET Lidocaine, epinephrine, tetracaine (LET) solution (N = 57) vs Tetracaine, adrenaline, cocaine (TAC) solution</td>
<td>“In the TAC and LET groups combined, 116 of the 151 patients (76.8%) received adequate anesthesia before suturing. There was no difference between TAC (79.5%) and LET (74.4%) (p = 0.46). There was no difference between TAC and LET in</td>
<td>“LET is an effective alternative to TAC for topical anesthesia during suturing of uncomplicated lacerations on the face and scalp in children.”</td>
<td>Applicability uncertain as population was pediatric with scalp/facial lacerations. May have had adult parents with needle phobia.</td>
</tr>
</tbody>
</table>

**Topical Agents**
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Participants</th>
<th>Intervention</th>
<th>Outcome</th>
<th>Type of Study</th>
<th>Key Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pryor 1980</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>N = 151 (gender not specified) with lacerations. Age range 1 to &gt;17, mean age 9 years.</td>
<td>Topical TAC (N = unknown) vs topical lidocaine (N = unknown) vs placebo for lacerations &lt;5 cm (N = unknown). Wound complications assessed at 48 to 72 hours.</td>
<td>Adequacy of anesthesia before suturing or duration of anesthesia during suturing of lesions located on the forehead/eyebrow or scalp area.</td>
<td>These was no significant difference between patients anesthetized with TAC (18%) and lidocaine (23%) in their need for additional lidocaine. Percentage of patients in the placebo group (83%) required supplemental lidocaine. Successful initial anesthesia did not differ significantly in any of the anesthetic groups. TAC produced initial anesthesia more often in extremity locations vs lidocaine or placebo.</td>
<td>Blinding only in TAC vs placebo group. Remarkably, 17% of topical placebo group did not require anesthesia. Study was pediatric population.</td>
<td></td>
</tr>
<tr>
<td>Zempsky 1997</td>
<td>RCT</td>
<td>Sponsored by by grant from the General Clinical Research Center, Children's Hospital of Pittsburgh. No mention of COI.</td>
<td>N = 32 (gender not specified) with lacerations. Ages 5 to 18 years.</td>
<td>EMLA without supplemental anesthetica (N = 16) vs TAC for suturing uncomplicated extremity wounds (N = 16). Mean time of anesthetic application in the EMLA-treated group was 55 minutes vs 29 minutes in TAC-treated group, (p &lt; 0.01).</td>
<td>&quot;85% of EMLA group had complete wound repair without supplemental anesthetica, compared with 7 of 16 patients (45%) in the TAC-treated group (p&lt;0.03). The mean time of anesthetic application in the EMLA-treated group was 55 minutes, compared with 29 minutes in the TAC-treated group (p&lt;0.01). The EMLA-and TAC-treated groups were not significantly different with regard to the VAS scores.</td>
<td>&quot;Our data show that extremity wounds treated with EMLA for 60 minutes require supplemental anesthesia less often than those wounds treated with TAC for 30 minutes.&quot;</td>
<td>No mention of control of other analgesics. May not be applicable to adults. Although inclusion criteria was up to 18 years old.</td>
<td></td>
</tr>
<tr>
<td>Kuhn 1996</td>
<td>RCT</td>
<td></td>
<td>N = 181 (gender not specified) with lacerations.</td>
<td>MAC (N = 95/114) vs &quot;There was no significant difference in the overall efficacy of the two solutions. MAC was significantly more effective in anaesthetizing wounds of the</td>
<td>&quot;MAC can be substituted for the less readily available TAC whenever expedient.&quot;</td>
<td>Purpose of study was to determine if acceptable alternative to tetracaine, which is not readily available in Australia. Allocation method and baseline comparability unclear.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vinci 1996 RCT</td>
<td>Age &gt;12 years.</td>
<td>TAC topical anesthesia for wound suturing (N = 37/66). Follow-up unclear.</td>
<td>head than of the extremities (p&lt;0.001), while TAC did not differ significantly in effectiveness between the two sites...Patients' preference for topical anesthesia in the future did not differ markedly between the two treatment groups: 70/86.&quot;</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0</td>
<td>N = 156 with lacerations. Age range 3-18 years.</td>
<td>Group I, TAC 11.8% cocaine (N = 49) vs Group II, TAC 4% cocaine (N = 49) vs Group III, tetracaine plus cocaine 4% for lacerations anesthesia (N = 58). First assessment after 15 minutes and 15 after second application.</td>
<td>“Solutions containing 11.8% cocaine (TAC 1) and 4% cocaine with adrenaline (TAC 2) were significantly more likely (p &lt; 0.001) to produce complete anesthesia then the solution with 4% cocaine without adrenaline...A second dose of TAC 3 was more often required to produce complete or partial anesthesia, (p&lt; 0.003).”</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>“The application of a TAC solution containing 4% cocaine is as effective as a TAC solution containing 11.8% cocaine; use of this 4% solution decreases the cost of the agent.”</td>
<td>No placebo group. Allocation unclear. Population 3-18 year olds.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
WOUND REPAIR
Wound repair is most commonly performed through primary closure (immediate approximation of the wound edges) to reduce discomfort and speed healing. Closure of most low-risk wounds can occur 12 to 24 hours after the injury. Contaminated wounds or those at high risk of infection should be closed within 6 hours. Wounds outside of these parameters can be treated by delayed primary intention after 2-3 days of antibiotics reducing risk for subsequent infection. Sutures are the most common method, followed by staples, adhesives, and tapes.

1. **Recommendation: Non-surgical Management of Non-complicated Hand Lacerations Less than 2cm in Linear Length**
   It is recommended that non-complicated linear lacerations of the hand less than 2cm be managed without suturing by healing via secondary intention for some workers. Wounds should be carefully selected, not have tension, including not overlying or near joints and not have tension applied due to manual labor.
   
   **Strength of Evidence** – Recommended, Evidence (C)
   **Level of Confidence** – Low

2. **Recommendation: Surgical Referral for Hand Lacerations with Evidence of Nerve Injury**
   Immediate referral to a surgeon is recommended if the laceration shows evidence of a nerve injury.
   
   **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   **Level of Confidence** – High

3. **Recommendation: Suture Repair for Hand or Forearm Lacerations**
   Suture repair is moderately recommended for lacerations of the hand or forearm as these lacerations respond well to common suture techniques and suture materials. There are no recommendations for one technique over another or for one suture material type over another.
   
   **Strength of Evidence** – Moderately Recommended, Evidence (B)
   **Level of Confidence** – Moderate

4. **Recommendation: Use of Tissue Adhesive, Staples, and Surgical Tape (Steri-Strips) for Uncomplicated Laceration Repair**
   Tissue adhesives, staples and surgical tape are moderately recommended for routine skin repair of non-complicated extremity lacerations within the limitations of repair strength equivalent to 5-0 suture material or higher.
   
   **Strength of Evidence** – Moderately Recommended, Evidence (B)
   **Level of Confidence** – Moderate

**Rationale for Recommendations**
There is one moderate-quality study comparing suture repair with non-surgical treatment (secondary intention) for hand lacerations less than 2 cm in length and uncomplicated by underlying joint, tendon, fracture, or nerve injury or medical conditions that would affect healing.(1392) There were no differences between the groups in cosmetic appearance, return to activity, or infection. As many hand lacerations are small and uncomplicated, this study suggests non-surgical management for non-gaping uncomplicated lacerations of the hand may be appropriate. Although, a comprehensive recommendation for working populations is not made as the provider should consider tensile forces on the wound and other environmental exposures resultant from occupational duties that likely reduce the ability to use non-surgical management for some patients in making a treatment decision.

There are no quality RCTs of upper extremity wound lacerations comparing suture repair with healing by secondary intent for gaping lacerations exceeding 2cm in linear length. However, wound closure most commonly by suture techniques has been long performed making suture repair the basis for other comparison studies. Therefore, although there is a lack of supporting studies, suturing is considered first line for laceration repair, with the strength of other repair recommendations made against using secondary intent in non-infected wounds.
Various suture techniques have been described to provide the approximation of skin margins. However, there is a relative lack of quality studies that are methodologically sound while also having sufficient follow-up time of greater than one year to derive robust conclusions regarding the relative merit of different suturing techniques. Optimal results are thought to be dependent on skin edge eversion to eliminate depressed scarring, elimination of dead space and minimization of tension of individual sutures to avoid tissue necrosis. Common techniques include simple interrupted, vertical mattress, and running sutures. There are two moderate-quality studies of suture techniques, although there were no direct comparisons between the common techniques. Two versions of vertical mattress were compared with no difference in outcomes in a low quality study.(1393) Simple running cutaneous suture was compared with running subcutaneous sutures (removed at 14 days and not removed non-absorbable suture) and subcutaneous polygalactin (absorbable) running suture for treatment of post-elliptical excisions, rather than traumatic lacerations.(1394) There were short and intermediate advantages of polygalactin vs. polypropylene subcutaneous sutures, which disappeared at 9 months. This study, however, may not be generalizable to laceration repairs. Comparison of single layer vs. bi-layer repair for minor lacerations showed no difference in cosmetic results, although this was a study of facial lacerations, and may not be applicable even though the face is considered cosmetically more sensitive than the extremity.(1395) Therefore, there is no quality evidence to recommend one technique over another, and there are multiple scenarios that one technique might offer technical advantage over another, so that the operator focus may best be on principles for assuring optimal results.

There is also a lack of quality data comparing suture types for extremity laceration repair. The available cosmetic studies are both methodologically weak and have inadequate follow-up times to derive clinically meaningful differences on cosmesis.(1396) One moderate-quality study showed comparability of absorbable catgut to nylon sutures for simple repair.(1397) A low-quality study showed no difference between absorbable suture with nylon suture.(1398) A systematic review in pediatric and adult populations of absorbable vs. non-absorbable sutures did not find superiority of one over the other.(1399) Another moderate-quality study compared Teno Fix® repair, which uses a multifilament stainless steel suture, to a simple repair with cruciate suture for flexor tendon lacerations and found that repairs with the Teno Fix® had lower rupture rates and similar functional outcomes when compared with conventional repair.(1400) Thus, there is insufficient quality evidence to make a recommendation for or against the use of absorbable or non-absorbable suture material for superficial closure of extremity lacerations.

In addition to evaluating different types of sutures, one moderate-quality study compared suturing to stapling and concluded that stapling is more cost-effective than sutures. However, no outcomes measures for cosmetic results or complications were presented.(1401)

There are 17 moderate-quality studies comparing tissue adhesives with standard suture repair of routine extremity lacerations that have shown at least equivalent or superior cosmetic results with no statistically significant increase in infections, dehiscence rates, or other complications.(151, 1402-1419) Advantages to using tissue adhesives also include elimination of the need for local anesthesia, significant reduction of repair time, and ease of performing the procedure saving on visit time over suture repair. Disadvantages of wound characteristics (especially depth and length) and occupational tasks requiring considerable tension are considerable in working populations and are discussed below.

The most commonly used tissue adhesive is octylcyanoacrylate also known as Dermabond®. The other major glue is N-butyl 2-cyanoacrylate, also known as Histoacryl® and Histoacryl Blue® which has a blue tint for reported easier application.(1420) are only two direct comparisons of the compounds, which showed no difference in outcomes measures.(1421, 1422)

In each of the studies which included traumatic and surgical wounds of the hands, upper extremities, trunk and face in both pediatric (1-18 years of age) and adult populations, wound characteristics were usually limited to non-crush injuries, less than 4 cm in length, less than 5mm deep, and without other complicating factors including history of keloid or other scarring disorders, diabetes mellitus,
corticosteroid or other immunosuppressant use, or debilitating illnesses. Thus, the results of equivalency in treatment may not be applicable to many with work-related upper extremity lacerations.

Tissue adhesive was also compared to the use of Steri-Strips in 7 moderate-quality studies as either a primary comparison(1418, 1419) or as part of the “standard care” treatment arm. (1403, 1407, 1410, 1413, 1414) In each trial, Steri-Strips were found to be equivalent in efficacy to tissue adhesive with the same inclusion and exclusion criteria. Tissue adhesive was also compared with the use of skin stapling in 5 moderate quality studies(151, 1404, 1407, 1413, 1414) as part of the non-surgical treatment arms. In each of these studies, the results were equivalent in all outcomes measures.

Therefore, there is strong evidence that tissue adhesives, skin stapling, and adhesive tapes are effective in the repair of routine lacerations of the upper extremity provided they are used on skin areas that are not subject to significant tension (i.e., joints, creases in hand, etc.). In appropriate cases, these have the added advantage of reduced operator or procedural time and material costs compared with suture repair.

Evidence for Wound Repair
There are 29 moderate-quality RCTs incorporated into this analysis.(151, 1392-1395, 1397, 1400, 1401, 1403-1406, 1408-1421, 1423-1425) There are 4 low-quality RCTs(1398, 1426-1428) in Appendix 2.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: wound repair, wound healing, laceration, wound, cuts, management, repair, care, upper extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 454 articles in PubMed, 95 in Scopus, 17 in CINAHL, 2 in Cochrane Library, 15062 in Google Scholar, and 0 from other sources. We considered for inclusion 20 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 9 Google Scholar, and 4 from other sources. Of the 34 articles considered for inclusion, 34 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quinn 2002</td>
<td>RCT</td>
<td>Sponsored by US National Institutes of Health. JQ was paid by Ethicon, for speaking and educational symposiums.</td>
<td>7.5</td>
<td>N = 91 (40 female and 51 male) with lacerations. Age in Suture and Conservative groups: 40 (16) and 38 (15).</td>
<td>Suturing method of securely closing wounds (N = 47) vs Conservative treatment of uncomplicated lacerations &lt;2cm (N = 48). Follow-up at 8 and 10 days.</td>
<td>Mean scores for cosmetic appearance; suturing vs. conservative treatment: Doctor scores 83mm vs. 80mm; patient scores 83mm vs. 82mm. One sutured wound treated with antibiotics for infection. No infections in conservatively treated wounds.</td>
<td>“Similar cosmetic and functional outcomes result from either conservative treatment or suturing of small uncomplicated lacerations of the hand, but conservative treatment is faster and less painful.”</td>
<td>Results are specific to hand lacerations &lt; 2 cm in linear length. The authors caution against generalization to cosmetically sensitive areas.</td>
</tr>
<tr>
<td>Singer 2005</td>
<td>RCT</td>
<td>No mention of sponsorship. No COI.</td>
<td>7.0</td>
<td>N = 65 (9 female and 56 male) with lacerations; mean age 18.5±20.0.</td>
<td>Single-layersutures (N = 32) vs Double-layer closure of facial lacerations (N = 33).</td>
<td>Mean number of deep sutures used in patients assigned to a 2-layer closure was 2.8 ± 1.4. Wound outcomes; Single vs. double-layer. No infections in either group.</td>
<td>“Single-layer closure of non-gaping, minor facial lacerations is faster than double-layer closure.”</td>
<td>Results may not be applicable to other body areas.</td>
</tr>
<tr>
<td>Alam 2006</td>
<td>RCT</td>
<td>Sponsored by by research funds from Department of Dermatology, Northwestern University.</td>
<td>7.0</td>
<td>N = 36 (21 female and 15 male) with lacerations. Age 18-65 years.</td>
<td>Simple running polypropylene sutures 14 days (N = unknown) vs Subcuticular running polypropylene sutures 14 days (N = unknown) vs Subcuticular running polypropylene sutures not removed (N = unknown) vs Subcuticular polyglactin 910 sutures left in place (N = unknown). Follow-up at 3 and 9 months.</td>
<td>No difference in suture at either 3 months or 9 months. Greater scar width at 3 and 9 months, with back wounds being wider, (p &lt; 0.001). No technique was superior.</td>
<td>“While scar width does not appear to vary significantly based on choice of epidermal closure, bilayered closures of the trunk and extremity have better overall appearance and less associated erythema at 3 and 9 months.”</td>
<td>Patient was both control and experimental arm with 2 lesions per person.</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Description</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------------------------</td>
<td>-------</td>
<td>-----------------------------------------------------------------------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jones 1993</td>
<td>7.0</td>
<td>N = 30 (gender not specified) with lacerations. Age for traditional and shorthand group: 27.9 ± 6.3 and 25.3 ± 5.5.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RCT</td>
<td></td>
<td>Shorthand vertical mattress sutures (N = 15) vs Classic mattress sutures for lacerations ranging from 2 to 9cm (N = 15). Follow-up 7 to 10 days for wound assessment.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td>“Suture repair times were significantly shorter using the shorthand vertical mattress stitch compared with the traditional method (88.4 vs. 45.6 sec/suture; p &lt; 0.05). No incidents of significant scar widening, cross-hatching, or prolonged inflammation were noted with the shorthand vertical mattress technique.”</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Karounis 2004</td>
<td>5.0</td>
<td>N = 95 (58 male and 37 female) with lacerations &lt; 12 hours old requiring suture repair. Mean age for groups A and B: 8.1 and 9.5 years.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RCT</td>
<td></td>
<td>Group A, absorbable catgut sutures (N = 50) vs Group B, non-absorbable nylon sutures (N = 45). Follow-up at 4 months.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sponsored by the Montreal Children’s Research Institute and the Canadian Association of Emergency Physicians. No mention of COI.</td>
<td></td>
<td>No differences were found in proportion of optimal WES (6/6) between Group A and NA (62% vs. 49%; relative risk = 0.73%; 95% CI = 0.45 to 1.17). No differences found between Group A and NA for rates of dehiscence (2% vs. 11%; p = 0.07).</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kundra 2010</td>
<td>4.5</td>
<td>N = 100 (21 male and 49 female) elective day case hand and wrist surgery. Mean age Absorbable/ non-absorbable group: 54.0 / 57.3.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RCT</td>
<td></td>
<td>Absorbable 3/0 Vicryl rapide™ (N = 37) vs Non-absorbable (3/0 nylon) for the wound closure (N = 33). Follow-up 6 weeks post-surgery.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td>Mean VAS score for wound satisfaction were 82.5 for non-absorbable group vs 80.4 for the absorbable group. Mean DASH scores were 21.7 vs 21.1 absorbable group.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orlinsky 1995</td>
<td>7.0</td>
<td>N = 141 with suturable linear lacerations of the extremities. Average age was 28 years and 29 for suture group.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RCT</td>
<td></td>
<td>Stapling (N = 78) vs Suturing for skin closure (N = 83). Follow-up until wound closure, average speed for stapling was 8.3 seconds per centimeter and suturing was 63.2 seconds.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td>“The average speed for stapling was 8.3 seconds per centimeter and for suturing was 63.2 seconds,(p = 0.0001) . Cost of labor was calculated to be 1.23 per minute….The relative labor cost of stapling versus suturing was 0.14 (76 cents v $ 5.31, p = 0.001). The speed of repair “We conclude that, with respect to emergency department repair of linear nonfacial lacerations, stapling is a less expensive means of skin closure than suturing.”</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Suture vs. Staples</strong></td>
<td></td>
<td>“The shorthand vertical mattress stitch is an efficient, alternative method for laceration repair that does not compromise wound eversion.”</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Randomization, allocation unclear. High drop-out rate.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Allocations unclear. No blinding.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Data suggest both suture types were comparable, but data based upon questionnaire responses.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Copyright© 2016 Reed Group, Ltd.**
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Sponsor</th>
<th>COI</th>
<th>Participants</th>
<th>Interventions</th>
<th>Outcome Measures</th>
<th>Findings</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Singer</td>
<td>1998</td>
<td>RCT</td>
<td>Sponsored by a grant from Closure Medical, Inc., Raleigh, NC. No mention of COI.</td>
<td></td>
<td>N = 124 (48 female and 76 male) with standard closure of traumatic lacerations. Range age 1-17 years.</td>
<td>Tissue adhesive Ocylcyanoacrylate (N = 63) vs Standard wound closure techniques for lacerations (N = 61).</td>
<td>Follow-up for 3 months.</td>
<td>Patients treated with ocylcyanoacrylate less frequently received local anesthesia (21% vs. 89%, p &lt; 0.001). Groups similar with respect to decontamination with normal saline (81% vs. 75%, p = 0.36), irrigation (50% vs. 65%, p = 0.13), and use of scrub (48% vs 31%, p = 0.08).</td>
<td>&quot;Wounds treated with Ocylcyanoacrylate and standard wound closure techniques have similar appearances 3 months later.&quot;</td>
</tr>
<tr>
<td>Quinn</td>
<td>1993</td>
<td>RCT</td>
<td>Sponsored by the Children's Hospital of Eastern Ontario Research Institute. No mention of COI.</td>
<td></td>
<td>N = 81 (34 male and 47 female) children with clean facial lacerations less than 4 cm in length and 0.5 cm in width. Age range, 0.7 to 16 years and 0.5 to 15 years.</td>
<td>Tissue adhesive Histoacryl Blue® (N = 37) vs Suturing with local anesthetic (N = 38).</td>
<td>Follow-up for 5 days.</td>
<td>Cosmetic outcomes; Histoacryl vs. suture: Mean visual analog scale score (mm) 60.6 vs. 57.2 p = 0.45</td>
<td>&quot;Histoacryl Blue® is a faster and less painful method of facial laceration repair that has cosmetic results similar to the use of sutures.&quot;</td>
</tr>
<tr>
<td>Holger</td>
<td>2004</td>
<td>RCT</td>
<td>Sponsored by HealthPartner s Research Foundation. No mention of COI.</td>
<td></td>
<td>N = 150 (108 male and 42 female) with facial lacerations. Mean age for those completing follow-up and did not: 70.2 and 28.6 (N = 84 and 66).</td>
<td>OC or octylcyano-acrylate tissue adhesive (N = 49) vs NL or 6-0 monofilament suture (N = 49) vs RG or Rapid 6-0 gut absorbable suture (N = 47).</td>
<td>Follow-up at 9 and 12 months.</td>
<td>No clinically significant differences in cosmetic outcome among the three groups at 9-12 months.</td>
<td>&quot;The use of either octylcyanoacrylate or rapid absorbing gut suture could be preferred in this setting (ED), eliminating the need for follow-up visits for suture removal.&quot;</td>
</tr>
<tr>
<td>Sinha</td>
<td>2001</td>
<td>RCT</td>
<td></td>
<td></td>
<td>N = 50 (9 male and 35 female) with variety of hand operations. Mean age for adhesive and suture groups: 49 (9) and 51 (17).</td>
<td>N-buty1 2-cyanoacrylate tissue adhesive (Indermil) (N = 20) vs Sutures (5-0 nylon) at 2 and 6 weeks</td>
<td>No significant difference in cosmetic outcome assessment, but 5 minor wound dehiscences (3 in tissue adhesive group, 2 in suture group).</td>
<td>&quot;Evaluation of patients in the two groups of our study showed similar wound outcomes.&quot;</td>
<td>Post-operative hand surgery wounds. Study limited to 6 week follow-up. Small sample size limits study power.</td>
</tr>
<tr>
<td>Study</td>
<td>Sponsorship</td>
<td>No mention of COI</td>
<td>N</td>
<td>Description</td>
<td>Methodology</td>
<td>Follow-up</td>
<td>Findings</td>
<td>Comments</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>-------------</td>
<td>------------------</td>
<td>---</td>
<td>-------------</td>
<td>-------------</td>
<td>-----------</td>
<td>----------</td>
<td>----------</td>
<td></td>
</tr>
<tr>
<td>Shamiyeh 2001</td>
<td>No mention of sponsorship. No COI.</td>
<td>N = 79 (24 male and 55 female) requiring varicose vein surgery. Are range for group S / T/ and TA: 26 – 70 / 16 – 72 / and 20 – 73.</td>
<td>6.5</td>
<td>S group or Suture 5-0 monofilament (N = 26) vs Group T or adhesive tape (N = 28) vs TA or octylcyanoacrylate tissue adhesive (N = 25).</td>
<td>Follow-up at 2 and 6 weeks.</td>
<td>There were no differences between the groups for dehiscence or infections. The scars were judged slightly better for cosmetic result in the suturing group, but scores were not statistically significant.</td>
<td>&quot;Comparing 5-0 monofilament sutures, tapes, and tissue adhesive for skin closure after phlebotomy, there was no difference in cosmesis, but closure with tape was by far the cheapest method.&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Singer 2002</td>
<td>Sponsored by a research grant from Closure Medical Corporation, Raleigh, NC, which developed TraumaSeal. Two of the authors (AJS, JEH) are on the speaker's bureau of Ethicon Inc. No mention of other COI.</td>
<td>N = 924 wounds of traumatic lacerations, excisions of skin lesions or scar revisions, minimally invasive surgeries, and general surgical procerus. Mean age 31.9 and 30.7 for standard group.</td>
<td>6.0</td>
<td>OCA or octylcyanoacrylate tissue adhesive (N = 406) vs Standard wound closure methods, sutures, adhesive tapes, or staples (N = 408).</td>
<td>Follow-up 5 to 10 days.</td>
<td>Wounds widely distributed over body. Many required subcutaneous sutures (55%). At 5-10 day follow-up, wound dehiscence and infection rates not significantly different between groups. At 3 months, no differences in wounds considered optimal (82% OCA vs. 83% other).</td>
<td>&quot;Repair of traumatic lacerations and surgical incisions with OCA is faster than with standard wound closure techniques, and cosmetic outcome is similar at 3 months.&quot;</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| Quinn 1997 | Sponsored by Closure Medical Corp. | N = 136 (101 male and 35 female) with lacerations requiring suture. Mean age 35.3 ± 14.1 and 36.9 ± 17.2 for suture group. | 6.0 | Skin closure with octylcyanoacrylate adhesive (N = 68) vs Monofilament suture (N = 68). | Follow-up for 3 months. | Octylcyanoacrylate vs. sutures. Mean VAS cosmesis scores, mm: 67 vs. 68 p = 0.65. Mean VAS pain scores, mm: 7.2 vs. 18.0 p <0.01; Infection, No.: 0 vs. 1; Dehiscence, No.: 3 vs. 1 | "Octylcyanoacrylate tissue adhesive effectively closes selected lacerations. This relatively painless and fast method of wound repair can replace the need for high dropout rate at 3 month follow-up."
<p>| | | | | | | | Cost argument is relative, as all treatment material costs were $11 or less. | Allocation unclear although baseline comparability was non-significant. Study included large number of wounds (surgical and traumatic) which may improve applicability. |</p>
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Randomized Controlled Trial (RCT)</th>
<th>Number and Gender</th>
<th>Treatment</th>
<th>Follow-up</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quinn 1998</td>
<td>6.0</td>
<td>Sponsored by a research gift from Closure Medical Corp. No mention of COI.</td>
<td>N = 136 (63 male and 13 female) with traumatic wounds. Mean age for OCT and Sutures groups: 37.4 ± 12.4 and 39.6 ± 18.3 years.</td>
<td>Octylcyano-acrylate tissue adhesive (N = 68) vs 5-0 or smaller monofilament suture (N = 68). Follow-up at 3 months and 1 year.</td>
<td>No differences found in demographic or clinical characteristics between groups. At 1 year, no difference found in optimal wound scores (73% vs. 68%, p = 0.60) or in visual analog scale cosmesis scores (69 vs. 69mm, p = 0.95) for octylcyanoacrylate.</td>
<td>“One year after wound repair, no difference is noted in the cosmetic outcomes of traumatic lacerations treated with octylcyanoacrylate tissue adhesive and sutures. The assessment of wounds 3 months after injury and wound repair provides a good measure of long-term cosmetic outcome.” One year follow-up to 1997 study.</td>
</tr>
<tr>
<td>Bruns 1996</td>
<td>6.0</td>
<td>No mention of sponsorship or COI.</td>
<td>N = 61 (49 male and 12 female) with lacerations less than 12 hours old. Between 1 and 18 years of age.</td>
<td>Histoacryl Blue (HAB) tissue adhesive (N = 30) vs Suture (N = 31) Follow-up at 1 week and 2 months.</td>
<td>Two plastic surgeons blinded to treatment. One rated no difference between groups, other favored HAB for better scar appearance.</td>
<td>“The use of HAB is an acceptable alternative to conventional suturing.” Pediatric population (&lt;18 years old). Allocation not well defined.</td>
</tr>
<tr>
<td>Simon 1998</td>
<td>6.0</td>
<td>Sponsored partially by Closure Medical</td>
<td>N = 61 (49 male and 12 female) with lacerations. Median age for with follow-up and without: 4.0 and 3.0.</td>
<td>Histoacryl Blue (HAB) tissue adhesive (N = 30) vs Sutures in facial lacerations (N = 31). Follow-up at 2 months and 1 year.</td>
<td>Overall ratings of cosmetic outcomes were comparable or better in appearance for HAB group by blinded plastic surgeons. When reviewed by Langer line orientation, cosmetic appearance of sutured lacerations worse against Langer lines vs. sutured with Langer line orientation. No difference in Langer orientation with HAB group.</td>
<td>“The cosmetic appearance of facial lacerations repaired with HAB was comparable to conventional suturing, and appears to be less affected by the initial orientation of the wound with Langers lines than with conventional suturing.” Second report of similar study group. Allocation unclear.</td>
</tr>
<tr>
<td>Toriumi 1998</td>
<td>5.5</td>
<td>Sponsored partially by Closure Medical</td>
<td>N = 111 (gender not specified) underwent surgical procedure for skin closure. Mean age was 41.2 years.</td>
<td>Octyl-2-cyanoacrylate (N = 57) vs 5-0 sutures (N = 54). Follow-up at 5, 7, and 90 days and 1 year.</td>
<td>Difference in time for skin closure between octyl-2-cyanoacrylate and sutures significant (p &lt; 0.0001). No significant difference on modified Hollander scale at 90 days (p = 0.51). However, at 1 year, mean VAS scale for cosmetic outcome showed improved cosmetic results for</td>
<td>“The lower visual analog scale score represented a superior cosmetic outcome at 1 year with the octyl-2-cyanoacrylate as compared with sutures.” Study population was post-operative plastic surgery for facial and neck lesions.</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>N</td>
<td>Setting</td>
<td>Interventions</td>
<td>Outcomes</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>--------</td>
<td>---</td>
<td>---------</td>
<td>--------------</td>
<td>----------</td>
</tr>
<tr>
<td>Simon 1997</td>
<td>5.0</td>
<td>No mention of sponsorship or COI.</td>
<td>43</td>
<td>N = 61 (49 male and 12 female) with lacerations. Median age for with follow-up and without: 4.0 and 3.0.</td>
<td>incisions treated with octyl-2-cyanoacrylate (p = 0.03).</td>
<td>Wounds evaluated at 2 months and 1 year. Wounds comparable in cosmetic appearance at 2 months by one rater and significantly better for HAB by second rater. At 1 year, wounds comparable by both raters.</td>
</tr>
<tr>
<td>Handschel 2006</td>
<td>5.0</td>
<td>No mention of sponsorship or COI.</td>
<td>45</td>
<td>N = 45 with an orbital floor fracture or facial wounds. The mean Age in the adhesive group was 47 years and 42 years in suture group.</td>
<td>Patients rated skin adhesive higher on VAS, whereas surgeons rated sutured wounds as best cosmetically based on photographs. The scar wound depth was statistically significantly greater in skin adhesive group than suture group.</td>
<td>“The adjustment of the edges of the wounds as measured by the depth of the scar is significantly worse with (Dermabond) than with thin sutures. The sutured wounds give better cosmetic results in younger patients in particular.”</td>
</tr>
<tr>
<td>Karcioglu 2002</td>
<td>4.0</td>
<td>No mention of sponsorship or COI.</td>
<td>92</td>
<td>N = 92 (male to female ratio 1.26) with lacerations equal to or shorter than 5 cm. Mean age 34 ± 11.04.</td>
<td>Tissue adhesive (HAB) tissue adhesive (N = 24) vs Suture repairs (N = 28).</td>
<td>“There were no statistically significant scores of cosmetic outcomes at the tenth day and third month. The ratio of patients who reported satisfaction from the method was significantly higher in the HAB group than the sutured group (p = 0.007). Costs of treatment were significantly lower than sutures (p = 0.000).”</td>
</tr>
<tr>
<td>Singer 1998</td>
<td>7.0</td>
<td>No mention of sponsorship or COI.</td>
<td>124</td>
<td>N = 124 (48 female and 76 male) with standard closure of traumatic lacerations; age 1-17 years.</td>
<td>Tissue adhesive (Octylcyanoacrylate) (N = 63) vs Standard wound closure techniques for lacerations</td>
<td>Patients treated with octylcyanoacrylate less frequently received local anesthesia (21% vs. 89%, p &lt;0.001). Groups similar with respect to decontamination with</td>
</tr>
</tbody>
</table>

### Tissue Adhesive vs. Adhesive Strips, Staples

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>N</th>
<th>Setting</th>
<th>Interventions</th>
<th>Outcomes</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simon 1997</td>
<td>5.0</td>
<td>No mention of sponsorship or COI.</td>
<td>43</td>
<td>N = 61 (49 male and 12 female) with lacerations. Median age for with follow-up and without: 4.0 and 3.0.</td>
<td>incisions treated with octyl-2-cyanoacrylate (p = 0.03).</td>
<td>Wounds evaluated at 2 months and 1 year. Wounds comparable in cosmetic appearance at 2 months by one rater and significantly better for HAB by second rater. At 1 year, wounds comparable by both raters.</td>
<td>“The use of HAB in an ideal alternative to conventional suturing for cutaneous closer of low-tension lacerations in children with a long term cosmetic outcome comparable to conventional suturing.”</td>
</tr>
<tr>
<td>Handschel 2006</td>
<td>5.0</td>
<td>No mention of sponsorship or COI.</td>
<td>45</td>
<td>N = 45 with an orbital floor fracture or facial wounds. The mean Age in the adhesive group was 47 years and 42 years in suture group.</td>
<td>Patients rated skin adhesive higher on VAS, whereas surgeons rated sutured wounds as best cosmetically based on photographs. The scar wound depth was statistically significantly greater in skin adhesive group than suture group.</td>
<td>“The adjustment of the edges of the wounds as measured by the depth of the scar is significantly worse with (Dermabond) than with thin sutures. The sutured wounds give better cosmetic results in younger patients in particular.”</td>
<td></td>
</tr>
<tr>
<td>Karcioglu 2002</td>
<td>4.0</td>
<td>No mention of sponsorship or COI.</td>
<td>92</td>
<td>N = 92 (male to female ratio 1.26) with lacerations equal to or shorter than 5 cm. Mean age 34 ± 11.04.</td>
<td>Tissue adhesive (HAB) tissue adhesive (N = 24) vs Suture repairs (N = 28).</td>
<td>“There were no statistically significant scores of cosmetic outcomes at the tenth day and third month. The ratio of patients who reported satisfaction from the method was significantly higher in the HAB group than the sutured group (p = 0.007). Costs of treatment were significantly lower than sutures (p = 0.000).”</td>
<td>“HAB is a cheaper method of laceration repair and results in greater satisfaction of both the patient and the physician. The cosmetic outcomes are the comparable.”</td>
</tr>
<tr>
<td>Singer 1998</td>
<td>7.0</td>
<td>No mention of sponsorship or COI.</td>
<td>124</td>
<td>N = 124 (48 female and 76 male) with standard closure of traumatic lacerations; age 1-17 years.</td>
<td>Tissue adhesive (Octylcyanoacrylate) (N = 63) vs Standard wound closure techniques for lacerations</td>
<td>Patients treated with octylcyanoacrylate less frequently received local anesthesia (21% vs. 89%, p &lt;0.001). Groups similar with respect to decontamination with</td>
<td>“Wounds treated with Octylcyanoacrylate and standard wound closure techniques have similar appearances 3 months later.”</td>
</tr>
</tbody>
</table>

Authors used standardized incision (periorbital) to control wound type. Lack of study details for randomization. Small sample size. Results may be more applicable to cosmetically sensitive areas (face).

Lack of study details. No baseline data presented. High dropout at follow-up visits at 10 days and 3 months.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>Patient Characteristics</th>
<th>Intervention</th>
<th>Outcome Measures</th>
<th>Key Findings</th>
<th>Funding Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bruns 1998</td>
<td>RCT</td>
<td>N = 83 (55 male and 28 female) with lacerations. Mean and median age for 2-OCA and Sutures / Staples: 3.5 (2.0, 5.0) and 4.0 (3.0, 6.0)</td>
<td>2-OCA or 2-Octylcyano-acrylate (N = 42) vs staples, steri-strips or monofilament sutures (N = 41). Follow-up at 3 months.</td>
<td>Length of time for cutaneous closure was decreased (median, 2-OCA 2.9 minutes vs. suture/staple 5.8 minutes; p &lt;0.001). Assessment of pain not significantly different between groups. 95% receiving 2-OCA would choose 2-OCA over standard wound closure at next visit for laceration repair. No significant differences in clinical characteristics between groups at 3 months.</td>
<td>&quot;2-OCA is an acceptable alternative to conventional methods of wound repair with comparable cosmetic outcome.&quot;</td>
<td>Sponsored in part by a grant from Closure Medical, Inc., Raleigh, NC. The authors also acknowledge the ED academic associates, nurses, and physicians for assistance in data collection. No mention of other COI.</td>
</tr>
<tr>
<td>Mattick 2002</td>
<td>RCT</td>
<td>N = 60 (28 male and 16 female) children with suitable lacerations. Between 1-14 years of age.</td>
<td>2-Octylcyano-acrylate or tissue adhesive (N = 30) vs Adhesive strips (N = 30). Follow-up at 3 and 12 months.</td>
<td>Evaluation at 3 and 12 months. &quot;Cosmetic outcome for both treatments was high, with no significance when viewed from the critical eye of both the parent and the plastic surgeon.&quot;</td>
<td>&quot;In conclusion, both tissue adhesives and adhesive strips are excellent &quot;no needle&quot; alternatives for the closure of suitable pediatric lacerations.&quot;</td>
<td>No sponsorship. Ethicon supplied the Dermabond tissue adhesive and the camera. The Steristrips were from</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Methodology</td>
<td>Participants</td>
<td>Intervention</td>
<td>Follow-up</td>
<td>Results</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>-------------</td>
<td>--------------</td>
<td>-------------</td>
<td>-----------</td>
<td>---------</td>
</tr>
<tr>
<td>Zempsky 2001</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>5.5</td>
<td>N = 97 (60 male 37 female) and with simple facial lacerations in children. Mean age for Steri-step group and Dermabond: 5.2 (2.7) and 5.3 (4.1) years.</td>
<td>3M Steri-Strip Closure, 2-Octylcyano-acrylate (N = 48) vs Dermabond or Adhesive strips (N = 49). Follow-up at 2 months.</td>
<td>Wound dehiscence occurred in 1 steri-strips and 5 dermabond patients. No difference in total complication rates between groups (p = 0.11). Wound scores for rating surgeons not significantly different.</td>
</tr>
<tr>
<td>Singer Plast Reconstr Surg 2002</td>
<td>RCT</td>
<td>Sponsored by a research grant from Closure Medical Corporation, Raleigh, NC, which developed TraumaSeal. Two of the authors (AJS, JEH) are on the speaker’s bureau of Ethicon Inc. No mention of COI.</td>
<td>5.0</td>
<td>N = 924 and 814 patients (542 male and 382 female) wounds. Mean age 31.3 (21.1) years.</td>
<td>Octylcyanoacrylate tissue adhesive (N = 455 wounds) vs Standard wound closure methods sutures, adhesive tapes, or staples (N = 469). Follow-up for 3 months.</td>
<td>Characteristics associated with suboptimal cosmetic appearance on multivariate analysis were presence of associated tissue trauma 3.9 (95 C.I. 1.4-10.7), use of electrocautery (OR 2.9, 95% CI 1.8-6.5), extremity location (OR 2.9, 95% CI 1.2-3.7), wound width (OR 1.08, 95% CI 1.01-1.14). Wound infection associated with tissue trauma (8.7% vs. 1.7, p = 0.04) and incomplete wound apposition (6.6 % vs. 0.5 %).</td>
</tr>
</tbody>
</table>

**Tissue Adhesive vs. Tissue Adhesive**

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Methodology</th>
<th>Participants</th>
<th>Intervention</th>
<th>Follow-up</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Osmond 1999</td>
<td>RCT</td>
<td>Sponsored by Closure Medical Corp., Raleigh, NC. No mention of COI.</td>
<td>7.0</td>
<td>N = 94 (37 female and 57 male) with facial lacerations. Age at least 18 years.</td>
<td>Octylcyano-acrylate (N = 47) vs butylcyano-acrylate for superficial linear facial lacerations (N = 47). Follow up at 3 months.</td>
<td>No difference between butylcyanoacrylate and octylcyanoacrylate in time of wound repair (4.2 vs. 4.0 min, p = 0.88), pain induced by the procedure (VAS score 24 vs. 15, p = 0.37), and ease of procedure as rated by study physician (12 vs. 15).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

"Steri-strips and Dermabond provide similar cosmetic outcomes for closure of simple facial lacerations." |

Lack of study details. No allocation and minimal baseline compatibility data provided. |

This is the second report of same population. Some methodology details lacking in this report. |

"Suboptimal wound appearance is increased with extremity wounds, wide wounds, incompletely apposed wounds, associated tissue trauma, use of electrocautery, and infection." |

Study population limited to pediatrics (<18 years old) with facial lacerations.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Sponsorship</th>
<th>COI</th>
<th>Sample Size &amp; Characteristics</th>
<th>Interventions</th>
<th>Outcomes</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Su 2005 RCT</td>
<td>5.0</td>
<td>N = 67 (67 male and 20 male) with 85 flexor tendon injuries digits 2-5. Zone II laceration of flexor digitorum profundus tendon with or without superficialis laceration. At least 18 years of age.</td>
<td>Teno Fix® repair (N = 29) vs Simple repair with cruciate suture (3-0/4-0 polypropylene) plus circumferential (6-0 monofilament nylon). Tendon had to be wide enough for use of the device. Rehabilitation with passive ROM first POD. Kleinert method for 1st 3 weeks (N = 38).</td>
<td>Excellent/good and fair/poor results in: Teno Fix vs. 67% and 33% vs. traditional suture 70% excellent/good and 30% fair/poor. Ruptures developed in 0% Teno Fix vs. 9/51 (18%) traditional suture (p = 0.01). No differences in pain, grip/pinch strength or DASH scores, (p &gt; 0.05).</td>
<td>&quot;Tendon repairs with the Teno Fix® have lower rupture rates and similar functional outcomes when compared with conventional repair, particularly in patients who are noncompliant with the rehabilitation protocol.&quot;</td>
<td>Some baseline differences may be due to 7 crossovers to control group for technical reasons. High dropouts in control group at 6 months. More smokers in control group combined with more ruptures in controls raise concern for potential confounding.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sener 2015 RCT</td>
<td>5.5</td>
<td>N = 54 (39 male and 15 female) with hand lacerations. Age range 18-65 years.</td>
<td>Local infiltration anesthesia or LIA; hydrochloride 2% and 27 gauge needles used (N = 23) vs Peripheral nerve block or PNB (N = 31).</td>
<td>Response to injection pain and suture pain, (p = 0.220 and p = 0.316). Patient satisfaction and need for additional local anesthetics, (p = 0.785 and p = 0.628). Difference statistically significant for time to loss of pinprick sensation in the local infiltration group 1.3 min vs 2.2 minutes in block group, (p &lt; 0.001). Significant difference regarding pain response to suturing; 8.8 vs 14.50, (p = 0.045).</td>
<td>&quot;In conclusion, LIA or PNB for hand laceration surgery is convenient and predictable.&quot;</td>
<td>Data suggest both groups with comparable efficacy except for time required to administer (nerve block 2.2 min and local anesthesia 1.3min)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Moazzam 2003 RCT</td>
<td>5.0</td>
<td>N = 20 (17 male and 3 female) undergoing free radial forearm flap surgery. Average age 58 years (range 28–84).</td>
<td>Cross-suturing, using a 4/0 gauge suture of Polyglyconate (N = 10) vs Control, the graft was applied without cross-suturing of the wound (N = 10).</td>
<td>Cross-suturing group had immediate reduction in size of 30-68%, the mean reduction of 53%. Reduction of area of the cross-sutured forearm scars made after 3-7 months from 40 to 77%, with a mean reduction of 65%. At 3-7 months after surgery in the control cases had a reduction in scar area ranging from 17 to 68%, the mean of 38%.</td>
<td>&quot;A cross-suturing technique is presented to reduce the deformity of the radial forearm flap donor defect.&quot;</td>
<td>Small sample size. Data suggest cross-suturing technique decreased size of forearm deformity when compared to controls (65% vs. 38%) as well as decreasing the area of the split skin donor site.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Follow-up Wound Care

1. **Recommendation: Semi-occlusive or Occlusive Dressing of Wounds**

   There is no recommendation for or against the use of semi-occlusive or occlusive dressing for wounds. The use of semi-occlusive dressings is commonly used although there is little evidence that this practice improves infection rate or cosmetic outcomes. Dressings may be more indicated based on potential contamination at work or other workplace exposures.

   - **Strength of Evidence** – No Recommendation, Insufficient Evidence (I)
   - **Level of Confidence** – Low

2. **Recommendation: Routine Wound Recheck by Health Professional**

   It is recommended that complicated wounds repaired with sutures or staples and heavily contaminated or infected at initial presentation be closely followed-up within 24 to 72 hours and at suture removal.

   - **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   - **Level of Confidence** – Moderate

**Rationale for Recommendations**

There is no quality evidence on proper wound dressing of upper extremity lacerations, the timing and necessity of wound recheck by a health professional, and the timing of suture removal. Upon completion of wound repair, common practice remotely was to cover the wound with semi-occlusive non-adherent dressing for 24 to 48 hours with topical antimicrobial product.(1429) Based on two reports from the 1960s, it was common practice is to keep the wound moist, which was thought to promote re-epithelialization, and reduce risk of infection. However, there are no quality trials supporting this practice and some question the concept.(1430-1432) Current practice is to minimize the use of these dressings for most lacerations to promote movement and use of injured body part(s) and frequently involves the use of bacitracin or poly-antibiotic ointment.

There is one related moderate-quality study comparing infection rates after dermatological excision and repair of wounds that were either left uncovered after 12 hours and allowing normal bathing vs. those that were kept dry under bandage for 48 hours. In this post-surgical population of 857 patients, there was no statistical difference in the infection rate, demonstrating that wounds can be uncovered and allowed to get wet in the first 48 hours without significant risk.(1433) However, it is unclear if traumatic lacerations would react different from surgically controlled wounds so no recommendation is made for or against use of occlusive or semi-occlusive dressings. Physician discretion is indicated dependent on the wound and characteristics of workplace exposures of the wound.

Wound care instructions are usually provided verbally or in written format including information on monitoring for signs of infection. There are no studies on post-repair infection rates comparing persons who have received verbal or written instructions with those that return in 24 to 48 hours for a wound check. However, there is one case series of 433 patients that on follow-up evaluation were asked to rate their wound based on wound care instructions provided for signs of infection. On physician examination, 21 were deemed to have a wound infection. Of these 21, 10 patients did not rate their wound as infected giving a false negative rate of 48% (10/21), although the false positive rate was low at 8%. It is, however, uncertain if these would have resolved or resulted in serious infection, as the follow-up visit occurred at different times, including suture removal. Thus, providing wound care instructions is likely useful, costs little except additional provider time, and may prevent serious infections from going undetected. Routine wound check at 24 to 72 hours is also a common practice and is recommended for complicated wound repair, those that are contaminated or with suspicion of retained foreign bodies, already infected at initial presentation, or if patient is working in unclean environments.(403)

Suture removal for optimal results in upper extremity lacerations is not well defined by quality studies. Common practice is removal of sutures or staples in cosmetically sensitive areas with low tension in 3 to 5 days, 1 week in lower tension areas on the upper extremities, and 10 to 14 days in high-tension areas.(151, 403, 1434) Wounds closed with cyanoacrylates or surgical tape are less likely to have
concerns and follow-up may not be needed except for documentation of healing for patients in workers compensation systems.

*Evidence for Follow-up Wound Care*
There is 1 moderate-quality RCT incorporated into this analysis.(1433)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: follow-up wound care, semi occlusive dressing, routine wound check, wound healing, laceration, wound, cuts, management, repair, care, upper extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 67 articles in PubMed, 84 in Scopus, 176 in CINAHL, 10 in Cochrane Library, 25 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, CINAHL, Cochrane Library, Google Scholar, and 0 articles from other sources. Of the 1 article considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heal 2006 RCT</td>
<td>6.0</td>
<td>N = 857 (600 male and 257 female) with wounds or minor skin excision. Mean age 44 years.</td>
<td>Intervention group, or wound kept dry and covered 48 hours (N = 450) vs Control group, dressing removal and bathing within 12 hours of repair (N = 420). Follow-up within 12 and 24 hours.</td>
<td>Infection rates: dry group 8.9% vs. no dressing and wet 8.4%, intervention rate ratio not inferior to control p &lt;0.05.</td>
<td>“Wounds can be uncovered and allowed to get wet in the first 48 hours after minor skin excision without increasing the incidence of infection.”</td>
<td>Wounds were post-surgical excision repairs, which may be different characteristically from traumatic laceration. No blinding.</td>
</tr>
</tbody>
</table>

*Prescription and Other Medications*

**ANTIBIOTIC PROPHYLAXIS**

*Recommendation: Antibiotic Prophylaxis in Uncomplicated Hand and Forearm Lacerations*

Routine antibiotic prophylaxis is not recommended for uncomplicated hand and forearm lacerations.

*Strength of Evidence – Not Recommended, Evidence (C)*  
*Level of Confidence – Moderate*

**TOPICAL ANTIMICROBIALS**

*Recommendation: Use of Topical Antimicrobials for Wound Care*

There is no recommendation for or against the use of topical antimicrobials for wound care as there is little evidence that this practice improves clinical infection rate or cosmetic outcomes.

*Strength of Evidence – No Recommendation, Insufficient Evidence (I)*  
*Level of Confidence – Low*

*Rationale for Recommendations*

There are two moderate-quality studies conducted over 25 years ago that demonstrated no difference in infection rates between no treatment or placebo and prophylactic oral doses of clindamycin, flucloxacillin, trilopen, and aerosolized povidine – iodine applied directly into the wound.(1435, 1436) However, one moderate-quality study did find that wound irrigation with penicillin provided reduced rates of wound infection.(1437) Each of these studies had significant weaknesses, and strong conclusions cannot be drawn. Two low-quality studies of cephalaxin and clindamycin demonstrated no improvement in infection rates.
rates but are excluded from the analysis because of lack of study details. There are no quality data or recent data on newer broad-spectrum antibiotics for prophylaxis. Adequate irrigation, cleansing, and debridement of non-complicated wounds is therefore recommended as first line treatment to prevent infection, whereas antibiotic prophylaxis is not recommended unless other complicating factors warrant.

The use of topical antimicrobials is also common, but it is controversial. A high-quality study demonstrated a lower infection rate in wounds treated with topical antibiotics vs. petroleum ointment, although the control group’s treatment may preclude strong conclusions. Although there was lower incidence of infection in the active antimicrobial arms vs. petrolatum, the infection rates were similar to other reported incidences that did not use any ointment. It is not possible to determine if the use of antimicrobial is efficacious, or if the use of non-antimicrobial ointment may increase infection risk. Thus, there is insufficient evidence to recommend for or against the use of topical antimicrobials, although they are generally inexpensive, easy to apply, and have relatively low risks for adverse effects. Wounds closed with tissue adhesives should remain uncovered, and application of ointments or antimicrobials should be avoided to reduce risk of dehiscence. An additional concern is that neomycin is considerably allergenic, thus neomycin-containing compounds may have a relative disadvantage.

Evidence for the Use of Antibiotic Prophylaxis
There is 1 high-quality RCT on topical antimicrobials and 3 moderate-quality RCTs on antibiotic prophylaxis that are incorporated into this analysis.

Antibiotic Prophylaxis
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Antibiotic, Prophylaxis, Wound, Healing, Laceration, Cuts, Management, Repair, care, Upper, Extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 5 articles in PubMed, 4 in Scopus, 8 in CINAHL, 8590 in Google Scholar, and 1 in Cochrane Library. We considered for inclusion 2 from PubMed, 0 from Scopus, 1 from CINAHL, 2 from Google Scholar, 1 from Cochrane Library and 0 from other sources. Of the 8608 articles considered for inclusion, 4 randomized trials and 6 systematic studies met the inclusion criteria.

Topical, Antimicrobials
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Topical, Antimicrobials, Wound, Healing, Laceration, Cuts, Management, Repair, care, Upper, Extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 58 articles in PubMed, 0 in Scopus, 8 in CINAHL, 5960 in Google Scholar, and 1 in Cochrane Library. We considered for inclusion 2 from PubMed, 0 from Scopus, 3 from CINAHL, 5960 from Google Scholar, 3 from Cochrane Library and 0 from other sources. Of the 6026 articles considered for inclusion, 0 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dire 1995</td>
<td>8.5</td>
<td>N = 426 (gender not specified) with hand lacerations. Age BAC/NEO/SIL/PTR group: 19.9 (15.1)/18.3 (13.7)/19.7 (14.1)/17.1 (13.1).</td>
<td>BAC or topical antimicrobials Neomycin (n = 109) vs. NEO group or bacitracin (n = 110) vs. SIL or silvadene group (n = 99) vs. PTR or petroleum ointment group (n = 108). Follow-up for 15 months.</td>
<td>Wounds were primarily head/neck followed by hand, lower extremity, and arm. Overall 42/426 infections (9.9%). Infection rates with 95% CI. Bacitracin 5.5% (2.0-11.6), Neomycin 4.5% (1.5-10.3), Silvadene 12.1% (6.4-20.2), Petrolatum 17.6% (10.9-26.1). Petrolatum was significantly higher (p = 0.0034) than others. No differences between other arms.</td>
<td>&quot;The use of topical antibiotics resulted in significantly lower infection rates than the use of a petrolatum control.&quot;</td>
<td>Study unable to address question of anti-microbial vs. no topical preparation. Infection rates in antimicrobial arms similar to previous studies using same techniques without antimicrobial treatment. Possible conclusion is that use of ointments without antimicrobial therapy increase risk of infection.</td>
</tr>
<tr>
<td>Lindsey 1982</td>
<td>6.0</td>
<td>N = 260</td>
<td>Gender and age were not disclosed.</td>
<td>0.9% NaCl vs. 5% sodium benzyl penicillin for lacerations Follow-up history not disclosed.</td>
<td>&quot;The study was terminated …after the inclusion of 260 lacerations, when the upper sloping boundary was crossed for late infections… [A]nalyses of the distribution of preferences in the data at the time of stopping the study indicated high levels of statistical significance in the early purulent infections as well.&quot;</td>
<td>&quot;It appears that two out of three or three out of four infections can be averted merely by flooding the wound with penicillin immediately before suture.&quot; Methodology details sparse. Analyses and results also sparse.</td>
</tr>
<tr>
<td>Roberts 1985</td>
<td>4.5</td>
<td>N = 418</td>
<td>Povidone Iodine average age 33.0 with 74.3% male and No treatment average age 28.1 with 71.2% males.</td>
<td>Povidine-iodine powder aerosol treatment of wound vs. none prior to suture repair Follow-up history not disclosed.</td>
<td>&quot;There was no significant difference in the infection and imperfect healing rates between the povidine iodine and control groups. Significant factors (P&lt;0.01) in the infected wounds were the condition of the dressing and part of the injured hand (palmar injuries). Neither the patients age, the time from injury to suturing or the number of sutures made a significant difference to the incidence of perfect healing.&quot;</td>
<td>&quot;This trial does not show a significant difference in infection rate with povidine iodine therapy. The number of infected cases which were statistically analyzed was small.&quot; No blinding of observer. Lack of study details. High dropout rate.</td>
</tr>
<tr>
<td>Roberts 1977</td>
<td>4.0</td>
<td>N = 368</td>
<td>patients with hand lacerations. Triplopen group mean age is 30.4,</td>
<td>Trilopen IM vs. Fluocoxacillin PO vs. Control (no antibiotics) Follow-up 7 days after suturing.</td>
<td>&quot;Chi-square analysis showed no significant difference in infection rate between the three groups (P &gt; 0.3), but the Triplopen-treated group healed better (P &lt; 0.05) than either of the other groups. Severe contamination of the original wound and a change of dressing carried</td>
<td>&quot;Overall infection rate was 9.8 %, lower than other published work. Our results show that a course of fluocoxacillin gave no improvement in wound healing over a policy of using no Lack of study details. No allocation or baseline compatibility data provided.</td>
</tr>
</tbody>
</table>

Copyright © 2016 Reed Group, Ltd.
| Flucloxacillin group mean age is 29.8, and No antibiotics group mean age is 33.8. No gender disclosed. | out at home were also found to be significant compared to controls.” | antibiotics. The other surprising fact…58% of patients said they had experienced no pain at all when the anesthetic had worn off.” |
**NSAIDs/ACETAMINOPHEN**

Recommendation: NSAIDs or Acetaminophen for Upper Extremity Post-laceration Repair

NSAIDs or acetaminophen are recommended to control pain associated with upper extremity post-laceration repair.

*Indications* – Pain due to upper extremity post-laceration repair.

*Frequency/Duration* – Scheduled dosage rather than as needed is generally preferable.

*Indications for Discontinuation* – Resolution of pain, lack of efficacy, development of adverse effects, particularly gastrointestinal.

*Strength of Evidence* – **Recommended, Insufficient Evidence (I)**

*Level of Confidence* – Moderate

**Evidence for the Use of NSAIDs/Acetaminophen**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDS, Wound Healing, Laceration, Lacerations, Wound, Cuts, Management, Repair, care, upper extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 3 articles in PubMed, 10 in Scopus, 0 in CINAHL, 1 in Cochrane Library, 2900 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**OPIOIDS**

See Opioids recommendations in Carpal Tunnel Syndrome section.

**Physical Methods/Rehabilitation**

**EXERCISE**

Exercise is not indicated acutely. For a few patients with major trauma, or complex wounds, exercise in the recovery period is necessary. For patients with residual deficits, particularly post-operatively, see section on Post-Operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders.

**Evidence for the Use of NSAIDs/Acetaminophen**

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: exercise, exercising, physical activity, wound healing, laceration, wound, cuts, management, repair, care, upper extremity, hand, arm, forearm; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 72 articles in PubMed, 39 in Scopus, 17 in CINAHL, 195 in Cochrane Library, 72,700 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.
**WOUND CULTURE**

*Routine Wound Culture and Sensitivity of Animal and Human Bites*

Routine culture and sensitivity of animal and human bite wounds is moderately not recommended as it has not been shown to be an effective predictor for infection or subsequent treatment of infected wounds.

**Strength of Evidence** – Moderately Not Recommended, Evidence (B)

**Level of Confidence** – Moderate

**Rationale for Recommendation**

There is 1 high-quality study and one moderate quality study of primarily animal, but also included some human bites where uncomplicated bite wounds were routinely cultured prior to treatment assignment.(162, 163) In both studies, there was no correlation between the pathogens that were cultured and any subsequent cultures from infected wounds.(162, 163) Another study also provided culture data, which confirmed expected flora, but no association was drawn in the analyses with subsequent infections.(1441) These analyses only apply to wounds that have no joint, tendon or tendon sheath involvement.

**Evidence for the Use of Bite Wound Cultures and Sensitivity of Animal and Human Bites**

There is 1 high-(163) and 2 moderate-quality(162, 1441) RCTs incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: wound culture, human, animal, dog, cat, bite, bites, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 1 articles in PubMed, 12 in Scopus, 0 in CINAHL, 17 in Cochrane Library, and 29,100 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 3 from other sources. Of the 3 articles considered for inclusion 3 diagnostic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skurka 1986</td>
<td>8.5</td>
<td>N = 39</td>
<td>Penicillin V-K (100,000 U/Kg/day q6hours x 2 days (n = 19) vs. Placebo (n = 20).</td>
<td>Overall infection rate 7.7%. Infection rate of antibiotic group = 5% vs. placebo = 10.5%, (p = NS).</td>
<td>“Prophylactic penicillin failed to prevent infection in dog bite wounds. Cultures showed various organisms but were of no predictive value for development of infection. It seems failure is better correlated to the quality of the local wound care than to prophylactic antibiotic.”</td>
<td>Small sample size. No control for co-interventions. Culture samples of infected wounds not resistant to penicillin. Sample size for wounds sutured too small for comparison (N = 2), although neither became infected.</td>
</tr>
<tr>
<td>Brakenbury 1989</td>
<td>5.0</td>
<td>N = 122 (42 female, 80)</td>
<td>Amoxicillin/clavulanate for 5 days vs. placebo</td>
<td>Non-significant trend toward faster healing with</td>
<td>Amoxicillin/clavulanate significantly reduced the wound infection rate</td>
<td>Study included a mixture of dog, human, and cat</td>
</tr>
<tr>
<td>RCT Beecham Research Laboratories sponsored the research and helped with the analysis.</td>
<td>male). Mean ages of antibiotic and placebo groups for general bites is 30 and 34. Mean ages for same groups for hand bites are 30 and 37.</td>
<td>in full thickness animal bite wounds.</td>
<td>amoxicillin/clavulanate. No difference in age subgroups in rate healing. In adults, 33% of wounds in antibiotic treatment group became infected vs. 60% receiving placebo (p = 0.009). In children, difference non significant (24% antibiotic vs. 20% placebo). Wound infection significantly reduced by antibiotics in wounds older than 9 hours, but not in fresher wounds.</td>
<td>patients with bites where the skin is broken and where the patient presented 9 to 24 hours after injury.</td>
<td>bites, although a majority was dog bites. Study included primarily bites to the hand.</td>
<td></td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td>4.0</td>
<td>N = 55 (gender not specified) with mean age for penicillin group and control group being 10.5 and 9.5 respectively.</td>
<td>Penicillin V 250mg PO QID for 5 days vs. no antibiotics</td>
<td>Overall infection rate 3.6%, with no significant difference between control and penicillin groups. No difference in types of organisms isolated prior to treatment.</td>
<td>Penicillin prophylaxis of superficial non-facial dog bites in children appears no better than local wound care alone when lesions are cleansed soon after occurring. Initial cultures of dog bite wounds have no value in predicting subsequent wound infection.</td>
<td>Quasi-randomization by odd-even day of admission. No blinding, non-placebo control group.</td>
<td></td>
</tr>
</tbody>
</table>

**Initial Care**

   For human bites, it is recommended that exposures that could be considered high risk for viral blood borne pathogen transmission be evaluated and treated according to blood borne pathogen protocols.

   **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   **Level of Confidence** – Moderate

   **Rationale for Recommendation**
   There is no quality evidence for or against implementing blood borne pathogens protocols for human bites. However, exposures that could be considered high risk for transmitting viral blood borne pathogens (HIV, HBV, HCV), such as a traumatic bite lacerations where the offender may have concurrent oral trauma (fight, accident, seizure) should be considered for testing and prophylaxis according to standard protocols.
particularly as needlestick injuries with HIV contaminated blood carry substantially reduced risk of transmission if prophylactic anti-virals are administered in a timely manner. Institutions where employees are at higher risk for human bites may consider implementing policies for this particular class of injuries. A recent study of police officer bite exposures reported an estimated exposure rate to possible viral transmission of 68/10,000/year. Of these measured exposures for this group, 89 (79%) sources were tested, finding 4% HBV-positive, 4% HIV-positive, and 18% HCV-positive. (161)

2. Recommendation: Prophylactic Antibiotics for Dog Bite Wounds

Prophylactic antibiotics are recommended for treatment of dog bite wounds.

Indication – All dog bites. It may be reasonable to omit antibiotics for minor wounds.

Dose/Frequency – Different antibiotics have been used in the quality studies, including penicillin VK, cloxacillin, dicloxacillin, erythromycin, co-trimoxazole, cephalxin, and amoxicillin/clavulanate. Strong Gram positive coverage is required. Tailoring the antibiotic selection to anticipated local antibiotic resistance profiles is advisable.

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – Low

Rationale for Recommendations
A pooled study of wound infection rates from dog bites was performed for this guideline that utilized the published data from all high- and moderate-quality studies antibiotics and showed a 37% reduction in wound infections compared with placebo (Odds Ratio 0.63, 95% CI 0.40, 0.97). These studies analyzed penicillin, (162, 163) penicillinase resistant penicillins, (1442-1444) sulfa compounds, (1445) erythromycin, (1442, 1444) or amoxicillin/clavulanate. (1441) Thus, there is no clear preferential antibiotic to recommend. The individual studies all failed to show statistically significant reductions in infections, but were likely underpowered as infections are relatively infrequent and the studies had modest sample sizes. Prophylactic antibiotics are not invasive, have low adverse effects and are low cost particularly for generic compounds and thus are recommended for treatment of dog bites.

Evidence for the Treatment of Dog Bites
There is 1 high-(163) and 5 moderate-quality (162, 1563, 1564, 1566, 1567) (Rosen 85) RCTs incorporated into this analysis. There is 1 low-quality RCT in Appendix 2. (1565)

Blood Borne Pathogen Protocol
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Blood borne pathogen protocol, Human bites, animal, dog, cat, bites, bite, Torso, Upper Extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 0 in Scopus, 0 in CINAHL, 1 in Cochrane Library, 618 in Google Scholar, and 7 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 7 from other sources. Seven articles met the inclusion criteria.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Prophylactic Antibiotics/ Cat bites, lacerations, upper extremity, bites, hand, arm, forearm; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 1 article in PubMed, 6 in Scopus, 2 in CINAHL, 9 in Cochrane Library, and 1542 in Google Scholar. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 in Google Scholar, and 0 from other sources. Of the 2 articles considered for inclusion, 0 randomized trials and 1 systematic study met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skurka 1986</td>
<td>8.5</td>
<td>N = 39 (gender not specified) with obviously infected wounds, allergy to penicillin, antibiotics administered within 3 days prior to bite. (aged one to 19)</td>
<td>Penicillin V (-\text{K}) (100,000 U/Kg a day 6 hours for 2 days (n = 19)) vs. Placebo (n = 20).</td>
<td>Overall infection rate 7.7%. Infection rate of antibiotic group = 5% vs. placebo = 10.5%. (p = \text{NS}).</td>
<td>&quot;Prophylactic penicillin failed to prevent infection in dog bite wounds.&quot;</td>
<td>Small sample size. No control for co-interventions. Culture samples of infected wounds were not resistant to penicillin. Sample size for wounds sutured too small for comparison (n = 2), although neither became infected.</td>
</tr>
<tr>
<td>Brakenbury 1989</td>
<td>5.0</td>
<td>N = 125 (42 females, 80 males for adults &amp; 20 females, 43 males for kids) with dog, human and cat bites. Mean age for adults: 33.5 years, and 9 for kids.</td>
<td>Augmentin (n=88) for 5 days vs. Placebo (n=97) in full thickness animal bite wounds.</td>
<td>Non-significant trend toward faster healing with amoxicillin/clavulanate. No difference in age subgroups in healing rates. In adults, 33% of wounds in antibiotic treatment became infected vs. 60% receiving placebo (p = 0.009). In children, difference non-significant (24% antibiotic vs. 20% placebo). Wound infection significantly reduced by antibiotics in wounds older than 9 hours, but not in fresher wounds.</td>
<td>&quot;Amoxicillin/clavulanate significantly reduced the wound infection rate in patients with bites where the skin is broken and where the patient presented 9-24 hours after injury.&quot;</td>
<td>Study included a mixture of dog, human, and cat bites, although a majority was dog bites. Study included primarily bites to the hand.</td>
</tr>
<tr>
<td>Jones 1985</td>
<td>4.5</td>
<td>N = 113 patients for dog bite wounds. Mean age, gender not specified.</td>
<td>5 day course of Co-trimoxazole 960 mg twice daily (n=58) wounds vs. placebo (n=55) wounds</td>
<td>Incidence of wound infection 13.8% in placebo vs. 5.5% in antibiotic group (p = 0.135). Hand wounds, infection rate 16.7% in placebo vs. 0% antibiotic (p = 0.0595).</td>
<td>&quot;In conclusion, we feel that the routine treatment of dog bite wounds with antibiotics is not justified, but that hand wounds should be considered for such treatment.&quot;</td>
<td>Thirty-five subjects who failed to return for follow-up were classified as non-infected. Study had low power (required 370 patients in each group for sufficient power).</td>
</tr>
<tr>
<td>Rosen 1985</td>
<td>4.5</td>
<td>N = 33 (66 wounds) (73 females, 77 males) with dog-bite wounds who were admitted within Prophylactic antibiotics (either cloxacillin, dicloxacillin or erythromycin) group receiving 250mg 4x times daily for 5 days (n)</td>
<td>Overall infection rate was 7.6% with 2/35 infections in antibiotics group, 3/31 in placebo group (p = \text{NS}). All infected wounds were of the hand/wrist vs. elsewhere (p &lt; 0.01).</td>
<td>&quot;Antibiotic administration does not reduce the likelihood of subsequent infection in the management of recent dog-bite wounds, or the incidence of infection when only hand wounds were considered.&quot;</td>
<td>Authors found higher risk for infection in hand/wrist wounds than other body parts. No information provided on compliance or other co-interventions.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Participants</td>
<td>Intervention</td>
<td>Outcome</td>
<td>Conclusion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>--------------</td>
<td>--------------</td>
<td>---------</td>
<td>------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Boenning 1983</td>
<td>N = 55 children with non-facial dog bites (gender: not specified) Mean age for penicillin group: 10.5; control group: 9.5 years</td>
<td>Penicillin V 250mg PO QID for 5 days (n=25) vs. no antibiotics only local wound care (n=30)</td>
<td>Overall infection rate 3.6%, with no significant difference between control and penicillin groups. There was no difference in types of organisms isolated prior to treatment.</td>
<td>Penicillin prophylaxis of superficial non-facial dog bites in children appears to be no better than local wound care alone when lesions are cleansed soon after they occur. Initial cultures of dog bite wounds have no value in predicting subsequent wound infection.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dire 1992</td>
<td>N = 185 (75 female/110 male). Mean age 9.0 for antibiotic group and 9.2 for placebo group.</td>
<td>Oral antibiotics (cephalexin, dicloxacillin or erythromycin) (n=89) vs. no antibiotic treatment. (n=96)</td>
<td>One wound (1.1%) in antibiotic group and 5 (5.1%) in control group became infected (p = 0.212). No partial-thickness wounds became infected. No difference in wound infection rates for sutured wounds in the two groups (p = 0.562).</td>
<td>“Our results do not show a significant difference in wound infection rates among all low-risk dog bite wounds with or without oral antibiotic use. Routine prophylactic antibiotics would not seem cost-effective in the low-risk dog bite population.”</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

8 hours of the incident. Mean age 27.8 for antibiotics group and 31.8 for placebo group. Placebo control group (n = 31 wounds). Both groups received standardized wound cleaning based on protocol. Follow-up at 2 or 3 days. Quasi-randomization by odd-even day of admission. No blinding, non-placebo control group. |

Boenning 1983 RCT No mention of sponsorship or COI. Penicillin prophylaxis of superficial non-facial dog bites in children appears to be no better than local wound care alone when lesions are cleansed soon after they occur. Initial cultures of dog bite wounds have no value in predicting subsequent wound infection. Sparse study details. No blinding or placebo. Wounds were irrigated with povidone-iodine.
3. **Recommendation: Prophylactic Antibiotics for Uncomplicated Human Bite Wounds**

   Prophylactic antibiotics are recommended for treatment of uncomplicated human bite wounds.

   **Strength of Evidence** – **Recommended, Evidence (C)**
   
   **Level of Confidence** – **Low**

**Rationale for Recommendation**

There is one moderate-quality study of human bites,(164) and another moderate-quality study that included human bites along with other animals(1441) comparing the utility of prophylactic antibiotics in preventing wound infections. However, despite a relatively modest sample size in the sole study addressing risk of infection from human bites, a broad-spectrum oral antibiotic or IV antibiotics was found to be effective in preventing infection.(164) The study, which included dogs, cats, humans, and other animals, did not find any significant differences using Augmentin®.(1441) Given the reported higher incidence of wound infections related to human bites and the sole quality study addressing this question documenting success, the balance of evidence suggests prophylactic treatment is appropriate. Pathogens are usually gram-positive bacteria; prophylactic coverage from a broad-spectrum oral antibiotic is suggested to cover most typical staphylococcal and streptococcal species.

**Evidence for the Treatment of Human Bites**

There is 1 moderate-quality RCT incorporated into this analysis.(164)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Prophylactic Antibiotics / Human bites, torso, Upper extremity, lacerations, antibiotics, Animal bites ;controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, random; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 8 in Scopus, 1 in CINAHL, 5 in Cochrane Library, and 3161 in Google Scholar. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 3 from Google Scholar, and 1 from other sources. Of the 5 articles considered for inclusion, 1 randomized trial and 3 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zubowicz 1991</td>
<td>5.0</td>
<td>N = 48</td>
<td>Ceclor 250mg po tid vs Kefzol 1gm IV q8h and penicillin G 1.2 million U IV q 6 h vs. placebo</td>
<td>Infection rate in placebo group was 47% (7/15) with no infections in oral or IV antibiotics groups (p &lt;0.05).</td>
<td>In uncomplicated human hand bite, wound toilet coupled with daily dressing changes and an oral prophylactic broad-spectrum antibiotic is satisfactory treatment in compliant patient.</td>
<td>Adult population. Sparse study details including lack of randomization and allocation methods. Patients admitted to hospital for control of co-interventions and compliance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>patient s  (23 males, 25 females). Mean age:26 years</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

4. **Recommendation: Prophylactic Antibiotics for Uncomplicated Cat Bite Wounds**

Prophylactic antibiotics are recommended for treatment of uncomplicated cat bite wounds.

   **Strength of Evidence** – **Recommended, Insufficient Evidence (I)**
   
   **Level of Confidence** – **Low**

**Rationale for Recommendation**

There are no quality studies of antibiotic prophylaxis for cat bites. Only one study was found, but was relatively unhelpful due to limited sample size.(1446) However, the study showed a high incidence of wound infection in the placebo group (4 of 6) compared to none in the oxacillin prophylaxis group. Reported incidence rates of infections from cat bites is 20 to 40%,(153) and complications related to cat bites may be more significant. Therefore, broad spectrum antibiotics that include coverage for
Pasteurella multocida, which is the most common pathogen contracted from cat bites, (1447) may be indicated.

Evidence for the Use of Prophylactic Antibiotics

There are no quality studies incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Prophylactic Antibiotics/ Cat bites, lacerations, upper extremity, bites, hand, arm, forearm; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 1 article in PubMed, 6 in Scopus, 2 in CINAHL, 9 in Cochrane Library, and 1542 in Google Scholar. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 1 in Google Scholar, and 0 from other sources. Of the 2 articles considered for inclusion, 0 randomized trials and 1 systematic study met the inclusion criteria.

5. Recommendation: Laceration Repair for Dog-Bite Wounds

Suturing of non-complicated dog bite wounds after adequate wound care is recommended as it may lead to a better cosmetic result and is not likely to result in increased wound infections over wounds allowed to heal by secondary intent.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendation

There is one moderate-quality study of laceration repair for dog bite wounds. (1442) There are no quality studies for human or cat bite lacerations. A low-quality study compared infection rates and cosmetic outcomes of dog bite wounds repaired with monofilament suture versus allowing to heal by secondary intent. (1448) There was no difference found in infection rates. Patients were less satisfied with the cosmetic outcome in the non-sutured group. No statistically significant difference was found in infection rates in sutured wounds. (1442) These and several other studies considered in the antibiotic prophylaxis recommendation section have concluded that wound care (irrigation, debridement, cleansing) is the primary factor for preventing infection.

Evidence for the Treatment of Bite Laceration Repair

There is 1 moderate-quality RCT incorporated into this analysis. (1442) There is 1 low-quality RCT in Appendix 2. (1448)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library and Google Scholar without date limits using the following terms: Suture, Bites, Human, Animal, Dog, Cat, Bite, Torso, Upper Extremity, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 2 in Scopus, 3 in CINAHL, 5 in Cochrane Library, and 50 in Google Scholar. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 1 from Cochrane Library, 2 from Google Scholar, and 2 from other sources. Of the 6 articles considered for inclusion, 4 randomized trials and 2 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dire 1992 RCT</td>
<td>4.0</td>
<td>N = 185 (75 female/110 male)</td>
<td>Oral antibiotics (cephalexin, dicloxacillin or erythromycin) vs. no</td>
<td>One wound (1.1%) in antibiotic group and 5 (5.1%) in control group became infected (p = 0.212). No partial thickness wounds became infected. No difference in</td>
<td>&quot;Our results do not show a significant difference in wound infection rates among all low-risk dog bite wounds with or without oral antibiotic use. Routine prophylactic antibiotics would not seem</td>
<td>Sparse study details. No blinding or placebo. Wounds irrigated with</td>
</tr>
</tbody>
</table>
Follow-up Visits
There are no quality studies on the frequency and timing of follow-up visits for animal or human bites, or the effectiveness of wound care instruction and education. As the incidence of infection related to human and cat bites is much higher than for dog bites, there may be a stronger argument for having these patients present for wound check in 48-72 hours post injury. Follow-up for non-routine wounds should be dictated by the clinical presentation, or by other indications such as blood-borne pathogens protocols and concurrent injury management.

Prescription Medications
There are no recommendations for the use of prescription medications except as noted for antibiotics and blood-borne pathogens elsewhere.

Hand/Finger Osteoarthrosis

Diagnostic Criteria
For most purposes, a history and physical examination is sufficient but sometimes x-rays are used. X-rays are sometimes used in medicolegal situations to document the degree and extent of involvement. However, x-rays can be negative in those with osteoarthrosis as well as show evidence of disease among those asymptomatic.

Special Studies and Diagnostic and Treatment Considerations

X-RAYS
Recommendation: X-rays to Evaluate Hand Osteoarthrosis
X-rays are recommended to define objective evidence of the extent of hand osteoarthrosis.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – High

Rationale for Recommendation
There are no quality studies evaluating x-rays for hand osteoarthrosis. Most patients do not require x-rays for diagnosis and can be managed clinically. However, in some cases, x-rays are helpful and may assist in some patients in diagnosing and treating the condition. Thus, x-rays are recommended for hand osteoarthrosis.

Evidence for the Use of X-rays
There are no quality studies incorporated into this analysis.
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: X-ray, radiography, x-rays, hand and finger osteoarthrosis, joint disease, osteoarthritis, diagnostic, diagnosis, sensitivity, specificity, positive predictive value, negative predictive value, and predictive value of tests, efficacy, and efficiency. We found and reviewed 0 articles in PubMed, 36 in Scopus, 0 in CINAHL, 1 in Cochrane Library, and 378 from Google Scholar. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

**Initial Care**

Relative rest, splints, ice, and heat have been utilized for treatment of hand osteoarthrosis. (1449-1451) Uncontrolled trials have reported splinting reduced the need for hand surgery. (910, 1452) Exercises have been recommended as well. (1451, 1453-1458)

1. **Recommendation: Relative Rest and Self-application of Ice for Chronic Hand Osteoarthrosis**
   Relative rest and self-application of ice are not recommended for chronic hand osteoarthrosis.
   
   Strength of Evidence – Not Recommended, Insufficient Evidence (I)
   Level of Confidence – Low

2. **Recommendation: Splinting for Acute Flares or Chronic Hand Osteoarthrosis**
   Splinting is recommended for acute flares or chronic hand osteoarthrosis.
   
   Indications – Hand osteoarthrosis symptoms insufficiently treated with NSAIDs, acetaminophen, and/or topical medications. Prefabricated or custom-made orthoses may be utilized.
   
   Strength of Evidence – Recommended, Evidence (C)
   Level of Confidence – Low

3. **Recommendation: Exercise for Acute Flares or Chronic Hand Osteoarthrosis**
   Exercise is recommended for treatment of acute flares or chronic hand osteoarthrosis.
   
   Indications – Hand osteoarthrosis symptoms insufficiently treated with NSAIDs, acetaminophen, and/or topical medications.
   
   Frequency/Dose – One or 2 appointments for teaching home exercises. An additional subsequent appointment or two a few weeks later may be helpful to reinforce exercises and techniques.
   
   Strength of Evidence – Recommended, Evidence (C)
   Level of Confidence – Low

4. **Recommendation: Self-application of Heat for Acute Flares or Chronic Hand Osteoarthrosis**
   Self-application of heat is recommended for acute flares or chronic hand osteoarthrosis.
   
   Indications – Hand osteoarthrosis symptoms insufficiently treated with NSAIDs, acetaminophen, and/or topical medications.
   
   Frequency/Dose – Self-applications of heat, most commonly 15 to 20 minutes, 3 to 5 times a day.
   
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Low

**Rationale for Recommendations**

There are no quality studies of these treatments other than splinting and exercise. Relative rest does not appear to improve the disease in any other joint in the body (see Hip and Groin Disorders Guideline), and instead may promote debility. Thus, while not invasive, potential adverse effects may occur. Although it is generally low cost provided the patient is able to continue to work, it is not recommended.

All quality studies of splinting addressed thumb CMC/trapeziometacarpal OA. There is one quality study evaluating splinting versus no splinting that suggested modest benefits, (1459) although that trial may have been biased by a non-interventional control. Two crossover trials of different splints suggest a
flexible splint or support across the thumb CMC joint is superior to other, more rigid splint options. (1449, 1460) A fourth study compared two different exercise and splint regimens and found no differences. (1461) thus whether splints are beneficial compared with no splint is unclear. Splinting, particularly with a soft elastic support, is not invasive, has few adverse effects, is generally low cost and thus is recommended.

Most patients find heat superior to cryotherapies; however, there are no quality studies of either for treatment of hand OA. Heat may help with symptomatic relief, is not invasive, has no adverse effects, is not costly when self-applied and thus is recommended.

Exercise has not been widely investigated for treatment of hand OA, but has not been found to be harmful for hip or knee osteoarthritis patients (see Hip and Groin Disorders Guideline) and those patients obtain superior benefits with active exercise, and by inference may suggest rest is not appropriate for hand osteoarthrosis patients. One quality study found a home exercise program performed daily after a single 30-minute training session superior to educational controls for treatment of hand osteoarthrosis. (1458) An uncontrolled trial found strength training increased grip strength and reduced pain; (1456) however, a subsequent moderate-quality crossover trial by the same researcher did not find an exercise regimen of range of motion and strengthening exercises superior to another. (1457) As well, a study of combined exercises and splints failed to find one program superior. (1461) However, it is possible the trial by Rogers et al that evaluated exercises placed emphasis on flexibility exercises, thus biasing towards the null when additional trials may demonstrate clinically meaningful results. Exercies are not invasive, have low adverse effects and are low cost after an appointment or two for teaching purposes and are recommended.

Evidence for Splinting and Exercise for Hand Osteoarthrosis
There are 10 moderate-quality RCTs and randomized crossover trials incorporated into this analysis. (1571, 1579-1587) (Bani 13; Becker 13; Carreira 10; Villafane 13) There are 4 low-quality RCTs and 1 low-quality controlled clinical trial (1572, 1574, 1588-1590) (Boustedt 09; Adams 14; Weiss 00) in Appendix 2.

Rest:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Rest, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 26 articles in PubMed, 20 in Scopus, 169 in CINAHL, 1 in Cochrane Library, 100 in Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Ice:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Ice, Cryotherapy, Cold Therapy, Ice Pack, Self-Applied Ice, Cold Pack, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis, controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 12 articles in PubMed, 22 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 47,970 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 0 from other sources. Of the 1 articles considered for inclusion, 0 randomized trials and 1 systematic studies met the inclusion criteria.

Splinting:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: splints, splint, splinting; hand, fingers, thumb, metacarpus, osteoarthritis, osteoarthrosis, degenerative arthritis; controlled clinical trial,
controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 63 articles in PubMed, 73 in Scopus, 18 in CINAHL, 57 in Cochrane Library, 15,710 in Google Scholar, and 0 from other sources. We considered for inclusion 8 from PubMed, 2 from Scopus, 1 from CINAHL, 2 from Cochrane Library, 4 from Google Scholar, and 0 from other sources. Of the 17 articles considered for inclusion, 10 randomized trials and 10 systematic studies met the inclusion criteria.

Exercise:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Exercise, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 10 articles in PubMed, 182 in Scopus, 5 in CINAHL, 184 in Cochrane Library, 150 in Google Scholar, and 2 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 2 Google Scholar, and 2 from other sources. Of the 5 articles considered for inclusion, 4 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Study Type</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bani 2013</td>
<td>5.0</td>
<td>Prefabricated thumb splints (N=12) vs Custom made thumb splint (N=12) vs Control group (N=11)</td>
<td>The control group reported no significant differences in pain, function, or grip and pinch strength at week 4. At week 6 pinch strength significantly improved (p=0.000). At week 10 pinch strength (p=0.000) and pain (p=0.05) were the only parameters to improve. Functionality scores were significant for prefabricated splints (p=0.018) but not for custom splints (p=0.232). All were compared to the control group. At week 6 pain was significantly different for the custom splint (p=0.049). Grip strength was not improved. Pinch strength was not improved.</td>
<td>Both splints increased pain, pinch strength and function compared to baseline and control group. We found no evidence that splints improved grip strength as compared to control group. There were no significant differences in function and pinch in comparing the splints. Pain was the only significant difference. The custom made splint demonstrated better results in pain reduction. It appears that these splints are helpful in the short-term in early CMC OA, particularly for pain.</td>
<td>Data suggest comparable efficacy with respect to functional outcomes but custom made splints were reported to be more comfortable. Small sample size. Crossover design.</td>
</tr>
<tr>
<td>Rannou 2009</td>
<td>7.0</td>
<td>N = 112 (56 females/56 males) base of thumb OA (trapeziometacarpal) Age = Mean of 63 for custom-made group, Mean of 63.5 for control group</td>
<td>Custom-made splint vs. no splint. Nocturnal use only prescribed for 12 months. Pain VAS (baseline/ change at 1 mo/change at 12 months): splint (45.5±19.9/-10.1±3.0/ -22.2±3.2) vs. control (47.7±19.8/-10.7±3.3/-7.9±3.5), p = 0.89 at 1 month and p = 0.002 at 12 months. Similar results for Cochin Hand Function Scale, patient-perceived disability. Pinch strength at 12 months splint: -5.4±7.1 vs. -14.4±7.7 (p = 0.38).</td>
<td>&quot;For patients with base-of-thumb osteoarthritis, wearing a splint had no effect on pain at 1 month but improved pain and disability at 12 months..&quot;</td>
<td>Subjects had severe disease. Baseline duration of disease worse in controls. More co-interventions in controls may have lessened differences. Post-traumatic disease excluded. No differences at 1 month vs. positive differences at 3 months difficult to resolve, particularly with nocturnal splint use.</td>
</tr>
</tbody>
</table>
was improved in both prefabricated (p=0.000) and custom (p=0.000) groups. Functionality also improved in custom group (p=0.026).

At week 10 both splints reduced pain (p=0.000 for prefabricated, p=0.000 for custom). Pinch strength and functionality were significant (p=0.000) for both.

There were no significant differences between the two types of splints for functionality (p=0.136), grip strength (p=0.528), or pinch strength (p=0.651). At week 10 pain levels significantly differed (p=0.024).

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Diagnosis</th>
<th>Interventions</th>
<th>Results</th>
<th>Quote</th>
</tr>
</thead>
<tbody>
<tr>
<td>Becker 2013</td>
<td>RCT</td>
<td>4.5</td>
<td>N = 62 (48 female/14 male) with diagnosis of trapeziometacarpal arthrosis</td>
<td>Pre-fabricated neoprene Comfort Cool_ Thumb CMC Restriction Splint (N=32) vs Customized 3.2 mm thick thermoplastic hand-based thumb spica splint (N=30)</td>
<td>Comfort was the only statistically significant variable between the two splints (p=0.048) with participants preferring the neoprene splint. There were no detectable differences between the splints for areas of functionality, pain, pinch strength, satisfaction, and grip strength.</td>
<td>“When compared to custom-made thermoplastic splints, pre-fabricated neoprene hand-based thumb spica splints are, on average, more comfortable, less expensive, and as effective in treating trapeziometacarpal arthrosis.”</td>
</tr>
<tr>
<td>Carreira 2010</td>
<td>RCT</td>
<td>5.0</td>
<td>N = 40 (38 female/2 male) with osteoarthritis in trapeziometacarpal joint in dominant hand, clinical and radiological diagnosis,</td>
<td>Splint group, thermoplastic splint, used splint from day 1 for daily activities (N = 20) vs Control group, thermoplastic splint, used only during evaluations and</td>
<td>Between day 0 and day 90 there was a statistically difference in pain level between the groups (p=0.003). This was also observed at day 45 (p=0.013) and day 90 (p=0.002). In the splint group the pain was significantly reduced when comparing levels from day 0 to 45 (p &lt; 0.001) and 0 and 90 (p &lt; 0.001).</td>
<td>“Splint use during activities of daily living for patients with trapeziometacarpal osteoarthritis reduces pain, but does not alter function, grip strength, pinch strength or dexterity.”</td>
</tr>
</tbody>
</table>

Data suggest that functional splints used for OA of the trapeziometacarpal joint “may” reduce pain but do not alter function (grip strength, pinch strength or dexterity).
<table>
<thead>
<tr>
<th>Study</th>
<th>Type</th>
<th>Sample Size</th>
<th>Design Details</th>
<th>Outcome Measures</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weiss 2004</td>
<td>Randomized Crossover Trial</td>
<td>N = 25 (21 females/4 males); all with first CMC joint OA</td>
<td>Prefabricated neoprene splint vs. custom thermoplastic short opponens splint for 1 week each</td>
<td>Pain at rest baseline 5.42 (SEM 0.48). Pain after CMT: 3.59 (0.44) vs. PFN: 2.29 (0.33), p &lt;0.05. Pain with pinching favored PFN splint (p &lt;0.05). “Long-term” patient preference 72% PFN vs. 24% CMT.</td>
<td>No significant difference between the groups in scores of the first (p=0.524) and second (p=0.893) question of the DASH scores. Scores differed significantly for question three (p=0.382). No significant difference was observed between groups for dexterity, grip strength, and pinch strength. Comparing score differences from day 0 and day 180 the only significant difference was for pain without a splint (p=0.009).</td>
</tr>
<tr>
<td>Buurke 1999</td>
<td>Randomized Crossover Trial</td>
<td>N = 10 (10 females/0 males) with OA of 1st CMC joint</td>
<td>3 thenar eminence orthoses [supple elastic (Uriel 25), elastic with semi-rigid thumb busk (Gibortho ref. 6302) vs. semi-rigid polyethylene (Sporlastic 07051)]; 4 weeks each splint</td>
<td>Wearing comfort: Uriel 62.5 vs. Sporlastic 28.6 vs. Gibortho 23.3 (p &lt;0.05). Order of preference Uriel then Gibortho/Sporlastic. Pain ratings: Uriel 47±34 vs. Sporlastic 55±37 vs. Gibortho 48±31. No preference for pain ratings.</td>
<td>“Eight out of 10 patients prefer the permanent use of a TE orthosis. Six patients chose the supple elastic orthosis and two chose the semi-rigid orthosis.”</td>
</tr>
<tr>
<td>Villafñne 2013</td>
<td>RCT</td>
<td>N = 60 diagnosed with CMC joint OA</td>
<td>Control (N = 30) – Placebo group, received detuned ultrasound therapy. vs Experimental (N = 30) – Received multimodal treatment protocol for CMC joint OA-related pain.</td>
<td>The experimental group (3.7, CI 95% 2.4, 3.8) had a significant greater reduction in pain than the control group (0.3, CI 95% 0). An ANOVA revealed no significant differences in pressure pain threshold between both groups (F=0.44, P=.72). There was no significant difference between the two groups in regards to grip strength (F=1.2, P = .31)</td>
<td>“This study provides evidence that a multimodal intervention consisting of joint mobilization, neural mobilization, and exercise is beneficial to reduce pain in patients with CMC joint OA.”</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Splint vs. Another Splint</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Study</th>
<th>Type</th>
<th>Sample Size</th>
<th>Design Details</th>
<th>Outcome Measures</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weiss 2004</td>
<td>Randomized Crossover Trial</td>
<td>No mention of COI. Funded by grant from the AAHS.</td>
<td>Prefabricated neoprene splint vs. custom thermoplastic short opponens splint for 1 week each</td>
<td>Pain at rest baseline 5.42 (SEM 0.48). Pain after CMT: 3.59 (0.44) vs. PFN: 2.29 (0.33), p &lt;0.05. Pain with pinching favored PFN splint (p &lt;0.05). “Long-term” patient preference 72% PFN vs. 24% CMT.</td>
<td>No significant difference between the groups in scores of the first (p=0.524) and second (p=0.893) question of the DASH scores. Scores differed significantly for question three (p=0.382). No significant difference was observed between groups for dexterity, grip strength, and pinch strength. Comparing score differences from day 0 and day 180 the only significant difference was for pain without a splint (p=0.009).</td>
</tr>
<tr>
<td>Buurke 1999</td>
<td>Randomized Crossover Trial</td>
<td>No mention of COI or sponsorship.</td>
<td>3 thenar eminence orthoses [supple elastic (Uriel 25), elastic with semi-rigid thumb busk (Gibortho ref. 6302) vs. semi-rigid polyethylene (Sporlastic 07051)]; 4 weeks each splint</td>
<td>Wearing comfort: Uriel 62.5 vs. Sporlastic 28.6 vs. Gibortho 23.3 (p &lt;0.05). Order of preference Uriel then Gibortho/Sporlastic. Pain ratings: Uriel 47±34 vs. Sporlastic 55±37 vs. Gibortho 48±31. No preference for pain ratings.</td>
<td>“Eight out of 10 patients prefer the permanent use of a TE orthosis. Six patients chose the supple elastic orthosis and two chose the semi-rigid orthosis.”</td>
</tr>
<tr>
<td>Villafñne 2013</td>
<td>RCT</td>
<td>No sponsorship or COI.</td>
<td>Control (N = 30) – Placebo group, received detuned ultrasound therapy. vs Experimental (N = 30) – Received multimodal treatment protocol for CMC joint OA-related pain.</td>
<td>The experimental group (3.7, CI 95% 2.4, 3.8) had a significant greater reduction in pain than the control group (0.3, CI 95% 0). An ANOVA revealed no significant differences in pressure pain threshold between both groups (F=0.44, P=.72). There was no significant difference between the two groups in regards to grip strength (F=1.2, P = .31)</td>
<td>“This study provides evidence that a multimodal intervention consisting of joint mobilization, neural mobilization, and exercise is beneficial to reduce pain in patients with CMC joint OA.”</td>
</tr>
<tr>
<td>Exercise vs. Sham</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Study</th>
<th>Type</th>
<th>Sample Size</th>
<th>Design Details</th>
<th>Outcome Measures</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weiss 2004</td>
<td>Randomized Crossover Trial</td>
<td>No mention of COI or sponsorship.</td>
<td>Prefabricated neoprene splint vs. custom thermoplastic short opponens splint for 1 week each</td>
<td>Pain at rest baseline 5.42 (SEM 0.48). Pain after CMT: 3.59 (0.44) vs. PFN: 2.29 (0.33), p &lt;0.05. Pain with pinching favored PFN splint (p &lt;0.05). “Long-term” patient preference 72% PFN vs. 24% CMT.</td>
<td>No significant difference between the groups in scores of the first (p=0.524) and second (p=0.893) question of the DASH scores. Scores differed significantly for question three (p=0.382). No significant difference was observed between groups for dexterity, grip strength, and pinch strength. Comparing score differences from day 0 and day 180 the only significant difference was for pain without a splint (p=0.009).</td>
</tr>
<tr>
<td>Buurke 1999</td>
<td>Randomized Crossover Trial</td>
<td>No mention of COI or sponsorship.</td>
<td>3 thenar eminence orthoses [supple elastic (Uriel 25), elastic with semi-rigid thumb busk (Gibortho ref. 6302) vs. semi-rigid polyethylene (Sporlastic 07051)]; 4 weeks each splint</td>
<td>Wearing comfort: Uriel 62.5 vs. Sporlastic 28.6 vs. Gibortho 23.3 (p &lt;0.05). Order of preference Uriel then Gibortho/Sporlastic. Pain ratings: Uriel 47±34 vs. Sporlastic 55±37 vs. Gibortho 48±31. No preference for pain ratings.</td>
<td>“Eight out of 10 patients prefer the permanent use of a TE orthosis. Six patients chose the supple elastic orthosis and two chose the semi-rigid orthosis.”</td>
</tr>
<tr>
<td>Villafñne 2013</td>
<td>RCT</td>
<td>No sponsorship or COI.</td>
<td>Control (N = 30) – Placebo group, received detuned ultrasound therapy. vs Experimental (N = 30) – Received multimodal treatment protocol for CMC joint OA-related pain.</td>
<td>The experimental group (3.7, CI 95% 2.4, 3.8) had a significant greater reduction in pain than the control group (0.3, CI 95% 0). An ANOVA revealed no significant differences in pressure pain threshold between both groups (F=0.44, P=.72). There was no significant difference between the two groups in regards to grip strength (F=1.2, P = .31)</td>
<td>“This study provides evidence that a multimodal intervention consisting of joint mobilization, neural mobilization, and exercise is beneficial to reduce pain in patients with CMC joint OA.”</td>
</tr>
</tbody>
</table>

Data suggest combination therapy (ie. Joint mobilization, neural mobilization and exercise) is better than sham for pain treatment in patients with CMC joint OA.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>Participants</th>
<th>Intervention</th>
<th>Follow-up</th>
<th>Outcomes</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stamm 2002</td>
<td>RCT</td>
<td>N = 40 with hand osteoarthritis Mean Age of 60.5 years 35 Females, 5 Males</td>
<td>Control (N = 20) Vs Joint protection and exercise (JPE) group (N = 20) No mention of Follow up</td>
<td>At baseline, grip strength was slightly, but not significantly, higher in the control group (0.43 ± 0.21 in JPE group and 5.4 ± 0.16 in the control group in the right hand and he left hand yielded 0.44 ± 0.19 and 0.53 ± 0.19, respectively. After 3 months, grip strength significantly improved in the JPE for both hands (P &lt; 0.0001 for the right hand and P = 0.0005 for the left hand, when compared to baseline). There was no significant improvement for the control group (P = 0.2335 for the right hand and P = 0.1612 for the left hand).</td>
<td>&quot;Joint protection and hand home exercises, easily administered and readily acceptable interventions, were found to increase grip strength and global hand function.&quot;</td>
<td></td>
</tr>
<tr>
<td>Rogers 2009</td>
<td>Randomized Clinical Trial</td>
<td>N = 46 subjects at least 50 years or older with radiographic OA. Mean age of 75 years old. 40 Females, 6 Males</td>
<td>Exercise Group – 16 weeks of daily hand exercise intervention. Vs Sham Group – 16 weeks with OTC nonmedicated hand moisturizing lotion. No mention of group distribution. Time of follow up not mentioned.</td>
<td>Changes in AUSCAN sub-scales did not differ between the two treatment groups. Grip and pinch measures improved after exercise but not sham.</td>
<td>&quot;The results of this investigation found that while a home-based daily 16-week regimen of hand strength and range of motion exercises modestly improved grip and pinch strength, this benefit was not sufficient to see an improvement in self-reported hand physical function or pain&quot;</td>
<td></td>
</tr>
<tr>
<td>Wajon 2005</td>
<td>RCT</td>
<td>N = 40 (31 females/9 males) All with Stage I-III trapeziometacarpal OA</td>
<td>Thumb strap splint plus abduction exercises vs. short opponens splint plus pinch exercises. Splints custom thermoaplast. Exercises (5-10 reps, 3 sessions a day) added after 2</td>
<td>VAS pain scores (weeks 0/2/6): thumb strap plus abduction exercises (3.0±1.9/2.1±1.8/1.3±2.2) vs. opponens splint plus pinch exercises (2.9±2.2/1.8±1.8/0.9±1.2).</td>
<td>&quot;While both groups improved, neither regimen is superior to the other in patients with trapeziometacarpal osteoarthritis.&quot;</td>
<td></td>
</tr>
</tbody>
</table>

**Data suggest comparable efficacy. Splint worn full time, which may reduce ability to work or perform other activities.**
| Age = 59.7 for thumb strap group, 61.2 for short opponens splint | weeks of splinting. Total 6 weeks treatment. |  |  |
Follow-up Visits

Many patients require no follow-up appointments as the main thrust of the initial treatment generally focuses on securing the diagnosis and initiating treatment. Patients may require a few follow-up appointments, depending on severity and need for workplace limitations.

Medications

**NSAIDS/ACETAMINOPHEN**

NSAIDs and acetaminophen are widely used to treat pain associated with osteoarthrosis (OA), and are considered highly efficacious, although most studies evaluating their use lasted not longer than 6 weeks. (1463-1466) Most quality studies evaluated NSAIDs and acetaminophen in hip and/or knee OA patients and some evaluated low back pain patients (see Hip and Groin, and Low Back Disorders Guidelines). Few have evaluated hand osteoarthrosis patients. (1467-1469)

1. **Recommendation: NSAIDs for Acute Flares, Subacute, or Chronic Hand Osteoarthrosis**

   NSAIDs are moderately recommended to control pain associated with acute flares, subacute, or chronic hand osteoarthrosis. [Evidence is robust and strongly recommended for the treatment of osteoarthrosis in other body regions – Strongly Recommended, Evidence (A) (see Hip and Groin Disorders Guideline). Evidence is also present for efficacy of these agents for treating symptoms from OA flares (see Hip and Groin Disorders guideline).]

   Strength of Evidence – Moderately Recommended, Evidence (B)
   Level of Confidence – High

2. **Recommendation: NSAIDs for Acute Flares, Subacute, or Chronic Hand Osteoarthrosis**

   There is no recommendation for or against one NSAID over another as there is no consistent quality evidence that one NSAID is superior to another.

   Strength of Evidence – No Recommendation, Insufficient Evidence (I)
   Level of Confidence – High

3. **Recommendation: NSAIDs for Acute Flares, Subacute, or Chronic Hand Osteoarthrosis**

   There is no recommendation for or against enteric-coated vs. sustained release preparations as there is no consistent quality evidence demonstrating superiority of one or the other (see Hip and Groin Disorders guideline).

   Strength of Evidence – No Recommendation, Insufficient Evidence (I)
   Level of Confidence – High

4. **Recommendation: Acetaminophen for Acute Flares, Subacute, or Chronic Hand Osteoarthrosis**

   Acetaminophen (or the analog paracetamol) may be a reasonable alternative for treatment of osteoarthrosis pain – Recommended, Evidence (C), (1470, 1471) although quality evidence is available that documents these are consistently less efficacious in comparison with NSAIDs [Evidence (A)](1472-1479) and at least two quality trials with placebo comparisons have been negative including one with a large sample size of 779 patients. (1474, 1480) Yet, acetaminophen may be preferable for initial treatment of elderly patients and others with risks for gastrointestinal bleeding.

   Level of Confidence – High

Risk of adverse events from chronic NSAID use should be incorporated, especially including risk of gastrointestinal bleeding. Risk assessment should particularly include: prior history of gastrointestinal bleeding and source, length of treatment, age, smoking, diabetes mellitus and other medical factors.

It is strongly recommended that patients with greater risk should be considered for treatment with either acetaminophen, NSAID plus misoprostol, proton pump inhibitors or a COX-2 selective agent – Strongly Recommended, Evidence (A) (see Hip and Groin Disorders Guideline). (1463, 1464, 1481-1484) While COX-2 selective agents have generally been
recommended as either third- or fourth-line medications for routine use in osteoarthrosis patients, when there is a risk of gastrointestinal complications, they are often preferred. Proton pump inhibitors and misoprostol are also gastro-protective and have quality evidence of efficacy and are recommended (see Hip and Groin Disorders Guideline),(1481, 1482, 1484-1493) while there is substantially less evidence in support of sucralfate.(1492) COX-2 selective agents may still be used for those with contraindications to other medications, especially those with a history of gastrointestinal bleeding or past history of peptic ulcer disease. For patients at high risk of gastrointestinal bleeding, there is evidence that a combination of proton pump inhibitor plus COX-2 selective agent is efficacious.(1487) Should rofecoxib become available, it is suggested that it be considered as a fourth- or fifth-line medication for treatment of osteoarthrosis, likely paired with low-dose aspirin, and be positioned as a second-line medication for those with contraindications for the first- and second-line medication and in whom acetaminophen and celecoxib appear ineffective.

**Indications** – For hand osteoarthrosis patients, NSAIDs and acetaminophen are recommended for treatment. Over-the-counter agents may suffice and may be tried first.

**Frequency/Duration** – As-needed use may be reasonable for many patients. However, nearly all trials used scheduled doses. While not evaluated in hand OA patients, there is evidence that nocturnal dosing is superior for treatment of hip OA if the patient primarily has morning or nocturnal pain,(1494) although the study was of indomethacin and may only apply to shorter half-life agents as reproducibility of these findings and generalizability to other NSAIDs such as celecoxib with a longer half-life has not been shown.(1495)

**Indications for Discontinuation** – Resolution of hand pain, lack of efficacy, or development of adverse effects that necessitate discontinuation.

- **Strength of Evidence** – Recommended, Evidence (C)
- **Level of Confidence** – High

5. **Recommendation: NSAIDs for Patients at Risk for GI Adverse Effects**
Concomitant prescriptions of cytoprotective medications are strongly recommended for patients at substantially increased risk for gastrointestinal bleeding. There are four commonly used cytoprotective classes of drugs: misoprostol, sucralfate, double-dose histamine Type 2 receptor blockers (famotidine, ranitidine, cimetadine, etc.), and proton pump inhibitors (esomeprazole, lansoprazole, omeprazole, pantoprazole, rabeprazole). There is not generally believed to be substantial differences in efficacy for prevention of gastrointestinal bleeding,(1496) although evidence for sucralfate is limited. There also are combination products of NSAIDs/misoprostol that have documented reductions in risk of endoscopic lesions (see Hip and Groin Disorders Guideline).

**Indications** – For patients with a high-risk factor profile who also have indications for NSAIDs, cytoprotective medications should be considered, particularly if longer term treatment is contemplated. At-risk patients include those with a history of prior gastrointestinal bleeding, the elderly, diabetics, and cigarette smokers.

**Frequency/Duration** – As recommended.

**Indications for Discontinuation** – Intolerance, development of adverse effects, or discontinuation of NSAID.

- **Strength of Evidence** – Strongly Recommended, Evidence (A)
- **Level of Confidence** – High

6. **Recommendation: Discussion Regarding NSAIDs for Patients at Risk for Cardiovascular Adverse Effects**
Patients with known cardiovascular disease or multiple risk factors for cardiovascular disease should have the risks and benefits of NSAID therapy for pain discussed.

**Frequency/Dose** – If needed, NSAIDs that are non-selective are preferred over COX-2 specific drugs. In patients receiving low-dose aspirin for primary or secondary cardiovascular disease prevention, to minimize the potential for the NSAID to counteract the beneficial effects of aspirin, the NSAID should be taken at least 30 minutes after or 8 hours before the daily aspirin,(1497)
Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

7. **Recommendation: Acetaminophen or Aspirin for Patients at Risk for Cardiovascular Adverse Effects**

*Acetaminophen or aspirin is strongly recommended as the first-line therapy for patients with known or multiple risk factors for cardiovascular disease.*

*Frequency/Dose – See above.*

Strength of Evidence – Strongly Recommended, Evidence (A)
Level of Confidence – High

8. **Recommendation: Acetaminophen for Acute Flares, Subacute, or Chronic Hand Osteoarthrosis**

*Acetaminophen is recommended to control pain associated with acute flares, subacute, or chronic hand osteoarthrosis pain, particularly for patients with contraindications for NSAIDs.*

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – High

**Rationale for Recommendations**

There is abundant quality evidence that COX-1 and COX-2 NSAIDs improve pain and produce higher functional status among chronic osteoarthrosis patients (see Hip and Groin Disorders Guideline), and two quality studies included hand OA patients. There are a few studies of osteoarthrosis flares that also consistently document benefits, although not involving hand OA patients. There are many quality trials comparing the various NSAIDs, however, there is no consistent quality evidence of superiority of one over another or of one class over another class. There is one quality study suggesting that evening dosing of indomethacin results in better pain control, but the study has not been replicated(1494) and there is no similar result with the longer half life agent celecoxib.(1495) There is quality evidence that NSAIDs are less impairing than opioids, yet with comparable efficacy (see Chronic Pain and Low Back Disorders guidelines).

Quality evidence documents NSAIDs as superior to acetaminophen for symptomatic relief of OA (see Chronic Pain, Hip and Groin Disorders, and Low Back Disorders Guidelines).(1472-1478, 1498)

However, quality evidence also indicates higher rates of gastrointestinal adverse effects among NSAID users and generally lower overall adverse effects profiles for acetaminophen, providing rational for utilization of acetaminophen to treat some patients, particularly the elderly and others prone to GI complications.

**Evidence for the Use of NSAIDs and Acetaminophen for Hand Osteoarthrosis**

There is 1 high-quality crossover trial(1499) and 5 moderate-quality RCTs(1467, 1500-1503) incorporated into this analysis. There is 1 low-quality RCT in Appendix 2.(1468)

**Acetaminophen:**

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: antiinflammatory agents, non-steroidal, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis, NSAIDS, Acetaminophen; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 42 articles in PubMed, 58 in Scopus, 11 in CINAHL, 3 in Cochrane Library, 24081 in Google Scholar, and 0 from other sources. We considered for inclusion 4 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 4 from other sources. Of the 8 articles considered for inclusion, 7 randomized trials and 0 systematic studies met the inclusion criteria.
Gastrointestinal tolerability:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: antiinflammatory agents, non-steroidal, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis NSAIDS, gastrointestinal tolerability; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 1 articles in PubMed, 8 in Scopus, 1 in CINAHL, 13 in Cochrane Library, 5496 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 0 from other sources. Of the 1 article considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

Cardiovascular tolerability:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: antiinflammatory agents, non-steroidal, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis, NSAIDS, cardiovascular tolerability; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 1 article in PubMed, 6 in Scopus, 3 in CINAHL, 10 in Cochrane Library, 5425 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.

Acetaminophen, Aspirin, cardiovascular tolerability:
A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: antiinflammatory agents, non-steroidal, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis, Acetaminophen, Aspirin, cardiovascular tolerability; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 6 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 5199 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 0 from other sources. Zero articles met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pope 2004</td>
<td>8.5</td>
<td>N = 51 (gender not specified) with hip, knee or hand OA. Mean age 54 ± 2.4 years in N of 1 group, and 59 ± 2.3 years in conventional therapy</td>
<td>N of 1 group or of diclofenac 50mg plus misoprostol 200µg (n = 24) vs. Conventional therapy or placebo for 2 week durations for 6 months (n = 27).</td>
<td>In one group 11 patients preferred diclofenac, none preferred placebo, and 11 had no preference. NSAID appeared to be effective in 81% of patients.</td>
<td>“N of 1 trials were time-consuming in these patients and are more expensive, but with slightly better outcomes. In addition, NSAID seem to be effective in a majority of subjects with OA who have been uncertain of their benefit.”</td>
<td>Subjects at enrollment “uncertain the nonsteroidal anti-inflammatory drugs were helpful.” Results suggest NSAIDs are efficacious for majority of patients who were uncertain if they were effective.</td>
</tr>
<tr>
<td>Barthel 2010</td>
<td>7.5</td>
<td>N = 783 (80.2% female and 19.8% male) with radiographically confirmed hand osteoarthritis. Mean age was 63.9 years.</td>
<td>Diclofenac Group-Diclofenac sodium 1% gel (4 g total, 2 g to each hand) (n = 400) vs. Placebo Group-Vehicle consisted of isopropyl alcohol, propylene glycol, cocoyl caprylocaprate, mineral oil, ammonia solution, perfume cream 45/3, carbomer homopolymer type C, polyoxyl 20 cetostearyl ether, and purified water (n = 383). Follow-up for 8 weeks.</td>
<td>There was no significant difference between groups for VAS pain intensity at 8 weeks, (p &gt; 0.05). There were also no significant differences between groups for changes in AUSCAN scores, (p &gt; 0.05) and global rating of disease, (p &gt; 0.05).</td>
<td>“Pain relief correlated with improvements in physical function, stiffness, and global rating of disease in patients with hand OA, irrespective of treatment.”</td>
<td>Combined analyses of 2 prospective RCTs suggesting that pain from hand OA is directly related to function, stiffness, disease status, and improvements in any of above is not dependent upon active vs. placebo treatment. Anticipation of pain is what limits function.</td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Participants</td>
<td>Intervention</td>
<td>Outcomes</td>
<td>Results</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>--------------</td>
<td>-------------</td>
<td>----------</td>
<td>---------</td>
<td></td>
</tr>
<tr>
<td>Grifka 2004</td>
<td>RCT</td>
<td>N = 594 (490 female and 401 male) with symptomatic osteoarthritis. Mean age 61.9 years.</td>
<td>200 mg Group- Lumiracoxib 200mg od (n = 205) vs. 400mg Group-Lumiracoxib 400mg od (n = 193) vs. Placebo (n = 196). Follow-up at 4 weeks.</td>
<td>At week 2, 200mg had pain intensity decrease of 21.3 points, 400mg group had decrease of 21.1 and placebo was 12.5. Both Lumiracoxib groups showed significant difference for pain intensity vs. placebo (p &lt;0.001). But differences not significant between lumiracoxib groups. At week 4, respective decreases 28, 30 and 19.3. Global assessment of disease activity also decreased at week 4; 16.3, 20.9 and 9.4 in 200, 400 and placebo groups.</td>
<td>“Lumiracoxib 200 and 400 mg od were effective and well tolerated treatments for OA of the hand. Lumiracoxib significantly improved overall OA pain intensity in the target hand versus placebo, with a tolerability profile similar to placebo.”</td>
<td></td>
</tr>
<tr>
<td>Widrig 2007</td>
<td>RCT</td>
<td>N = 204 (147 female and 57 male) with hand osteoarthritis. Mean age was 64 years.</td>
<td>Ibuprofen Group- 4cm strip of gel, applied 4x a day for 3 weeks. (n = 99) vs. Arnica gel 4cm strip of gel applied 4x a day for 3 weeks (n = 105). Follow-up for 3 weeks.</td>
<td>Pain intensity and hand function very similar in both groups, (p &gt;0.05). No significant differences between groups for secondary outcomes of number of painful joints, intensity and duration of morning stiffness, (p &gt;0.05).</td>
<td>“Our results show that short-term use, up to three weeks, of arnica gel improves pain and function in hand OA, indistinguishably from ibuprofen gel.”</td>
<td></td>
</tr>
<tr>
<td>Gabay 2011</td>
<td>RCT</td>
<td>N=162 patients (42 males, 120 females) with hand OA. Mean age 63.9±8.5 years for CS group and 63.0±7.2 years for placebo group.</td>
<td>CS group: (n=80) 800 mg tablet of chondroitin sulfate with glass of water taken for 6 consecutive months. Vs. Placebo Group: Placebo same size tablet as CS group. (n=82)</td>
<td>Improvement in patient hand pain was significantly better for the CS group than the placebo group (p=0.016). The decrease in FIHOA score showed a similar pattern (p=0.008). Presence of erosive OA was significantly associated with higher FIHOA score (p=0.005), but not with global pain intensity (p=.75). Hand function improved significantly more in the CS groups than in the placebo group (p=0.008). There was a statistically significant difference.</td>
<td>“This study demonstrates that CS improves hand pain and function in patients with symptomatic OA of the hand and shows a good safety profile.”</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Sponsorship</td>
<td>N</td>
<td>Characteristics</td>
<td>Intervention</td>
<td>Outcomes</td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>-------------</td>
<td>---</td>
<td>----------------</td>
<td>-------------</td>
<td>----------</td>
</tr>
<tr>
<td>Smith 2010</td>
<td>RCT</td>
<td>Sponsored by past Peacock Trust. No COI.</td>
<td>7.0</td>
<td>N = 40 (35 female/5 male) with osteoarthritis in first carpometacarpal joint. Mean age 66.9 years.</td>
<td>Treatment group up to 20ml 0.5% sodium salicylate injected on any 1 occasion, given all in 1 large patch or divided between 2-4 smaller patches (n = 20) vs. Control Group: blunt 23-gauge probe pressed on skin over each patch as if patch injected (n = 20). Assessments at weeks 3, 7, and 13 years.</td>
<td>Patients assessed for pain, tenderness and disability using the VAS scale. The difference was 1.9 cm between the groups for VAS pain at the final follow-up in favor of the active group, (p = 0.007). The difference for VAS tenderness score was also significant in favor of the active group, 1.4 cm, (p = 0.02).</td>
</tr>
<tr>
<td>Lisse 2003</td>
<td>RCT</td>
<td>Sponsored by Merck &amp; Co., Inc. No COI.</td>
<td>7.0</td>
<td>N = 5,557 (3948 female/1609 male) with knee, hip, hand or spine OA. Mean age 63 years.</td>
<td>Rofecoxib 25mg a day (n = 2785) vs. Naproxen 500mg twice daily for 3 months. Double dummy (n = 2772).</td>
<td>Discontinuation due to adverse GI events lower in rofecoxib group (5.9% vs. 8.1%), RR = 0.74 (95% CI 0.60-0.92, p = 0.005). Similar findings in low-dose ASA takers. Less use of GI meds in rofecoxib group (9.1% vs. 11.2%, p = 0.014). Two perforations, ulcers or bleeding episodes in rofecoxib vs. 9 in naproxen (RR = 0.22, p = 0.038).</td>
</tr>
</tbody>
</table>

**Gastrointestinal Complications**

Small sample size. Data suggest injection of subcutaneous sodium salicylate effective in thumb OA vs. sham.
TOPICAL NSAIDS

Topical NSAIDs are used for treatment of hand osteoarthrosis (1504-1506) that include over-the-counter products.

Recommendation: Topical NSAIDs for Hand Osteoarthrosis

Topical NSAIDs are recommended to control pain associated with hand osteoarthrosis.

Indications – Mild, moderate, or severe hand osteoarthrosis.

Frequency/Duration – See manufacturer’s recommendation.

Indications for Discontinuation – Resolution, intolerance, adverse effects, or lack of benefits.

Strength of Evidence – Recommended, Insufficient Evidence (I)

Level of Confidence – Low

Rationale for Recommendation

Topical NSAIDs have been widely used. There are two quality studies are single-application studies precluding an evaluation is a regular treatment regimen, although they do suggest weak efficacy. (1504, 1505) Thus, there are not quality studies, and they appear to have quality evidence of efficacy for conditions with target tissue that is close to the skin, such as lateral epicondylitis (see Elbow Disorders Guideline) which is analogous to the skin in the dorsal hands. These medications are generally well tolerated, have few adverse effects, and are not costly when generic prescriptions are used, although they can be costly with name-brand prescription use over time. These medications are recommended.

Evidence for the Use of Topical NSAIDs for Hand Osteoarthrosis

There are 4 moderate-quality RCTs or crossover trials (1629, 1633, 1634, 1636) (Rothacker 94; Altman 09; Barthel 10) incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Topical NSAIDs, Topical non steroidal anti-inflammatory drug, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 32 in Scopus, 9 in CINAHL, 150 in Google Scholar, and 2 from other sources. We considered for inclusion 0 from PubMed, 0 from Scopus, 1 from CINAHL, 0 from Cochrane Library, 3 Google Scholar, and 2 from other sources. Of the 6 articles considered for inclusion, 4 randomized trials and 2 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rothacker 1994</td>
<td>RCT Crossover Trial</td>
<td>Sponsored in part by Thompson Medical Company, West Palm Beach, Florida. No mention of COI.</td>
<td>7.5</td>
<td>N = 50 (41 female/8 male) with hand OA. Mean age 66 years.</td>
<td>Trolamine salicylate 10% cream single application (n = 24) vs. Placebo single application (n = 25).</td>
<td>Changes in right hand pain severity (0/45/120 minutes): Trolamine salicylate (-0.2/-1.3/-1.4) vs. placebo (-0.2/-0.9/-1.1), p = 0.60, p = 0.08, p = 0.32. Mean change in pain relief scores at 45 minutes p = 0.047, with other times not significant.</td>
<td>“Trolamine salicylate has been shown to be both safe and effective in this single-application study of patients suffering from morning pain and stiffness associated with osteoarthritis in the hands.”</td>
<td>Ultra-short term study, single application. Suggests weak efficacy that is not long lasting.</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Diagnosis and Demographics</td>
<td>Intervention</td>
<td>Outcome Measures</td>
<td>Findings</td>
<td>Notes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>---</td>
<td>-----------------------------</td>
<td>-------------</td>
<td>------------------</td>
<td>----------</td>
<td>-------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Altman 2009</td>
<td>7.5</td>
<td>N = 385 diagnosed with OA in their primary hand. Mean age of 64.1 years old. 296 Females, 89 Males</td>
<td>Diclofenac Sodium Gel Group (N = 198) – Patients were given a topical 1% diclofenac sodium gel.  vs Vehicle Group (N = 187) – Patients were given a placebo gel. Follow up 1, 2, 4, 6, 8 weeks after gel given.</td>
<td>At week 8, the diclofenac sodium gel group stayed significantly superior to the vehicle group on the AUSCAN stiffness and functional indices (P&lt;0.048 and P&lt;0.017, respectively). Diclofenac sodium gel decreased pain intensity by 42.3%, total AUSCAN score by 35% and global rating of disease by 36.1%.</td>
<td>&quot;Topical diclofenac sodium gel was generally well tolerated and effective in primary hand OA.&quot;</td>
<td>Data suggest topical diclofenac gel was superior to placebo suggesting efficacy.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Barthel 2010</td>
<td>7.5</td>
<td>N = 783 diagnosed with primary hand OA by American College of Rheumatology criteria Mean age of 63.9 years old. 628 Females, 155 Males</td>
<td>Diclofenac Sodium Gel Group (N = 400) – Received 4g of 1% diclofenac sodium gel.  vs Vehicle Group (N = 383) – Received 4g of vehicle gel. Follow up 1, 2, 4, 6, 8 weeks after gel given.</td>
<td>Patients with at least 70% improvement from baseline score in VAS pain intensity had large mean improvements in AUSCAN pain, function, stiffness, and global rating of disease. Those that worsened also experienced a decrease in AUSCAN pain, function, stiffness, and global rating of disease. Change in VAS is correlated with AUSCAN pain, function, stiffness, and global rating of disease (P&lt;0.001).</td>
<td>&quot;Diclofenac sodium 1% gel is indicated for relief of OA pain in joints amenable to topical treatment, such as the hands and knees.&quot;</td>
<td>Data suggest pain relief correlates with improved hand function in OA patients irrespective of treatment.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rothacker 1998</td>
<td>6.5</td>
<td>N = 86 with hand OA.</td>
<td>Trolamine salicylate 10% cream vs. placebo. Single applications of each.</td>
<td>Sum of pain intensity differences scores: Trolamine salicylate -3.44 vs. -2.45, p = 0.072. Combined hands analysis p = 0.049.</td>
<td>&quot;10% trolamine salicylate cream was shown to be safe and effective for the temporary relief of minor pain and stiffness associated with osteoarthritis in the hands.&quot;</td>
<td>Data suggest efficacy over very short-term from single application.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**OPIOIDS – Oral, Transdermal, and Parenteral (Includes Tramadol)**
See Opioids recommendations in Carpal Tunnel Syndrome.

**COMPLEMENTARY/ALTERNATIVE THERAPIES**
Glucosamine, chondroitin sulfate, methyl-sulfonyl methane, diacerein (diacerhein, diacetylrhein), harpagophytum, avocado soybean unsaponifables, ginger, oral enzymes, and rose hips are often classified as complementary and alternative therapies that are sometimes used by patients for treatment of osteoarthrosis. (These are reviewed in detail in the Hip and Groin Disorders guideline.)

Copyright © 2016 Reed Group, Ltd.
1. **Recommendation: Capsaicin for Chronic Hand Osteoarthrosis or Acute Flares of Osteoarthrosis**  
Capsaicin is recommended for treatment of chronic hand osteoarthrosis or acute flares of osteoarthrosis.

**Indications** – Hand osteoarthrosis pain or acute flares (study has also included rheumatoid arthritis patients).(1507, 1508)

**Frequency/Duration** – Up to 4 times a day.

**Dose** – Fixed dose per manufacturer.

**Indications for Discontinuation** – Excessive burning of the skin or other intolerance. Not recommended for continual use, rather periods without use have been recommended.

**Strength of Evidence** – **Recommended, Evidence (C)**  
**Level of Confidence** – Low

2. **Recommendation: Yoga for Chronic Hand Osteoarthrosis or Acute Flares of Osteoarthrosis**  
Yoga is recommended for treatment of chronic hand osteoarthrosis or acute flares of osteoarthrosis.

**Indications** – Hand osteoarthrosis pain in motivated patients.

**Frequency/Duration** – Self-directed program after up to 8 supervised sessions.(1509)

**Indications for Discontinuation** – Intolerance, non-compliance.

**Strength of Evidence** – **Recommended, Insufficient Evidence (I)**  
**Level of Confidence** – Low

3. **Recommendation: Glucosamine, Chondroitin Sulfate, Methyl-sulfonyl Methane, Diacerein (Diacerhein, Diacetylrhein), Harpagophytum, Avocado Soybean Unsaponifiables, Ginger, Oral Enzymes, Nettle Leaf, or Rose Hips for Chronic Hand Osteoarthrosis or Acute OA Flares**  
There is no recommendation for or against use of glucosamine, chondroitin sulfate, methyl-sulfonyl methane, diacerein (diacerhein, diacetylrhein), harpagophytum, avocado soybean unsaponifiables, ginger, oral enzymes, nettle leaf, or rose hips for treatment of chronic hand osteoarthrosis or acute flares.

**Strength of Evidence** – **No Recommendation, Insufficient Evidence (I)**  
**Level of Confidence** – Low

**Rationale for Recommendations**

There is one quality study of capsaicin for treatment of these patients and it suggests benefits over a 4-week trial, thus it is recommended.(1507) There is one low-quality study of yoga,(1509) that suggested benefits. As yoga is not invasive, has few adverse effects, and is low cost, it is recommended for select, motivated patients.

There are more than 30 quality studies reviewed in the Hip and Groin Disorders Guideline. The largest volume of studies addresses glucosamine and chondroitin sulfate. This quality literature mostly addresses hip or knee osteoarthrosis or low back pain. Of the 5 quality, double-blinded studies that used x-rays for evaluation of glucosamine/chondroitin, three have documented delayed progression of joint space narrowing. There are 3 low-quality studies of chondroitin sulfate for treatment of hand arthrosis with one suggesting delay of hand x-ray changes.(1510) Yet, there are quality studies of knee and hip OA that have been both sizable and negative. However, glucosamine and chondroitin have problems with lack of standardization of doses. Nettle leaf(1511) has an additional problem of relative unavailability. This problem affects the other, less studied agents in this group of treatments. Consequently, although these agents are not invasive, have low adverse effects profiles, and may be costly over time, there is no recommendation for or against these agents.

**Evidence for the Use of Complementary and Alternative Therapies for Hand Osteoarthrosis**
There is 1 high-(1642) (Reeves 00) are 4 moderate-quality RCTs and crossover trials incorporated into this analysis.(1637, 1638, 1641, 1643) (Shin 13) There are 4 low-quality RCTs(1639, 1640, 1644, 1645) in Appendix 2.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Complementary therapy, alternative therapy, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 55 in Scopus, 6 in CINAHL, 70 in Cochrane Library, 150 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, 1 from Scopus, 1 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 7 from other sources. Of the 9 articles considered for inclusion, 9 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Capsaicin vs. Placebo</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>McCarthy 1992</td>
<td>RCT</td>
<td></td>
<td>5.0</td>
<td>N = 21 OA (14) and RA (7)</td>
<td>Capsaicin 0.075% vs. placebo QID for 4 weeks</td>
<td>VAS pain scores were (baseline vs. weeks 1/2/4): Capsaicin -10% vs. placebo -11%/-35% vs. -10%/-55% vs. -18% (p &lt;0.02) (graphic interpretations).</td>
<td>“[T]opical capsaicin is a safe and potentially useful drug for the treatment of painful OA of the hands.”</td>
<td>Blinding questionable. Suggests capsaicin reduces pain.</td>
</tr>
<tr>
<td>Schnitzer 1994</td>
<td>RCT</td>
<td></td>
<td>4.0</td>
<td>N = 59 Hand OA</td>
<td>Study began with all on capsaicin 0.025% vs. placebo and all QID dosing for 3 weeks, then BID for 6 weeks.</td>
<td>Capsaicin superior to placebo at Weeks 1 and 3 for pain responses (p = 0.046 and p = 0.018). Articular tenderness also favored capsaicin at all times except 6 weeks.</td>
<td>“[I]t may be prudent to taper the regimen gradually to avoid the decrease in pain relief seen with an abrupt decrease in dosage.”</td>
<td>Data suggest capsaicin effective, however study both decreased treatment frequency and randomized to placebo vs. treatment, thus somewhat limiting conclusions.</td>
</tr>
<tr>
<td><strong>Stinging Nettle vs. Non-Stinging Nettle</strong></td>
<td></td>
<td></td>
<td>7.0</td>
<td>N = 27 (23 female/4 male) with OA base of thumb or index finger. 2RA, 1 AS. Age range 45-82 years.</td>
<td>Stinging Urtica dioica (n = 13) vs. non-stinging nettle leaf Lamium album (n = 14).</td>
<td>VAS pain scores (baseline/post): stinging nettle (38.3/23.67) vs. non-stinging nettle (36.59/37.04), p = 0.026. Daily NSAID use: nettle (1.04/0.70) vs. non-stinging nettle (0.93/0.93), p &gt;0.05. Health assessment scores improved more with stinging nettle (p = 0.003).</td>
<td>“After one week’s treatment with nettle sting, score reductions on both visual analogue scale (pain) and health assessment questionnaire (disability) were significantly greater than with placebo.”</td>
<td>Success of blinding questionable. Patients applied the plant leaf themselves.</td>
</tr>
<tr>
<td>Reeves 2000</td>
<td>Prospective RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>8.0</td>
<td>N = 27 patients with osteoarthritis in the hands. Mean age of 64.2 years old. 16 Females, 11 Males</td>
<td>Dextrose Group (N = 13) – Received 0.5 mL of 10% dextrose or 0.075% xylocaine in bacteriostatic water. vs Control Group (N = 14) – Received 0.075% xylocaine in bacteriostatic water.</td>
<td>Flexion range improved significantly (P = 0.003) in dextrose treated joints compared to placebo-treated joints. After 6 months, the control group received dextrose injections and improved pain reduction from 18% to 54% in the average joints and 9.7% to 38% in total joint collection.</td>
<td>“Dextrose prolotherapy was clinically effective and safe in the treatment of pain with joint movement and range limitation in osteoarthritic joints.”</td>
<td>Data suggest at 12 months, ROM, pain level and PRWE and DASH scores equivalent. Patients in surgical group reported better grip strength throughout trial.</td>
</tr>
<tr>
<td>Study</td>
<td>Authors</td>
<td>Year</td>
<td>Sample Size</td>
<td>Intervention</td>
<td>Follow-up</td>
<td>Results</td>
<td>Conclusion</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>---------</td>
<td>------</td>
<td>-------------</td>
<td>--------------</td>
<td>-----------</td>
<td>---------</td>
<td>-----------</td>
<td></td>
</tr>
<tr>
<td>Shin 2013</td>
<td>RCT</td>
<td>7.0</td>
<td>N = 86 patients fulfilled the American College Board of Rheumatology criteria for hand OA. Mean age of 57.8 years old. 83 Females, 3 Males</td>
<td>Diacerein Group (N=42) – Received Diacerein 50 mg BID or 12 weeks vs Placebo Group (N=44) – Received placebo BID for 12 weeks</td>
<td>Follow up 4 and 12 weeks after initial enrollment.</td>
<td>There are no significant difference in change in AUSCAN pain score at 4 weeks (Diacerein vs placebo, P = 0.507). Diacerein was significantly improved (P = 0.004) for the physician global assessment. Adverse events occurred 38 (90%) times in the diacerein group and 29 (67%) in the placebo group.</td>
<td>“The results of this trial indicate that the safety profile of diacerein 50 mg BID is acceptable, although the regimen may be unsuccessful in controlling the symptoms of hand OA.”</td>
<td>Data suggest comparable efficacy between groups.</td>
</tr>
</tbody>
</table>
LOW-LEVEL LASER THERAPY
Low-level laser therapy has been used for treatment of hand osteoarthrosis patients, although the evidence has been noted to conflict.(1514-1516)

Recommendation: Low-Level Laser Therapy for Hand Osteoarthrosis
Low-level laser therapy is moderately not recommended for treatment of hand osteoarthrosis. (1517) Low-level laser therapy is not invasive and has low adverse effects, but it is costly and in the absence of efficacy is therefore not recommended.

Strength of Evidence – Moderately Not Recommended, Evidence (B)
Level of Confidence – Moderate

Rationale for Recommendation
There is one high-quality study that suggests low-level laser therapy is ineffective for treatment of hand osteoarthrosis. (1517) Low-level laser therapy is not invasive and has low adverse effects, but it is costly and in the absence of efficacy is therefore not recommended.

Evidence for the Use of Low-Level Laser Therapy for Hand Osteoarthrosis
There is 1 high-quality RCT incorporated in this analysis. (1517)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Low Level Light Therapy, LLLT, Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 9 articles in PubMed, 18 in Scopus, 1 in CINAHL, 0 in Cochrane Library, 150 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 0 from other sources. Of the 1 articles considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brosseau 2005</td>
<td>RCT</td>
<td>Sponsored by Ontario Arthritis Society, Ontario Ministry of Health and Long-Term Care, University Research Chair, and Ministry of Human Resources. No mention of COI.</td>
<td>9.0</td>
<td>N = 88 patients diagnosed with OA. Mean age of 65.7 years old. 69 Females, 19 Males</td>
<td>Low Level Laser Therapy Group (N = 42) – Received inactive LLLT vs Sham Low Level Laser Therapy Group (N = 46) – Received Gallium Aluminum Arsenide LLLT</td>
<td>Follow-up 6 and 18 weeks after last treatment of LLLT. There was no significant difference in VAS scores and morning stiffness. Grip strength significantly improved for participants in the active LLLT group (P = 0.041) and a significant reduction in finger distance between thumb and the base of the fifth metacarpal (P = 0.011). No significant differences were found in other outcomes.</td>
<td>“LLLT is no better than placebo at reducing pain, morning stiffness, or improving functional status for OA-hand patients.”</td>
<td>Suggests LLLT not effective.</td>
</tr>
</tbody>
</table>
Injections
INTRAARTICULAR INJECTIONS
Intraarticular glucocorticosteroid and hyaluronidate injections are sometimes performed to attempt to deliver medication with minimal systemic effects to the arthritic joint, particularly when acetaminophen and NSAIDs have failed. These injections are generally performed without fluoroscopic or ultrasound guidance in the distal upper extremity.

1. Recommendation: Intraarticular Glucocorticosteroid Injection for Subacute or Chronic Hand Osteoarthrosis

Intraarticular glucocorticosteroid injections are recommended for the treatment of subacute or chronic hand osteoarthrosis.

Indications – Moderately severe or severe hand osteoarthrosis pain with insufficient control with NSAID(s), acetaminophen, and potentially splinting and/or exercise. Its usual purpose is to gain sufficient relief to either resume medical management or to delay operative intervention.

Frequency/Duration – One (1) injection should be scheduled, rather than a series of 3. Various medications have been used, as well as adjuvant anesthetic agents. There are no head-to-head comparisons in quality studies of different medications to ascertain optimum medication(s).

Dose – Various doses have been utilized without evidence to identify an ideal dose for hand or phalangeal joints.

Indications for Discontinuation – A second glucocorticosteroid injection is not recommended if the first has resulted in significant reduction or resolution of symptoms. If there has not been a response to a first injection, there is generally less indication for a second. If the physician believes the medication was not well placed and/or if the underlying condition is so severe that one steroid bolus could not be expected to adequately treat the condition, a second injection may be indicated. If placement is thought to be difficult, ultrasound or fluoroscopic guidance may be particularly indicated for a second injection. In patients who respond with a pharmacologically appropriate several weeks of temporary partial relief of pain, but who then have worsening pain and function and who are not (yet) interested in surgical intervention, a repeat injection is an option. There are not believed to be benefits beyond approximately 3 injections in a year. Patients requesting a 4th injection should have reassessment of non-operative management measures and be counseled for possible surgical intervention.

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – Low

2. Recommendation: Intraarticular Hyaluronate Injection for Subacute or Chronic Hand Osteoarthrosis

Intraarticular hyaluronate injections are recommended for the treatment of subacute or chronic hand osteoarthrosis.

Indications – Hand osteoarthrosis pain with insufficient control with NSAID(s), acetaminophen, and potentially splinting and/or exercise. Its usual purpose is to gain sufficient relief to either resume medical management or to delay operative intervention.

Frequency/Duration – Number and frequency of injections are unclear (one trial found no differences between 1, 2, or 3 injections). Most physicians perform 3 injections.

Dose – See manufacturer’s recommendations.

Indications for Discontinuation – Sufficient relief to not require additional injection(s), failure to improve, or allergic reactions.

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – Low

Rationale for Recommendations
There are several quality studies for treatment of hand osteoarthrosis with glucocorticosteroids. However, the studies conflict regarding the length of benefits. However, nearly all studies have suggested benefits. No studies have suggest prolonged benefits with more than approximately 3 months benefits, thus this intervention is a short to intermediate term intervention.
Optimal glucocorticoid doses and preferable adjuvant anesthetic agents are unclear. These injections are invasive, have low adverse effects, and are moderately costly. They are recommended as an option for treatment of hand OA patients particularly after inadequate results from NSAID trials or other non-operative interventions.

There are a few quality studies of hyaluronate injections for treatment of hand osteoarthrosis. These suggest benefits. Duration of improvement is uncertain, although one trial suggested pain relief as far as 26 weeks. These injections are invasive, have moderate adverse effects and are costly. In select cases where other treatments have failed, these injections are recommended.

Evidence for the Use of Intraarticular Injections
There is 1 high-(1527) and 3 moderate-quality RCTs(1528-1530) incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Intraarticular Injections, glucocorticosteroid, hyaluronate injection; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 22 articles in PubMed, 9 in Scopus, 3 in CINAHL, 0 in Cochrane Library, 9928 in Google Scholar, and 0 from other sources. We considered for inclusion 7 from PubMed, 1 from Scopus, 1 from CINAHL, 0 from Cochrane Library, 0 Google Scholar, and 0 from other sources. Of the 9 articles considered for inclusion, 6 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wenham 2012</td>
<td>RCT</td>
<td>Sponsored by Arthritis Research UK. E.H. is part-funded by Arthritis Research UK. COI, A.J.G. has received payment from GE ultrasound division for lecturing. All other authors have declared no COI.</td>
<td>8.5</td>
<td>N = 70 (57 female/13 male) with symptoms of OA hand pain. Mean age for placebo group was 61.1 years and for PNL group 61.9 years.</td>
<td>Prednisolone (PNL) 5mg (n = 35) vs. placebo capsules (n = 35). Follow-up at 4 and 12 weeks.</td>
<td>At 4 weeks the adjusted mean reduction in pain VAS was 19.9mm (PNL group) and 16.8mm (placebo group) (p = 0.54). There were no statistically significant differences in VAS, Australian/Canadian Hand Osteoarthritis Index or joint counts between placebo and PNL groups at 4 or 12 weeks.</td>
<td>“This is the first randomized controlled trial of low-dose corticosteroid alone for painful hand OA, which demonstrated that short-term low-dose oral PNL is not an effective analgesic treatment for hand OA.”</td>
<td>Data suggest PNL compared to placebo is not effective for hand OA pain.</td>
</tr>
<tr>
<td>Spolidoro Paschoal Nde 2015</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>6.0</td>
<td>N=60 patients (2 males, 58 females) with diagnosis of OA involving PIP or DIP joints. Mean age: 60.7 years(±8.2)</td>
<td>TH/LD Group: One injection 20 mg/ml TH and 2% lidocaine w/o epinephrine. IAI administered in 0.3 ml dose of TH for PIP and 0.2 ml of TH for DIP both with 0.1 ml of 2% LD. Vs. LD Group: One injection of lidocaine. IAI administered with only 0.1 ml 2% LD w/o epinephrine in IP joint.</td>
<td>Significant difference was observed for VASm and physician VASs variables. The TH/LD group showed better performance than LD group for VASm and physician VASs (p=0.014 and 0.022 respectively) from first week to end of study. No statistical significance between groups or adverse effects.</td>
<td>“The IAI with TH/LD has been shown to be more effective than the IAI with LD for pain on movement and joint swelling in patients with OA of the IP joints. Regarding pain at rest, there was no difference between groups.”</td>
<td>Data suggest IAI with TH/LD better than IAI with LD foredema reduction and increased movement.</td>
</tr>
<tr>
<td>Figen-Ayhan, 2009</td>
<td>RCT</td>
<td>No mention of industry sponsorship. No COI.</td>
<td>6.0</td>
<td>N= 33 patients (0 males, 33 females)(66 hands) with thumb base osteoarthritis (OA). Mean age was 62.6 years.</td>
<td>Hylan Group- Injection of Hylan G-F 20 solution (N = 33 Hands) vs Saline Group- Injection of saline (N = 33 Hands). Follow-up at 6 and 24 weeks.</td>
<td>Both groups significantly improved from baseline to 6 weeks in pulp to pulp pinch, tripod pinch and key pinch strength (p≤ 0.04). Both groups also showed a significant improvement in VAS pain scores between baseline and the 6th week (p≤ 0.02). The Hylan group also showed significant improvement with regards to VAS pain score at the 24th week (p= 0.01). This difference was not</td>
<td>“Although short-term placebo analgesic effect could not be ignored, intra-articular hylan was effective on pain, pinch strength, and function at the 24th week.”</td>
<td>Each patient received both treatments (one on each hand). The hands were randomized. Data suggest HA may be an alternative method to improve thumb OA as there was improvement of function and grip strength at 24 weeks compared to placebo.</td>
</tr>
<tr>
<td>Study</td>
<td>Score</td>
<td>Patients</td>
<td>Intervention</td>
<td>Outcomes</td>
<td>Key Findings</td>
<td>Notes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------------------------------------------------------------------</td>
<td>-------</td>
<td>----------</td>
<td>--------------</td>
<td>---------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------</td>
<td>-------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Monfort, 2015 RCT</td>
<td>6.0</td>
<td>N= 88 patients (11 males, 77 females) with osteoarthritis of the thumb. Mean age was 62.8 years.</td>
<td>Hyaluronic acid Group-Ultrasound-guided intra-articular treatment with hyaluronic acid. Total of 3 injections (5 mg) (N=48) Vs. Betamethasone Group-ultrasound-guided intra-articular treatment with betamethasone. 1.5 mg of betamethasone disodium phosphate and 1.5 mg of betamethasone acetate. (N=40) Follow-up at 7, 14, 30, 90 and 180 days.</td>
<td>There was a significantly larger change from the baseline FIHOA score in the HA group vs. the Betamethasone group at Day 14; 3 vs. 1 (p&lt;0.05), Day 90; 5.5 vs. 1 (p&lt;0.005) and Day 180; 7 vs. 2 (p&lt;0.005). There was also a significantly larger change from the baseline VAS pain score in the HA group vs. the Betamethasone group at Day 180; 3.12 vs. 1.60 (p&lt;0.05).</td>
<td>&quot;Both hyaluronic acid and betamethasone were effective and well-tolerated for the management of rhizarthrosis. Hyaluronic acid was more effective over time and more efficiently improved functionality and pain in patients with more severe symptoms.&quot;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bahadir, 2009 RCT</td>
<td>5.0</td>
<td>N= 40 patients (0 males, 40 females) with stage II or III trapeziometacarpal joint osteoarthritis. Mean age was 62.9 years.</td>
<td>Steroid group- One injection of 20 mg triamcinolone acetonide (N=20) Vs. Hyaluronate group- 3 injections of 5 mg sodium hyaluronate at 1-week intervals. (N=20) Follow-up for 12 months.</td>
<td>The VAS pain score decreased significantly for both groups after treatment throughout the 12 months. The VAS pain score was significantly lower in the Steroid group compared to the hyaluronate group at 1 month after treatment; 3.1 vs. 4.7 (p=0.003), and at 6 months; 3.5 vs. 5.7 (p=0.002). The difference was not significant at 12 months; 4.9 vs. 6.0 (p=0.128).</td>
<td>&quot;Our findings showed that both intra-articular injection of steroid and sodium hyaluronate are effective in trapeziometacarpal joint osteoarthritis. However the steroid injection was found to be superior to sodium hyaluronate injection in reducing pain and improving hand function.&quot;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stahl 2005 RCT</td>
<td>4.5</td>
<td>N=52 patients (6 males, 46 females) with trapeziometacarpal MA Group: 40 mg Methylprednisolone intraarticular injection Vs.</td>
<td>Significant improvement in pain for both groups (p&lt;0.001). No significant difference in improvement of VAS values between 2</td>
<td>&quot;Steroids and hyaluronate injections were found effective in reducing pain. Hyaluronate was more</td>
<td>Data suggest comparable efficacy with some improved function in HA group.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No mention of sponsorship or COI.</td>
<td>joint grade II arthritis. Mean age: 62 years.</td>
<td>SH Group: 15 mg Sodium hyaluronate intraarticular injection groups. Grip strength improved significantly for steroid group. SH group showed improvement in grip after 6 months and pinch and PPT after 3 months.</td>
<td>effective in the improvement of some aspects of fine hand function.&quot;</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
PROLOTHERAPY INJECTIONS
Recommendation: Prolotherapy Injections for Subacute or Chronic Hand Osteoarthrosis

There is no recommendation for or against the use of prolotherapy injections for treatment of subacute or chronic hand osteoarthrosis.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation

Prolotherapy injections are invasive as they require numerous, repeated injections in phalangeal joints. The magnitude of the purported benefits is modest. The results of Reeves’ 2000 study suggesting some benefits compared with placebo injections(1531) needs to be replicated, including with a larger sample size, evaluation of functional outcomes and with sufficient follow-up duration to allow for adequate assessment of risks and benefits of these procedures prior to a recommendation in favor of this treatment.

Evidence for the Use of Injections for Hand Osteoarthrosis

There are 2 high-(1642, 1654) quality and 5 moderate-quality RCTs and crossover trials incorporated into this analysis.(1651-1653, 1655, 1664) (Jahangiri 14) There is 1 low-quality RCT in Appendix 2.(1656)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Prolotherapy Injections OR Proliferative Therapy AND Hand, Fingers, Thumb, Metacarpus, Osteoarthritis, Osteoarthrosis; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 2 in Scopus, 1 in CINAHL, 2 in Cochrane Library, 997 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 1 from Scopus, 1 from CINAHL, 0 from Cochrane Library, 1 Google Scholar, and 4 from other sources. Of the 8 articles considered for inclusion, 8 randomized trials and 2 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meenagh 2004</td>
<td>RCT</td>
<td>No mention of sponsorship or COI</td>
<td>8.5</td>
<td>N = 40 (36 female/4 male) with CMC joint OA. Age range 41-71.</td>
<td>Triamcinolone hexacetonide 0.25mL, 5mg (n = 20) vs. sterile saline. Fluoroscopically guided injections (n = 20).</td>
<td>VAS pain changes (4/12/24 weeks): placebo (18.5/23.3/14.0) vs. steroid (10.5/3.5/0.0), NS. Patient and physician global assessments improved in both groups at 4-12 weeks.</td>
<td>“No clinical benefit was gained from intra-articular steroid injection to the CMCJ in moderate to severe osteoarthritis compared with placebo injection.”</td>
<td>VAS pain ratings suggest trend towards modest pain reductions especially at 4 weeks, but none at 24 weeks. Suggests steroid injection relatively ineffective.</td>
</tr>
</tbody>
</table>

### Different Types of Glucocorticosteroid Injections (No Placebo)

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jalava 1983</td>
<td>Crossover Trial</td>
<td>No mention of sponsorship.</td>
<td>5.0</td>
<td>N = 24 (12 females, 12 males); 120 injected DIP, PIP joints, yet study describes RA patients; Mean Age 48.6 (24-73)</td>
<td>Triamcinolone hexacetonide vs. methyl-prednisolone 0.2-0.3mL/joint</td>
<td>Follow-Up at baseline, week 1, 4, 12, and 24.</td>
<td>Effect at 6 months: TH: 21.0% Unchanged; 3.5% Worse, p&lt;0.05. MP: 32.0 % unchanged; 10.0 % worse</td>
<td>“All injections produced clinically significant effects. There were no significant differences between the two treatment groups at the start of the treatment, but Crossover trial. States RA patients, but DIP/PIP joint injections. Multiple injections in multiple digits of same patient. No placebo group, thus conclusion on benefit for all not clearly clear.”</td>
</tr>
</tbody>
</table>
Glucocorticosteroid vs. Viscosupplementation Injections

Fuchs 2006
RCT
Sponsored by TRB Chemedica AG, Richard-Reitzner-Allee. No COI.

6.0
N = 56 (11 males/45 females) thumb CMC joint OA; Median Age, Group 1: 59.5 Group 2: 61.0
Three intraarticular injections of: Group 1 (N=28) Sodium hyaluronic acid (SH) 10mg vs. Group 2 (N=28) triamcinolone acetonide TA 10mg injections. Imaging not used; Follow Up at baseline, 3, 14, and 26 weeks.

VAS pain assessment (visits 1/3/5/6/7): SH (65.5/54.0/34.0/35.0/30.0) vs. TA (63.5/46.0/20.0/22.0/45.5).

“A single course of three SH injections is effective in relieving pain and improving joint function in patients with OA of the CMC joint of the thumb. Although in comparison with triamcinolone its effects are achieved more slowly, the results indicate a superior long-lasting effect of hyaluronan at 6 months after end of treatment period.”

No placebo group. Data suggest effect of steroid largely gone at 6 months, but not for visco-supplementation.

Viscosupplementation vs. Glucocorticosteroid vs. Placebo

Heyworth 2008
RCT
Sponsored by by a grant from Wyeth-Ayerst Pharmaceuticals and Genzyme Corporation. No mention of COI.

7.5
N = 60 (52 female/2 male) with basal joint OA. Mean age 63 ± 1.
(2) 1-mL injections of hylan G-F 20 1 week apart (n = 20) vs. Steroid1mL betamethasone (n = 22) vs. 2 placebo saline injections (n = 18). All received 2 injections, 1 week apart.

Data graphically presented; suggest grip strengths worse for saline than other 2 groups. However, not statistically significant between groups. Within groups, steroid superior at Weeks 2 and 4 to baseline and Hylan better at Weeks 2, 4, 12, 26 compared with baseline. No between-group VAS differences, but lower VAS pain compared with baseline for controls and steroid at Weeks 2 and 4, however for hylan, reductions were at Weeks 2, 12, 26 compared with baseline.

“There were no statistically significant differences among hylan, steroid, and placebo injections for most of the outcome measures at any of the follow-up time points. However, based on the durable relief of pain, improved grip strength, and the long-term improvement in symptoms compared with preinjection values, hylan injections should be considered in the management of

Trend towards Hylan relief lasting longer than glucocorticosteroid injection. States no baseline difference but stats for age are dissimilar. Dropout rate unclear.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Sponsorship</th>
<th>Participants</th>
<th>Intervention</th>
<th>Endpoints</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roux 2007</td>
<td>2007</td>
<td>No mention of sponsorship or COI.</td>
<td>N = 42 (4 males/38 females); Mean Age 64.8 ± 8.0</td>
<td>1 ml sodium hyaluronidate (Sinovial) 1 injection vs. 2 injections vs. 3 weekly injections using image intensifier</td>
<td>1 injection VAS (1 month, 3 months): 58.4±16.2, 43.1±22.8; 2 injections: 54.6±18.9, 39.5±28.6; 3 injections: 60.1±17.0, 29.8±21.9</td>
<td>&quot;No significant differences were found between each group over the study period for pain relief and function. But the intra groups analysis results show that intra-articular sodium hyaluronidate injections into the carpometacarpal joint of the thumb in osteoarthritis can be efficacious on pain and functionality.&quot;</td>
</tr>
<tr>
<td>Reeves 2000</td>
<td>2000</td>
<td>No mention of sponsorship or COI.</td>
<td>N = 27 (16 female/11 male) with 150 joints DIP, PIP and thumb CMC joint OA. Age for dextrose and control group: 64.5 (9.2) and 63.9 (9.4).</td>
<td>0.5ML of 10% dextrose plux 0.075% xylocaine vs. 0.075% xylocaine injections into medial and lateral aspects of each joint. Injections at 0, 2, 4 months</td>
<td>VAS after 3 injections improved 37% in active treatment vs. 18% controls (NS). Pain with rest and grip non-significant trend towards dextrose. Pain with movement improved with dextrose (59 to 67 vs. 57 to 48 in controls) (p = 0.027)</td>
<td>&quot;Dextrose prolotherapy was clinically effective and safe in the treatment of pain with joint movement and range limitation in osteoarthritic finger joints.&quot;</td>
</tr>
<tr>
<td>Jahangiri 2014</td>
<td>2014</td>
<td>No sponsorship or COI.</td>
<td>N=60 (44 female/16 males) patients with osteoarthritis in the first carpometacarpal joint (CMC); Mean Age 63.6 ± 9.7.</td>
<td>Local corticosteroid (LC) group, had placebo injections of 1 ml 0.9 % saline were administered (for masking) followed by a single dose of 40 mg methylprednisolone acetate (0.5 ml) mixed with 0.5 ml of 2 % lidocaine in the 3rd month Vs. Group 2: Dextrose Prolotherapy (DX) group, had 0.5 ml of 20 % DX mixed with 0.5 ml of 2 % lidocaine was injected; Follow-Up at baseline 1, 2, and 6 months.</td>
<td>LC - DX difference, Hand Assessment Questionnaire Disability Index (HAQ-DI) scores (Mean Difference (95% CI)), two months: 1.0 (0.2-1.9) (p=0.01). 6 months: 1.0 (0.2 – 1.8) (p=0.01). Pain, Visual Analogue Scale (VAS), 2 months: 1.0 (0.1-2.0) (p=0.01). 6 months: 1.1 (0.2-2.0) (p=0.02). Pinching, 1 month: 2.9 (0.9-4.9) (p=0.005). Both groups improved significantly within themselves and was significant in all three categories listed above.</td>
<td>&quot;Both LC and DX can relieve pain and suppress inflammatory processes. Furthermore, DX has been suggested to strengthen soft tissue too. There are some reports indicating improvement in ligament laxity after DX prolotherapy.&quot;</td>
</tr>
</tbody>
</table>
Various surgical procedures are utilized to treat patients with hand osteoarthrosis. Among these are arthrodesis, arthroplasty and various other reconstructive procedures, although many have been developed and utilized to primarily treat patients with rheumatoid arthritis.

1. **Recommendation: Reconstructive Surgery for Select Patients with Trapeziometacarpal Arthrosis**
   
   Reconstructive surgery is recommended for treatment of select patients with trapeziometacarpal arthrosis.
   
   **Strength of Evidence** – Recommended, Evidence (C)
   
   **Level of Confidence** – Low

2. **Recommendation: Trapeziectomy with Ligament Reconstruction and Tendon Interposition for Thumb CMC Joint Osteoarthritis**
   
   Trapeziectomy with ligament reconstruction and tendon interposition (LRTI) is not recommended for treatment of thumb CMC joint osteoarthritis.
   
   **Strength of Evidence** – Not Recommended, Evidence (C)
   
   **Level of Confidence** – Low

3. **Recommendation: Fusion for Select Patients with Hand Osteoarthrosis**
   
   Fusion is recommended for treatment of select patients with hand osteoarthrosis.
   
   **Strength of Evidence** – Recommended, Insufficient Evidence (I)
   
   **Level of Confidence** – Low

Rationale for Recommendation

There are several moderate-quality studies evaluating surgery for hand osteoarthrosis, all of which concern the basal thumb joint (trapeziometacarpal joint). (1532-1534, 1536, 1537, 1540, 1543-1545, 1549, 1550) There are a few quality studies of surgery for rheumatoid arthritic joints, such as MCP joint replacement. (1551-1554) However, these are beyond the scope of this document. Most of the OA studies address a comparison between trapeziectomy and trapeziectomy with ligament reconstruction or arthroplasty versus tendon interposition arthroplasty. Regardless, ligament reconstruction and tendon interposition procedures do not appear to be superior to the simpler trapeziectomy. (1534, 1535, 1541, 1549, 1550, 1555, 1556) and some studies suggest longer recovery and higher complication rates with the more extensive procedures. Thus, the ligament reconstruction with tendon interposition procedure is generally not recommended. One moderate-quality study has concluded that earlier mobilization is superior to delayed mobilization after surgery. (1537)

Surgery is often career ending for patients who perform manual labor or requires cessation of manual tasks. Thus patients should be appropriately counseled as they may decide that the fulfillment from performing physical labor outweighs the discomfort. There are no quality studies of joint fusion. However, joint fusion is generally helpful for patients with significantly symptomatic osteoarthrosis who fail to achieve sufficient relief from other treatments.

Evidence for the Use of Surgery for Hand Osteoarthrosis

There are 6 moderate-quality RCTs incorporated into this analysis. (1534, 1537, 1549, 1550, 1555, 1557)
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prosser 2014</td>
<td>6.5</td>
<td>N = 56 (11 males, 45 females) with osteoarthritis of TMC joint underwent TMC arthroplasty allocated to either rigid orthotic or Semi-rigid orthotic groups. Mean age: 66.9±8.5 years.</td>
<td>Trapeziectomy vs. Trapeziectomy plus Palmaris Longus Tendon Interposition</td>
<td>Both groups performed equally well. There was no significant between-group difference for PRWHE scores (0.47, CI -11.5 to 12.4), including subscales for pain and function, or for any of the secondary outcomes at one year follow-up</td>
<td>&quot;The rigid orthosis and semi-rigid orthosis (allowing more wrist and thumb motion) used from 2 to 6 weeks following TMC arthroplasty performed equally well in this study. There was no significant difference between the two groups at one year for the primary outcome of PRWHE scores or for any secondary outcome. Clinically, either orthosis could be recommended. Patient comfort, cost and availability may determine choice between orthoses in clinical practice.&quot;</td>
<td>Data suggest comparable efficacy between rigid vs semi-rigid orthotics post TMC arthroplasty.</td>
</tr>
<tr>
<td>Davis 2004</td>
<td>6.5</td>
<td>N = 162 patients (0 males, 162 females) with painful trapeziometacarpal osteoarthritis; 183 thumbs; 183 surgeries</td>
<td>Simple trapeziectomy vs. trapeziectomy with Palmaris longus interposition vs. trapeziectomy with ligament reconstruction and tendon interposition using 50% of flexor carpi radialis tendon. All thumbs splinted for 6 weeks.</td>
<td>82% good pain relief and 68% regained sufficient strength for normal activities of daily living at 1-year follow-up. No differences in pain levels at 3 months (p = 0.58) or 1 year (p = 0.4). Pain levels at 3 months (No pain or restriction): T = 12, T+PL = 9, T+LRTI = 10. Discomfort with use but no restriction: T = 24, T+PL = 20, T+LRTI = 19.</td>
<td>&quot;The outcomes of these 3 variations of trapeziectomy were very similar at 1-year evaluation. In the short term at least there appears to be no benefit to tendon interposition or ligament reconstruction.&quot;</td>
<td>Includes patients in other report below; 21 bilateral cases – did not always crossover. Results suggest no differences in outcomes.</td>
</tr>
</tbody>
</table>
| Davis 2009 | 6.5          | N = 113 patients (5 males, 103 females) (133 thumbs; 20 bilateral) | Trapeziectomy with Flexor carpi radialis ligament reconstruction, tendon interposition and Kirschner wire insertion followed by splintage for 6 weeks vs. excision of trapezium with no Kirschner wire and immobilization of thumb in soft bandage for 3 weeks. | At 1 year, 81% of trapeziectomy had no pain or only discomfort after use with no activity restrictions vs. 67% of trapeziectomy with LRTI (p = 0.1). DASH scores [baseline (95% CI)/3 months/1 year]: Trapeziectomy [65(58-72)/52(44-59)/34 (26-42)] vs. Trapeziectomy and LRTI [65(59-72)/42 (35-"

"[T]his study found that the results of simple excision of the trapezium, as described by Gervis (1949), are similar to those produced by excision of the trapezium with ligament reconstruction and tendon interposition."

Suggests no short or intermediate term (1 year) benefits demonstrable of more extensive procedures and trend of benefit for trapeziectomy alone. |
<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Patients Characteristics</th>
<th>Procedure Details</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Davis 1997 RCT</td>
<td>4.5</td>
<td>N = 76 patients (0 males, 76 females)</td>
<td>Trapeziectomy vs. trapeziectomy with soft tissue interposition and tendon interposition</td>
<td>RSD complications: T = 0, T+STI = 0, T+LRTI = 2. Thumb key pinch strengths (baseline/3 months/1 year): T (3.7/3.4/4.8) vs. T+STI (4.0/3.1/4.6) vs. T+LRTI (3.4/3.1/4.4). Hand grip strengths [mean (range in kg)]: T [14.8 (4-46)/14.7(2-40)/19.2] vs. T+STI [12.4 (4-25)/10.8 (2-27)/16.9] vs. T+LRTI [11.3 (1-22)/14.0 (2-25)/19.1].</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>&quot;In the short term at least, tendon interposition and ligament reconstruction do not improve the results of trapeziectomy.&quot;</td>
</tr>
<tr>
<td>Kriegs-Au 2004 RCT</td>
<td>4.0</td>
<td>N = 43 patients (6 males, 25 females) (52 thumbs)</td>
<td>Trapezial excision with ligament re-construction vs. trapezial excision with tendon interposition</td>
<td>Long-term outcome (Buck-Gramcko Score): 51.3 vs. 44.6 points. Strength measures Group I (ligament reconstruction) vs. (pre-op and final follow-up) vs. Group II (ligament reconstruction and tendon interposition): Mean tip-pinch strength (bar[Pa]): 0.21, 0.32; 0.23, 0.25; Mean grip strength bar [Pa]): 0.52, 0.46; 0.52, 0.44; Mean palmar abduction (degree): 10.7, 3.6;2.4; 11.9, 4.1;2.9.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>&quot;Tendon interposition does not affect the outcome after the ligament reconstruction for the treatment of osteoarthritis of the thumb carpometacarpal joint. Furthermore, proximal migration of the thumb metacarpal does not appear to influence the functional outcome.&quot;</td>
</tr>
</tbody>
</table>

Some baseline differences. Results suggest trapeziectomy equivalent to combined ligament reconstruction procedure or soft tissue interposition.
Post-operative Rehabilitation and Rehabilitation of Patients with Functional Deficits: CTS and Other Disorders

Post-operative rehabilitation and rehabilitation for patients with distal upper extremity musculoskeletal disorders has long been prescribed. Post-operative splinting was previously widely used as evidenced in the older quality literature. (911, 915, 937, 1558, 1559) But, plaster casts have been replaced by splints which were later replaced by soft bandages and dressings (907, 918, 922-924, 928, 932, 935, 953, 958) which has also coincided with, or been facilitated by, less invasive and smaller incisions.

SOFT BANDAGES AND SPLINTS

1. Recommendation: Soft Bandages During Post-operative Rehabilitation
   Soft bandages are recommended during post-operative rehabilitation.
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Low

2. Recommendation: Splints During Post-operative Rehabilitation
   Splints are recommended during post-operative rehabilitation for select patients.
   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Low

Rationale for Recommendations
As surgery has become less invasive, the degree or whether to splint, has become questionable as splints encourage lack of mobility which likely impairs or delays recovery with potentially increasing risk of complex regional pain syndrome, debility and delayed recovery. Three low-quality studies all suggest that splints are not effective; (784, 1559, 1560) however, there is no quality data and some splints appear indicated for select patients. Thus, there are limited indications for splints in patients with select diagnoses generally involving more extensive surgical procedures or other needs to utilized splints for protective purposes.

Evidence for the Use of Post-operative Soft Bandages and Splints
There are 7 moderate-quality RCTs (1459, 1561-1566) incorporated into this analysis. There are 4 low-quality RCTs in Appendix 2 (963, 1559, 1560, 1567)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Soft bandage, splint, splinting, immobilization, Postoperative Period, post-operative, rehabilitation, upper, extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 120 articles in PubMed, 12 in Scopus, 35 in CINAHL, 1 in Cochrane Library and 18800 in Google Scholar. We considered for inclusion 7 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library 0 from Google Scholar, and 1 from other sources. Of the 18968 articles considered for inclusion, 11 randomized trials and 1 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crowley 2013</td>
<td>RCT</td>
<td>No sponsorship or COI.</td>
<td>4.0</td>
<td>N = 12 with ulnar collateral ligament (UCL) injuries of the thumb who underwent UCL repair with Mitek bone anchor. Median age 42 years.</td>
<td>SR: standard rehabilitation for 4 weeks of immobilization in POP thumb spica then 2 weeks of flexion, extension, opposition, abduction, and adduction of thumb and ultrasound, scar massage, light function for ADLs, and splint at night and out of home; therapy continued for 2-4 more weeks (N = 6) vs. (EAM) early controlled active mobilization 3-5 days postop; given custom-made thermoplastic splint; first 4 weeks exercises emphasized flexion, extension, opposition, abduction and adduction of thumb ; next 2 weeks same are SR group (N = 6). Study duration, 8-12 weeks or until participants could resume full ADLs.</td>
<td>There were no significant differences between groups.</td>
<td>&quot;Our results suggest that there may be a benefit in early active mobilization over standard rehabilitation but that a larger randomized control trial is needed to assess this more accurately.&quot;</td>
<td>Pilot study of 12 patients. Data suggest early active mobilization lead to earlier restoration of hand function as well as an earlier return to work but no difference between groups in final ROM. A larger study would support preliminary findings.</td>
</tr>
<tr>
<td>Germann 2001</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>4.0</td>
<td>N = 20 with extensor indicis proprius transfer for extensor pollicis longer (EPL) tendon rupture. Mean age dynamic motion 52 years, immobilization 42 years.</td>
<td>Dynamic motion protocol (DG group): 2 days after surgery were forearm splint with limited but progressive increase in active flexion of interphalangeal (IP) joint plus passive extension through wire-rubber band system for 3 weeks (N = 10) vs. immobilization protocol (IG group): forearm cast with 20º wrist extension</td>
<td>Active ROM of IP joint after 4 weeks: DG 74º vs. IG 50º (p&lt;0.05). Grip strength (DG vs. IG): 3 weeks 49% vs. 27% (p&lt;0.05); 4 weeks 45% vs. 60% (p&lt;0.05); 6 weeks 44% vs. 65% (p&lt;0.05). Pinch grip (DG vs. IG): 3 weeks 36% vs. 20% (p&lt;0.05).</td>
<td>&quot;The dynamic protocol can therefore be considered as an important factor for a considerable reduction of overall treatment cost. Although all parameters plateaued after 6 and 8 weeks, the early dynamic motion protocol is the superior concept and has become standard procedure for these patients.&quot;</td>
<td>Small sample. Data suggest early dynamic motion group had better ROM of the interphalangeal joint grip and pinch strength at 3 weeks compared to immobilization group. Hand function was comparable between groups at 6 and 8 weeks but the shortened total rehab...</td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Splint Type</td>
<td>Control Type</td>
<td>Duration</td>
<td>Sample Size</td>
<td>Outcome Measures</td>
<td>Results</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>-------------</td>
<td>--------------</td>
<td>----------</td>
<td>-------------</td>
<td>-----------------</td>
<td>---------</td>
<td></td>
</tr>
<tr>
<td>Sillem 2011</td>
<td>RCT/Crossover</td>
<td>Comfort Cool™ prefabricated neoprene splint (N = 59) vs. Hybrid custom-made splint (N = 59)</td>
<td>Time in dynamic motion</td>
<td>6.0</td>
<td>N = 59 with carpometacarpal (CMC) OA of the thumb. Mean age 64 years.</td>
<td>Follow-up at 3, 4, 6, and 8 weeks after surgery</td>
<td>Mean±SD mean difference Australian Canadian Hand Osteoarthritis Hand Index (AUSCAN): 3.7±11.13 in favor of Hybrid splint (p=0.02). “The Hybrid and Comfort Cool™ splints had an equivalent therapeutic effect on hand function, grip strength, and lateral pinch strength.” Crossover equivalence trial. Data showed comparable results for hand function, grip strength and lateral pinch strength but the Hybrid splint was better at decreasing pain compared to Comfort Cool™.</td>
<td></td>
</tr>
<tr>
<td>Sillem 2011</td>
<td>RCT/Control</td>
<td>Comfort Cool™ prefabricated neoprene splint (n = 59) vs. Control group: usual care (n = 55).</td>
<td></td>
<td>7.5</td>
<td>N = 112 (101 female/11 male) with base-of-thumb osteoarthritis. Mean age splint group 63.0±7.9 years, control 63.5±7.6 years.</td>
<td>Follow-up: 1, 6, and 12 months.</td>
<td>Intervention group had intervention in VAS pain score/reduction in disability by Cochin Hand Function Scale score/patient-perceived disability at 12 months: -22.2 vs. -7.9; -14.3 [CI: -23.4 to -5.2]; p = 0.002/ -1.9 vs. 4.3; -6.3 [CI: -10.9 to -1.7]; p = 0.008/-11.6 vs. 1.5; -13.1 [CI: -21.8 to -4.4]; (p = 0.003). Intervention group experienced statistically significant improvements (61% vs. 38%, &gt;10-mm [p = 0.014]; 56% vs.31% &gt;15-mm [p = 0.007]; and 54% vs. 25% &gt;20-mm [p = 0.002]). For patients with base-of-thumb osteoarthritis, wearing a splint had no effect on pain at 1 month but improved pain and disability at 12 months.” Crossover equivalence trial. Data suggest wearing a splint for base of thumb OA had no effect on pain reduction at one month but at 12 months there was pain and function improvement.</td>
<td></td>
</tr>
<tr>
<td>Rannou 2009</td>
<td>RCT</td>
<td>Intervention group: custom-made neoprene splint (n = 57) vs. Control group: usual care (n = 55).</td>
<td></td>
<td>7.5</td>
<td>N = 112 (101 female/11 male) with base-of-thumb osteoarthritis. Mean age splint group 63.0±7.9 years, control 63.5±7.6 years.</td>
<td>Follow-up: 1, 6, and 12 months.</td>
<td>There were no significant differences between groups.</td>
<td></td>
</tr>
<tr>
<td>Hermann 2014</td>
<td>RCT</td>
<td>Orthosis group: soft thumb base orthosis and hand exercises focused on increasing joint mobility, grip strength, and stability of CMC joint 2 sessions per day (n = 30) vs. Control group: usual care (n = 30).</td>
<td></td>
<td>7.5</td>
<td>N = 59 (58 female/1 male) with hand osteoarthritis (HOA). Mean age 70.5±6.7 years.</td>
<td>There were no significant differences between groups.</td>
<td>“[A] soft orthosis seems to have an immediate pain-relieving effect when worn, but no general effect in terms of reduced pain, or improved hand strength. Data suggest a soft orthosis has immediate pain relieving benefits when worn but no benefit in terms of pain reduction.”</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Sponsorship</td>
<td>COI</td>
<td>Conclusion</td>
<td>Notes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>--------</td>
<td>-------------</td>
<td>-----</td>
<td>------------</td>
<td>-------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jerosch-Herold 2011</td>
<td>RCT</td>
<td>Sponsored by Action Medical Research Charity and National Institute for Health Research (NIHR). No COI.</td>
<td>-</td>
<td>“Contrary to the widespread belief in the value of postop night splinting for up to 6 months after fasciectomy or dermofasciectomy we found no evidence of its short or long-term effect.”</td>
<td>Data suggest comparable results from self-reported outcomes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cook 1995</td>
<td>RCT</td>
<td>No mention of sponsorship of COI.</td>
<td>-</td>
<td>“We conclude that splinting the wrist following open release of the flexor retinaculum is largely detrimental, although it may have a role in preventing the rare but significant complications of bowstringing of the tendons or entrapment of the median nerve in scar tissue. We recommend a home physiotherapy programme in which the wrist and fingers are exercised separately to avoid simultaneous finger and wrist flexion, which is the position most prone to cause bowstringing.”</td>
<td>Sparse details. Full open incision suggests splints not appropriate post-operatively.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Therapy Association, Norwegian Rheumatism Association, and the Norwegian Women’s Public Health Association. No COI.

Control group: hand exercises only (n = 29). Study duration 2 months. Follow-up at 2 months. or activity performance in participants with CMC-OA when not worn.

Control group: hand exercises only (n = 29). Study duration 2 months. Follow-up at 2 months.

Hand therapy only (n = 77) vs. hand therapy with night splinting worn for 6 months (n = 77). Follow-up for 12 months after surgery.

There were no significant differences between groups.

N = 154 undergoing fasciectomy of dermofasciectomy for Dupuytren’s disease. Mean age hand therapy only 67.5±9.2 years, splint 67.2±10.0 years.

N = 50 patients having undergone CTR. Patient’s age and gender are not disclosed.

Volar splint vs. soft bulky dressing removed 1st post-op day. 1 month follow-up.

Excellent results (14 days/1 month): unsplinted 9/25 (36%)/12/25 (48%) vs. splinted 1/25 (4%)/2/25 (8%). More rapid RTW in unsplinted (15 days vs. 24 days, p = 0.01). Return to full work in 17v27days, p = 0.005.
NSAIDS/ACETAMINOPHEN

1. Recommendation: NSAIDs During Post-operative Rehabilitation

NSAIDs are moderately recommended to control pain during post-operative rehabilitation.

Indications – All hand, wrist, forearm post-operative patients may be candidates other than those with contraindications for use.

Frequency/Duration – Scheduled dosage rather than as needed is generally preferable initially.

Indications for Discontinuation – Resolution of pain, lack of efficacy, development of adverse effects, particularly gastrointestinal.

Strength of Evidence – Moderately Recommended, Evidence (B)
Level of Confidence – High

2. Recommendation: Acetaminophen During Post-operative Rehabilitation

Acetaminophen is recommended to control pain during post-operative rehabilitation.

Indications – All hand, wrist, forearm post-operative patients may be candidates other than those with contraindications for use.

Frequency/Duration – Scheduled dosage rather than as needed is generally preferable initially.

Indications for Discontinuation – Resolution of pain, lack of efficacy, development of adverse effects, particularly gastrointestinal.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Moderate

Rationale for Recommendations

Acetaminophen has been shown to be less efficacious than naproxen, but is recommended due to its lower adverse effects.(639)

Evidence for the Use of NSAIDs Post-operatively

There are 1 high-(639) and 9 moderate-quality(972, 1568-1575) RCTs incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: NSAIDs, Anti-Inflammatory Agents, Non-Steroidal, acetaminophen, Agents, Non-Steroidal, Postoperative, Period, post-operative, rehabilitation, upper, extremity;controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 40 articles in PubMed, 0 in Scopus, 0 in CINAHL, 0 in Cochrane Library and 13502 in Google Scholar. We considered for inclusion 10 from PubMed, 0 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 0 from other sources. Of the 13542 articles considered for inclusion, 10 randomized trials and 0 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Husby 2001</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>8.0</td>
<td>N = 42 (9 female/33 male) due to be operated on for DC or CTS. Mean age 61 years.</td>
<td>Post-op naproxen (500mg BID) vs. Paracetamol (1000mg QID) vs. Placebo for 3 days immediate post-op CT release surgery. Second trial 35 with Dupuytren’s contracture.</td>
<td>Post-op CTS swelling as percentage of pre-op volume 3.5±3.3 vs. 4.8±3.2 vs. 3.8±2.6. For Dupuytren’s contracture releases 5.6±3.8 vs. 6.9±3.7 vs. 8.2±5.1. Additional analgesics used 0, 2, and 8 in naproxen, paracetamol and placebo groups.</td>
<td>“Naproxen might have a clinical relevant effect on swelling when used on minor surgery in the hand, unlike paracetamol. Naproxen might be a useful analgesic during the immediate postoperative phase.”</td>
<td>Results suggest a beneficial effect that the studies were not powered to detect.</td>
</tr>
<tr>
<td>Sen 2006</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>7.5</td>
<td>N = 45 (24 female/21 male) ASA I-II undergoing hand or forearm surgery. Mean age control 45 years, L-IVRA 42 years, L-IV 39 years.</td>
<td>Control group: IV saline 0.9% 2 ml + intravenous regional anesthesia (IVRA) with lidocaine 0.5% and saline (n = 15) vs. L-IVRA group: IV saline + IVRA lidocaine 0.5% with lornoxicam 8mg (n = 15) vs. L-IV group: intravenous lornoxicam 8mg + IVRA lidocaine 0.5% and saline (n = 15). Follow-up for 24 hours post-op.</td>
<td>Mean±SD intraoperative fentanyl (control vs. L-IVRA vs. L-IV): amount (µg) 23.3±25.8 vs. 3.3±12.9 vs. 19.4±18.6 (p = 0.014); requirement time (min): 15.8±6 vs. 28±9 vs. 13.6±8 (p = 0.042). Mean VAS (control vs. L-IVRA vs. L-IV): tourniquet release 3.3±3 vs. 1.7±3 vs. 3.13 (p = 0.003); tourniquet release after 2 hour 2.6 ± 2 vs. 2.0 ± 2 vs. 2.93 (p = 0.031). Mean±SD time to first postoperative analgesic request, minutes (control vs. L-IVRA vs. L-IV): 28±20 vs. 229±85 vs. 95±24 (p = 0.0038). Mean±SD diclofenac mg (control vs. L-IVRA vs. L-IV): 85±26 vs. 15±31 vs. 67±36 (p &lt;0.001). Mean±SD paracetamol consumption mg (control vs. L-IVRA vs. L-IV): 1400±207 vs. 200±253 vs. 1100±320 (p &lt;0.0001).</td>
<td>“[A]ddition of lornoxicam to lidocaine in IVRA shortens sensory and motor block onset times, prolongs sensory and motor block recovery times, and improves tourniquet pain while it prolongs first analgesic requirement time, and decreases total amount of analgesic.”</td>
<td>Pilot study. Data suggest adding lornoxicam to lidocaine for intravenous regional anesthesia shortens the onset of sensory and motor block, decreases tourniquet pain and improves post-op analgesia. However, data suggest recovery times were prolonged in lornoxicam plus lidocaine group.</td>
</tr>
<tr>
<td>Ashworth 2002</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>7.0</td>
<td>N = 47 (20 female/29 male) scheduled for inpatient elective hand surgery. Mean age systemic presurgery 57 years, regional presurgery 54.7 years, systemic</td>
<td>Systemic presurgery group: ketorolac 20mg intravenously in non-operative arm before surgery (n = 15) vs. regional presurgery group: ketorolac 20mg intravenously to operative arm after tourniquet inflation (n = 15) vs. systemic postsurgery group: ketorolac 20mg intravenously in non-</td>
<td>VAS score 24 hours after surgery: 12.2 mm higher in systemic postsurgery group vs. systemic presurgery group (p=0.037).</td>
<td>“[T]here seems no benefit to be gained by giving ketorolac as intravenous regional anaesthesia compared with the usual method of giving it intravenously into the general circulation before the operation.”</td>
<td>Data suggest no benefit in the administration of ketorolac post surgery.</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Age (mean ±SD)</td>
<td>Anesthesia</td>
<td>Pain Management</td>
<td>Results</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>---------------------</td>
<td>---</td>
<td>----------------</td>
<td>------------</td>
<td>----------------</td>
<td>---------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rivera 2008</td>
<td>7.0</td>
<td>N = 60 (20 female/35 male) undergoing hand surgery. Mean age ketorolac 39.5±13.6 years, placebo 37.58±12.2 years.</td>
<td>Bier block of 50mL of 0.5% lidocaine + 20mg ketorolac (n = 30) vs. Bier block of 50mL of 0.5% lidocaine + normal saline (n = 30). Follow-up 48 hours after discharge.</td>
<td>VAS post anesthesia care unit (PACU) ketorolac vs. control: 30 min 0.48 vs. 2.20 (p&lt;0.05); 45 min 0.38 vs. 2.23 (p&lt;0.05); 60 min 0.45 vs. 2.50 (p&lt;0.05). Median time (minutes) to second request of postop analgesic (ketorolac vs. placebo): 1102 vs. 505 (p=0.048).</td>
<td>“Based on the results of this study we recommend that 20 mg ketorolac be considered in intravenous regional anesthesia.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sai 2001</td>
<td>6.5</td>
<td>N = 120 (gender not specified) undergoing hand surgery with brachial plexus block. Mean age 43 years.</td>
<td>Amproficamic 27mg orally vs. alegioxa 100mg, orally vs. placebo 3 hours before surgery. Follow-up when each patient requested an analgesic suppository.</td>
<td>Median pain scores at time of first analgesic request (analgesic vs placebo): 1.0 vs. 4.0 (p&lt;0.0001). Median pain scores at 24 hours after operation (analgesic vs. placebo): 0 vs. 2.0 (p&lt;0.0001). Number of patients requiring analgesic suppositories (analgesic vs placebo): 6 vs 44 (p&lt;0.0001).</td>
<td>“We suggest that preoperative administration of amproficamic improves pain control during the early post-operative phase.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cornesse 2010</td>
<td>6.0</td>
<td>N = 60 undergoing minor hand surgery (carpal tunnel release or synovial cyst resection) under intravenous regional anesthesia. Mean age 1 g 51±15 years, 2 g 55±18 years.</td>
<td>1 g intravenous paracetamol before surgery (n = 30) vs. 2 g intravenous paracetamol before surgery (n = 30). Discharged after 4 hours. Once at home, patients instructed to take 1 g of paracetamol orally every 6 hours. Follow-up for 24 hours after surgery.</td>
<td>Pain scores: lower in 2 g paracetamol intravenous group vs. 1 g paracetamol intravenous (p=0.04).</td>
<td>“[A]n intravenous loading dose of 2 g paracetamol provides better analgesia than 1 g in adult patients undergoing minor hand surgery.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rawal 2001</td>
<td>6.0</td>
<td>N = 120 ASA I-II undergoing ambulatory hand surgery with IV regional anesthesia.</td>
<td>Group T: tramadol 100 mg orally every 6 hours (n = 40) vs. Group M: metamizol 1 g every 6 hours (n = 40) vs. Group P: paracetamol 1 g every 6 hours (n = 40)</td>
<td>Mean ±SD number of study tablets (tramadol vs. metamizol vs. paracetamol) day 1/ day 2: 5.5±1.1/ 5.0±2.6 vs. 4.9±1.1/ 6.0±2.9 vs. 2.8±1.2/ 3.1±0.6 (p&lt;0.05 metamizol vs. tramadol on day of surgery;</td>
<td>“None of the study drugs provided adequate analgesia for all patients, as about 40% required rescue analgesia.”</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Data suggest tramadol most effective in pain relief of ambulatory hand surgery patients. It was associated with the greatest number</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Design</td>
<td>Sponsorship</td>
<td>COI</td>
<td>Methodology</td>
<td>Patients</td>
<td>Intervention</td>
<td>Follow-up</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>-----</td>
<td>--------</td>
<td>-------------</td>
<td>-----</td>
<td>-------------</td>
<td>----------</td>
<td>--------------</td>
<td>-----------</td>
</tr>
<tr>
<td>Spagnoli 2011 RCT</td>
<td>6.0</td>
<td>N = 114 with postoperative pain following hand and foot surgery under brachial plexus block. Mean age 56 years.</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td>Group TP: tramadol/paracetamol 37.5/325mg (n = 57) vs. Group P: paracetamol monotherapy 1000 mg (n = 57) 2 tablets a day for 3 days. Follow-up 7 days after discharge.</td>
<td>Mean VAS (paracetamol vs. tramadol/paracetamol): post-op 0-6 hours 1.92 vs. 0.40 (p &lt; 0.005). Number requiring extra dose analgesic post-op (paracetamol vs. tramadol/paracetamol): 0-6 h 32 vs. 4 (p&lt;0.005); 6-12 h 11 vs. 0 (p &lt;0.005); 12-24 h 7 vs. 0 (p &lt;0.01).</td>
<td>“The association of tramadol and paracetamol appears to have more efficacy when compared with paracetamol monotherapy for acute postoperative pain after hand and foot surgery.”</td>
</tr>
<tr>
<td>Jankovic 2008 RCT</td>
<td>5.5</td>
<td>N = 45 ASA physical status I-II undergoing ambulatory hand surgery. Mean age Group L 34±12 years, Group LK 33±12 years, Group LDK 35±13 years.</td>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td>Group L: 3mg/kg 2% lidocaine for intravenous regional anesthesia (IVRA) (n = 15) vs. Group LK: 3mg/kg 2% lidocaine + 30 mg ketorolac for IVRA (n = 15) vs. Group LDK: 3 mg/kg 2% lidocaine + 8 mg dexamethasone + 30 mg ketorolac for IVRA (n =15). All groups received 0.9% NaCl added for total volume of 40mL. All patients allowed 10mg ketorolac every 6 hours as needed at home. Follow-up 24 hours after surgery.</td>
<td>Median postoperative VAS scores post anesthesia care unit admittance (PACU ad.) 120 min (Group L vs. Group KL vs. Group LDK): 3.8±1.3 vs. 2.2±1.6 vs. 1.3±0.6 (p&lt;0.05 Group LDK vs. Group L; p&lt;0.05 Group LDK vs. Group KL).</td>
<td>“The addition of both ketorolac and dexamethasone to lidocaine IVRA provided improved tourniquet tolerance, prolonged analgesia in the postanesthesia care unit during the first 2 h after the medical procedure, and diminished the need for analgesic supplements during the first day after ambulatory hand surgery.”</td>
</tr>
<tr>
<td>Reuben 1995 RCT</td>
<td>5.0</td>
<td>N = 60 undergoing hand surgery (carpal tunnel release).</td>
<td></td>
<td></td>
<td>Control group: 0.9% intravenous (IV) saline 2mL and intravenous regional anesthesia (IVRA) with saline</td>
<td>Median postoperative pain scores (control vs. IV-K vs. IVRA-K): VAS 30: 1.1 vs. 0.9 vs. 0.3 (p&lt;0.0001 IVRA-K vs. other groups); VAS 60: 1.0±1.6±0.7 (p = 0.0131 IVRA-K vs.</td>
<td>“[T]he addition of K to 0.5% lidocaine for IVRA provided better control of intraoperative tourniquet pain, improved analgesia in</td>
<td>Sparse methods. Data suggests addition of ketorolac to 0.5% lidocaine provided better</td>
</tr>
<tr>
<td>No mention of sponsorship or COI.</td>
<td>excision of a ganglion cyst, or tenolysis). Mean age control 49±17 years, IV-K 46±21 years, IVRA-K 50±19 years.</td>
<td>added to it (n = 20) vs. Group IV-K: ketorolac 60 mg IV and saline added to IVRA solution (n = 20) vs. Group IVRA-K: saline IV and ketorolac 60 mg added to IVRA solution (n = 20). All patients allowed Tylenol No. 3 tablets every 4 hours as needed for pain at home. Follow-up 24 hours.</td>
<td>other groups). Mean±SD 24 hour total medicine (control vs. IV-K vs. IVRA-K): 4.6±1.3 vs. 3.0±1.1 vs. 1.9±1.4 (p=0.0003 IVRA-K vs. other groups). Mean±SD time to first medicine (control vs. IV-K vs. IVRA-K): 281±2.44 vs. 356±255 vs. 653±501 (p=0.0241 IVRA-K vs. other groups).</td>
<td>the PACU during the first postoperative hour, and diminished the need for analgesic supplements during the first postoperative day.”</td>
<td>control of intraoperative tourniquet pain, improved PACU pain relief during the first hour post-op and up to 24h post-op compared to either lidocaine alone or placebo.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
ARNICA

**Recommendation: Arnica During Post-operative Rehabilitation**

Arnica is **not recommended during post-operative rehabilitation**.

**Strength of Evidence** – Not Recommended, Evidence (C)

**Level of Confidence** – Low

**Rationale for Recommendation**

Arnica has been utilized for post-operative recovery in CTS patients, (772, 1576) with the two quality studies conflicting. However the higher quality study suggests a lack of efficacy, thus there is overall weak evidence that arnica is ineffective, and thus it is not recommended.

**Evidence for the Use of Arnica Post-Operatively**

There is 1 high-(772) and 1 moderate-quality(1576) RCT incorporated into this analysis.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Arnica, Montana, Postoperative Period, post-operative, rehabilitation, upper, extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 2 articles in PubMed, 9 in Scopus, 19 in CINAHL, 6 in Cochrane Library and 144 in Google Scholar. We considered for inclusion 1 from PubMed, 0 from Scopus, 0 from CINAHL, 2 from Cochrane Library 0 from Google Scholar, and 0 from other sources. Of the 180 articles considered for inclusion, 2 randomized trials and 2 systematic studies met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stevinson</td>
<td>RCT</td>
<td>Sponsored by Dr Susil</td>
<td>8.5</td>
<td>N = 62</td>
<td>Arnica 30C (n = 21) vs. Arnica 6C (n = 21) vs. Placebo TID for 7 days pre-op and 14 days post-op (n = 22).</td>
<td>No pain differences (p = 0.79) and bruising (p = 0.45) at Day 4. Swelling and analgesic use did not differ. Adverse events reported by 2 patients in arnica 6C group, 3 in placebo, 4 in arnica 30C. Results do not suggest homeopathic arnica has an advantage over placebo in reducing post-op pain, bruising and swelling in patients undergoing elective hand surgery.</td>
<td>“Since the experiences of patients who receive no benefit from Arnica are less likely to be reported, the myth becomes reinforced.”</td>
<td>One surgeon operated. Data suggest no efficacy.</td>
</tr>
<tr>
<td>Jeffrey</td>
<td>RCT</td>
<td>Ian Wiggle and Weleda</td>
<td>6.0</td>
<td>N = 32</td>
<td>Arnica D6 3 tablets TID plus Arnica 5% ointment TID vs. double placebo Follow-up was 2 weeks after surgery</td>
<td>Wrist circumferences and grip strengths both non-significant. Pain reduced in Arnica compared with placebo at 2 weeks (p &lt;0.03).</td>
<td>“The role of homeopathic and herbal agents for recovery after surgery merits further investigation.”</td>
<td>Baseline data not given and 1 week data suggest trend. Possible inadequate randomization. Objective measures showed no differences.</td>
</tr>
</tbody>
</table>

Stevinson 2003

RCT

Sponsored by Dr Susil Kumar and Jamila Mitra Charitable Trust (UK); homeopathic and placebo tablets supplied by A Nelson & Co Ltd. No mention of COI.

Jeffrey 2002

RCT

Ian Wiggle and Weleda Ltd provided arnica and placebo preparations. No mention of COI.
CRYOTHERAPY/COOLING BLANKET

1. Recommendation: Cryotherapy During Post-operative Rehabilitation

Cryotherapy is recommended for post-operative rehabilitation for carpal tunnel release patients.

Strength of Evidence – Recommended, Evidence (C)
Level of Confidence – Low

2. Recommendation: Cooling Blanket During Post-operative Rehabilitation

A cooling blanket is recommended during post-operative rehabilitation.

Strength of Evidence – Recommended, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendations
Cryotherapy has been shown to be effective for post-carpal tunnel release patients and is therefore recommended during post-operative rehabilitation. The evidence is in favor of a cooling blanket versus ice therapy and therefore, a cooling blanket is recommended during post-operative rehabilitation.(1577)

Evidence for the Use of Cryotherapy/ Cooling Blanket During Post-operative Rehabilitation
There is 1 moderate-quality RCT incorporated into this analysis.(1577)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: Cryotherapy OR Cooling Blanket / Post-operative rehabilitation and rehabilitation of patients with functional deficits: CTS and other disorders; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found, reviewed and considered for inclusion 17 articles in PubMed, 0 in Scopus, 2 in CINAHL, 0 in Cochrane Library, 3883 in Google Scholar, and 0 in other sources. One RCT met the inclusion criteria.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hochberg</td>
<td>RCT</td>
<td>No sponsorship or COI.</td>
<td>4.0</td>
<td>N = 72 (46 males/26 females) adults presenting Carpal Tunnel Syndrome who were eligible for single open surgical procedures; No specification on mean age of study sample.</td>
<td>Temperature-controlled cooling blanket vs. standard ice pack for 3 days. Follow-Up immediate following post-op and after three days.</td>
<td>Pain ratings (baseline/day 3): cooling blanket (8.3±1.8/4.5±2.3) vs. ice (8.3±1.3/7.3±2.5), p &lt;0.001.</td>
<td>Use of a “(temperature-controlled cooling blanket) compared with traditional ice therapy, provides patients with greater comfort and lessens the need for narcotics.”</td>
<td>Incisional length of 6cm large compared with most recent trials which may have affected results and limits study generalizability to treatment of larger open CTR incisions.</td>
</tr>
</tbody>
</table>
ACTIVITY/EXERCISE/PHYSICAL OR OCCUPATIONAL THERAPY

1. Recommendation: Activity During Post-operative Rehabilitation for Patients with Functional Deficits
   It is recommended that post-operative patients or those with functional deficits stay as active as possible and use the hand as much as possible post-operatively or post-injury.

   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – High

2. Recommendation: Exercise During Post-operative Rehabilitation for Patients with Functional Deficits
   It is recommended that post-operative patients or those with functional deficits perform graded, increased exercises post-operatively or post-injury. A home exercise program may accomplish this for many patients.

   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – High

3. Recommendation: Formal Physical or Occupational Therapy During Post-operative Rehabilitation for Patients with Functional Deficits
   Post-operative patients should be observed particularly for failure to progress as expected, as well as for complex regional pain syndrome (see Chronic Pain guideline) or other complications, and it is recommended that there should be a low threshold for institution of formal physical or occupational therapy for rehabilitation. Patients with functional deficits should have a home exercise program with low threshold to refer to therapy for formal treatment if deficits are considerable or there is a failure to progress as expected with a home exercise program.

   Indications – Failure to progress, or moderate to severe functional deficits.
   Frequency – Five to 6 appointments with objective measures followed. Patients demonstrating continued improvements in objective measures may require additional 5 to 6 appointments.
   Discontinuation – Achievement of goals, failure to progress, adverse effects, non-compliance.

   Strength of Evidence – Recommended, Insufficient Evidence (I)
   Level of Confidence – Moderate

Rationale for Recommendations
Most of the quality studies that have described post-operative rehabilitation components have not prescribed formal physical or occupational therapy for rehabilitation.(907, 918, 923, 1558) Instead, most instructed patients to “keep fingers moving” or perform finger exercises,(922, 928) perform mobility exercises,(946) use the hand daily as tolerated,(922) use “as comfort allowed,”(917) or “use as much as possible” or “as soon as possible.”(778, 940, 953) Unfortunately, there is little quality evidence comparing approaches and there is likely a significant difference between using the hand “as tolerated” and “as much as possible.” There also are no quality studies comparing specific exercises for rehabilitation of patients with deficits compared with no treatment or home exercise programs. Quality studies are needed to address these issues, particularly as they may impact the sizable lost-time problems. In the absence of quality evidence, but inferring from numerous other MSD diagnoses that suggest activity is helpful, it is recommended that patients stay as active as possible and use the hand as much as possible post-operatively, as well as for those with functional deficits, and that there should be a low threshold for institution of formal physical or occupational therapy for rehabilitation.

Evidence for Mobilization During Post-operative Rehabilitation
There are 12 moderate-quality RCTs(958, 1578-1588) incorporated into this analysis. There are 4 low-quality RCTs(1589-1592) in Appendix 2.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: exercise, physical therapy, occupational therapy, upper extremity, postoperative period, postoperative, post-operative, rehabilitation, upper extremity; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic,
systematic review, retrospective, and prospective studies. We found and reviewed 1,005 articles in PubMed, 6,515 in Scopus, 53 in CINAHL, 499 in Cochrane Library, 50,100 in Google Scholar, and 0 from other sources. We considered for inclusion 5 from PubMed, 1 from Scopus, 0 from CINAHL, 0 from Cochrane Library, 0 from Google Scholar, and 13 from other sources. Of the 119 articles considered for inclusion, 17 randomized trials and 2 systematic studies met the inclusion criteria.
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rath 2009</td>
<td>5.0</td>
<td>N = 50 (11 female/39 male) with supple claw hand deformities, ulnar nerve paralysis for &gt;1 year and completion of multi-drug therapy for Hansen’s disease undergoing tendon transfer. Mean age IAMP 31±10 years, immobilization group 28±10</td>
<td>Immediate active motion protocol (IAMP) 2 days after tendon transfer for 3 weeks (n = 25) vs. immobilization after tendon transfer for 3 weeks with therapy beginning 22 days after surgery (n = 25). Follow-up monthly for 3 months after discharge and then at 3 month intervals for 1 year, then once a year. Mean±SD PIP joint angles in open hand position: total digits at discharge IAMP 1±9º vs. immobilization 5±9º (p = 0.005). Mean±SD PIP joint angles in the intrinsic plus position: total digits at discharge IAMP 10±10º vs. immobilization 16±10º (p = 0.00). Mean±SD zero pain level (VAS scores) achieved, week: IAMP 3±1 vs. immobilization 6±1 (p &lt;0.001).</td>
<td>“The current study demonstrates that an early motion protocol results in quicker restoration of function.”</td>
<td>Data suggest IAMP group yields earlier pain relief and quicker restoration of function.</td>
<td></td>
</tr>
<tr>
<td>Giessler 2008</td>
<td>4.0</td>
<td>N = 21 (10 female/11 male) with a closed extensor pollicis longus (EPL) tendon rupture in zones T4 and T5 treated with tendon transfer. Mean age DY 51 years, AC 59 years.</td>
<td>Dynamic extension splinting (DY) starting 2 days postoperative with limited ROM of IP joints vs. early active (AC) protocol starting 2 days postoperative: early active thumb extension with limited flexion in a splint. Both groups wore a dynamic extension splint between exercises and saw hand therapist at least 3 times a week. Splints completely removed after 3 weeks. Follow-up 3, 4, 6, 8 weeks post-op. Total ROM in IP joint at 3 weeks (splint removal): higher in DY group vs. AC group (p=0.027). Relative ROM of CMP and IP joints vs. contralateral thumb week 3: active ROM of IP joints DY group 72% of contralateral side vs. AC group 49% of contralateral side (p=0.005).</td>
<td>“Considering the small group sizes, both regimens (dynamic vs early active) achieved comparable clinical results. The early active protocol does not have a notably higher complication rate but fails to accelerate rehabilitation.”</td>
<td>Small sample (N = 21). Data suggest comparable efficacy between groups although the early active protocol reported a higher complication rate without rehabilitation rate acceleration.</td>
<td></td>
</tr>
<tr>
<td>Souer 2011</td>
<td>5.0</td>
<td>N = 94 with unstable distal</td>
<td>Surgeon-directed independent exercises: wrist splint until full 3 month outcomes (independent exercise vs.</td>
<td>“[T]his clinical trial supports the concept</td>
<td>Data suggest formal, prescribed PT is not as</td>
<td></td>
</tr>
</tbody>
</table>
| RCT | Radial fracture treated with open reduction and volar locking plate fixation and screws alone within 4 weeks of injury.  
No mention of gender distribution.  
Mean age occupational therapy 50.7 years, independent exercise 48.6 years. | Finger and forearm motion and then wean out wearing splint to regain wrist motion; performed exercises for finger flexion, forearm supination and pronation 3-4 times a day for at least 30 minutes (n = 48) vs. occupational therapy: supervised exercises to regain digit, wrist, and forearm motion and strengthen hand (n = 46). Follow-up at 6 weeks, 3 months and 6 months after surgery. | Occupational therapy: grip strength (lb) 55±22.6 vs. 45±17.4 (p <0.05); grip strength (% of value on uninjured side) 81±18.9 vs. 62±16.0 (p <0.05); pinch strength (% of value on uninjured side) 90±23.7 vs. 80±22.7 (p <0.05); Garlant and Werley score (points) 2±1.3 vs. 2±2.2 (p <0.05). 6 month outcomes (independent exercise vs. occupational therapy): wrist flexion-extension arc (deg) 129±22.6 vs. 118±17.7 (p <0.05); Wrist flexion-extension arc (% of value on uninjured side) 88±11.7 vs. 84±7.3 (p <0.05); Wrist extension (deg) 62±13.7 vs. 55±10.2 (p <0.05); Ulnar deviation (deg) 40±9.2 vs. 32±12.1 (p <0.05); Ulnar deviation (% of value on uninjured side) 93±19.4 vs. 82±29.2 (p <0.05); Supination (deg) 90±0.9 vs. 84±13.1 (p <0.05); Grip strength (% of value on uninjured side) 92±19.8 vs. 81±16.4 (p <0.05); Mayo wrist score (points) 83.4±12.7 vs. 79.0±9.9 (p <0.05). | That patient education and independent exercises are, on the average, adequate for optimal recovery from a distal radial fracture treated with open reduction and volar plate fixation. |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Krischak 2009 RCT</td>
<td>N = 46 (30 female/16 male) with distal radius fractures undergoing internal fixation with locking plates after open reduction.</td>
<td>Physical therapy, 12 sessions lasting 20-30 minutes each, 6 weeks (n = 23) vs. unassisted home exercise program for 6 weeks, detailed instructions and demonstrations given after surgery (n = 23). All put in splint after surgery for 2 weeks. Splint removed for therapy and then back on afterward. Follow-up 1 week after surgery and after 6 weeks of treatment.</td>
<td>Mean±SD Patient Rated Wrist Evaluation (PRWE) score at 6 weeks: home exercise 18.5±13.9 vs. physical therapy 61.1±13.9 (p &lt;0.001). Mean grip strength relative to opposing healthy side 6 weeks: home exercise 54% of starting base value vs. physical therapy 32% of starting base value (p =0.003). ROM of extension and flexion after 6 months: physical therapy 91.4±19.8 vs. home exercise 87.2±17.8 (p = 0.004).</td>
<td>Data suggest PT after volar plating of wrist fractures is effective for post-op rehab although data in study is self-reported in a training diary.</td>
</tr>
<tr>
<td>Study</td>
<td>Mean age</td>
<td>Weeks of treatment: home exercise 79% of uninjured side vs. physical therapy 52% of uninjured side (p &lt;0.001). Ulnar and radial abduction compared to uninjured side at 6 weeks: home exercise 70% vs. physical therapy 59% (p = 0.013).</td>
<td>Physical Therapy/ Occupational Therapy</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>----------</td>
<td>--------------------------------------------------------------------------------</td>
<td>--------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Pomerance 2007 RCT</td>
<td>Mean age home exercise 53.7±17.9 years, physical therapy 56.0±11.1 years.</td>
<td>Therapy (2 week course, 6 sessions, nerve gliding, ROM, strengthening) (n = 73) vs. No therapy. No restrictions to motion and no splints either group. RTW allowed at first post-op visit (N = 77).</td>
<td>“The current randomized study failed to show benefit in a 2-week course of hand therapy after carpal tunnel release using a short incision. The cost of supervised therapy for an uncomplicated carpal tunnel release seems unjustified.”</td>
<td></td>
</tr>
<tr>
<td>Provinciali 2000 RCT</td>
<td>Mean age 46 years.</td>
<td>Multimodal rehabilitative treatment vs. progressive home exercise program</td>
<td>“A rehabilitation approach after hand surgery is clinically relevant to accelerate recovery but neither modifies functional recovery nor reduces symptom occurrence.”</td>
<td></td>
</tr>
<tr>
<td>Mitsukane 2015 RCT</td>
<td>Mean age 63±13.0 years.</td>
<td>Experimental group: 30 repetitive wrist extensions of injured wrist with maximal isometric contraction for 3 seconds followed by 3 seconds of rest repeated 10 times for 1 minute with a minute rest, sequence repeated 3 times during a 6 minute period (n = 14) vs. control group: no exercises, 6 minutes of rest (n = 14). Follow-up after Mean±SD change in grip strength (kg) post intervention (experiment vs. control): 16.4±9.9 (p=0.01) vs. 15.3±8.2 (p=0.26). Mean±SD change in VAS (mm) post intervention (experiment vs. control): 2.3±5.1 (p=0.03) vs. 13.3±23.0 (p=0.13).</td>
<td>“This study suggests that repetitive maximal wrist extension is useful in physical examinations to reveal the maximal grip force of patients with DRF, and it is effective as a warm-up training procedure in preparation for Small sample, sparse methods. Data suggest grip strength increased in experimental group immediately after repetitive wrist extension but not in control group. Pain decreased in experimental group vs. control group.”</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>N</td>
<td>Intervention</td>
<td>Conventional Grip Strength Exercises</td>
</tr>
<tr>
<td>---------------</td>
<td>------</td>
<td>------------</td>
<td>--------------</td>
<td>--------------------------------------</td>
</tr>
<tr>
<td>Rostami 2013</td>
<td>4.5</td>
<td>N = 23 (17 female/6 male) with active ROM impairment of hand after orthopaedic injuries. Mean age 38 years.</td>
<td>Mirror therapy (MT): concentrating on ROM exercises on unaffected hand in mirror while performing ROM exercises with impaired hand not in mirror 30 minutes a day, 5 days a week for 3 weeks plus half hour of conventional rehab (tendon gliding exercises, blocking exercises, place-and-hold exercise, PNF techniques, dynamic splinting, functional activities, and ADLs) after each MT session (n = 15) vs. control group: conventional rehabilitation for 30 minutes plus 30 minutes direct observation of affected hand performing movements 5 days a week for 3 weeks (n = 15). Both groups performed a 15 minute home program, MT for MT group and active range of motion (AROM) for control group. Assessments at baseline and day after 3 week intervention ended.</td>
<td>Mean±SD change total active motion (TAM) pre to post/post to follow-up (MT vs. control): 154±32 vs. 61±24 (p = 0.001) NS. Mean±SD change DASH score pre to post/post to follow-up (MT vs. control): -34±7 vs. -15±11 (p = 0.001)/ -5±4 vs. -10±6 (p = 0.02).</td>
</tr>
<tr>
<td>Guzelkucuk 2007</td>
<td>4.0</td>
<td>N = 36 with functional loss due to hand injury. Bone, tendon, peripheral nerve injuries, with impaired hand function.</td>
<td>Controls: rehab program (physical therapy, passive, active assist, active ROM, strengthening, BID) vs. therapy plus therapeutic exercises (same exercises plus 1 session of therapeutic activities). Sessions 30 minutes, 5 days a week for 3 weeks. HEP after 3 weeks; 2 month follow-up.</td>
<td>Grip strength (baseline/post/follow-up): Control (10±9/10±9/11±10) vs. therapeutic exercises (7±5/13±6/23±14), p &lt;0.001. Pinch strength, Jebsen tests also all p &lt;0.001.</td>
</tr>
</tbody>
</table>

“Findings suggest that adding a regular and scheduled programme of MT to classic rehabilitation techniques is effective for early and maximum improvement of motor recovery and functional abilities in the patients with orthopaedic injuries.”

Data suggest MT plus conventional rehab was better than control group.

“Because of the complex anatomy, determination of the most appropriate treatment may not be easy in an injured hand. Our results showed that the therapeutic activities that mimick the ADL improve the functions of the hand more effectively.”

Some sparse details. Heterogeneous disorders. Seen 1.5-6 months after injuries. More contact time in exp. group. Trend to longer time since injury in controls. Also suggests benefits of therapy with emphasis on functional exercise.
<table>
<thead>
<tr>
<th>Study</th>
<th>Methodology</th>
<th>n</th>
<th>Description</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnus 2013 RCT</td>
<td></td>
<td>4.0</td>
<td>N = 51 females with unilateral distal radius fracture &lt;2 weeks old. All &gt;50 years of age. Mean age 63.0±10.0 years. Standard rehabilitation: forearm casting; 6 visits to clinic at weeks 1, 3, 6, 9, 12, and 26 post-fracture; and adoption of 3 exercise protocols targeting the fractured side; active ROM of neck, shoulder, elbow, fingers, and thumb while in cast; cast removed – exercises focused on improving active and passive ROM of fractured wrist and hand; stretching and strengthening with encouragement to continue at home after 12 weeks, control (n = 24) vs. standard rehabilitation + strength training of nonfractured arm for 26 weeks, train (n = 27). Follow-up for 26 weeks.</td>
<td>Mean±SD handgrip strength of fractured arm 12 weeks postfracture (training vs. control): 17.3±7.4 kg vs. 11.8±5.8 kg (p = 0.017). Mean handgrip strength of nonfractured arm at 12 weeks postfracture (training vs. control): 30.7±6.5 vs. 24.9±4.4 (p = 0.017). Mean±SD ROM data (degrees) 12 weeks postfracture (training vs. control): flexion/extension 100.5±19.2 vs. 80.2±28.7 (p = 0.017). “This intervention study found that strength training the nonfractured limb was associated with significantly improved strength and ROM in the fractured limb via cross-education in the early stages of rehabilitation.”</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dilek 2013 RCT</td>
<td></td>
<td>6.0</td>
<td>N = 46 (40 female/6 male) with bilateral hand osteoarthritis. Mean age paraffin 58.87±9.47 years, control 59.95±8.71 years. Group 1: dip-wrap paraffin bath therapy at 50ºC 10 dips followed by 15 minutes in a plastic bag until paraffin cooled 5 times a week for 3 weeks for both hands (n = 29) vs. Group 2: control (n = 27). All patients received education about disease and joint protection techniques and allowed paracetamol. Follow-up at 3 and 12 weeks.</td>
<td>Median pain at rest: 3 weeks paraffin group 2.00 vs. control 4.00 (p = 0.01); 12 weeks 0.00 vs. 5.00 (p &lt;0.001). Median grip strength: right hand 12 weeks paraffin group 20.00 vs. control 13.33 (p = 0.004); left hand 12 weeks 18.00 vs. 12.00 (p = 0.010). Median pinch strength: right hand chuck pinch 12 weeks 5.33 “Paraffin bath therapy seems to be effective both in reducing pain and tenderness and maintaining muscle strength in hand osteoarthritis.”</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Cross-Education**

No mention of gender distribution.

Average age of 23±3 years.

**Paraffin Bath Therapy**

No sponsorship or COI.

Median pain at rest: 3 weeks paraffin group 2.00 vs. control 4.00 (p = 0.01); 12 weeks 0.00 vs. 5.00 (p <0.001). Median grip strength: right hand 12 weeks paraffin group 20.00 vs. control 13.33 (p = 0.004); left hand 12 weeks 18.00 vs. 12.00 (p = 0.010). Median pinch strength: right hand chuck pinch 12 weeks 5.33 “Paraffin bath therapy seems to be effective both in reducing pain and tenderness and maintaining muscle strength in hand osteoarthritis.”

Data suggest paraffin bath therapy had significant benefit in hand OA both for pain reduction and muscle strength retention suggesting paraffin may be a short term therapy option.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>Setting</th>
<th>Participants</th>
<th>Interventions</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Massage Therapy</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Field 2011</td>
<td>RCT</td>
<td>Johnson &amp; Johnson Pediatric Institute and Massage Envy</td>
<td>N = 46 with hand pain.</td>
<td>Massage therapy once a week for 15 minutes for 4 weeks and taught self-massage to be done once daily vs. standard treatment control. Assessments on the first and last days of the 4 week study</td>
<td>First day post: mean pain massage 2.4 vs. control 2.6 (p &lt;0.05); mean grip strength 7.7 vs. 6.3 (p &lt;0.05); mean anxiety 27.19 vs. 30.2 (p &lt;0.001); mean depression 1.9 vs. 3.9 (p &lt;0.01). Last day post: mean pain 1.3 vs. 2.8 (p &lt;0.01); mean grip strength 8.5 vs. 6.7 (p &lt;0.005); mean anxiety 28.4 vs. 29.7 (p &lt;0.01); mean depression 1.4 vs. 3.9 (p &lt;0.05).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No mention of COI.</td>
<td>Mean age 50 years.</td>
<td>vs. 3.66 (p = 0.03); right hand lateral pinch 12 weeks 6.00 vs. 4.33 (p = 0.01); left hand chuck pinch 4.83 vs. 3.66 (p=0.01); left hand lateral pinch 12 weeks 5.15 vs. 4.33 (p=0.05). Median painful joint: 12 weeks 3.00 vs. 10.00 (p = 0.04).</td>
<td></td>
</tr>
<tr>
<td><strong>Physical Therapy/Mobilization</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wakefield 2000</td>
<td>RCT</td>
<td>No sponsorship OR COI.</td>
<td>N = 96 (72 female/9 male) with fracture of distal radius, previously treated by plaster immobilization</td>
<td>Taught and given standard sheet of home exercises by physiotherapist, referred for course of physiotherapy (n=49) vs. Taught and given standard sheet of home exercises only</td>
<td>Only flexion/extension at 26 weeks was significantly different (p=0.001) in the two group comparison via ANOVA. No significant differences were observed in parameters between groups. The physiotherapy group displayed significantly higher flexion/extension improvement at six months (p=0.044). There were no significant differences between each group at six months.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mean age of 72 years (55 – 90).</td>
<td>Follow up Week 6, Month 3, Month 6</td>
<td>“Our study has shown that home exercises are adequate rehabilitation after uncomplicated fracture of the distal radius, and routine referral for a course of physiotherapy should be discouraged. The role of physiotherapy in patients at high risk of a poor outcome requires further investigation.</td>
</tr>
</tbody>
</table>

Data suggest massage therapy group experience less pain, greater grip strength and a more positive mood vs. control group causing less anxiety and better sleep. Data suggest home exercises for uncomplicated fractures are beneficial.
Appendix 1. Dupuytren’s Disease

Dupuytren’s disease is a disorder of the hand involving the formation of fibrosis (scar tissue) in the palm and digits with subsequent contractures.(1593) It has strong age and inheritance patterns;(1594-1598) thus, it is generally thought to be non-occupationally related.(1599, 1600) Purported risks include the use of alcohol, smoking, diabetes mellitus, and epilepsy.(1594) However, although there are no quality studies involving occupational factors, there are some reported associations with both heavy(1601) and manual work.(1602) Therefore, to help provide improved care for patients, this disorder is included as an appendix to the Hand, Wrist, and Forearm Disorders Guideline.

Many treatments have been used for patients with Dupuytren’s disease, including radiotherapy, dimethylsulfoxide injections, topical applications of vitamins A and E, physical therapy, ultrasound, corticosteroid injections, 5-Fluouracil, and gamma interferon injections. Almost all of these treatments have been found ineffective.(1603) While surgery is currently the most effective treatment for Dupuytren’s disease, the contracture often reoccurs with time.

RADIOThERAPY
Radiotherapy has been used to attempt to slow or prevent the progression of Dupuytren’s disease.(1604) Treatment involves irradiating the nodules and cords associated with Dupuytren’s with x-rays or electrons.

Recommendation: Radiotherapy for Prevention of Progression of Dupuytren’s Disease
There is no recommendation for or against the use of radiotherapy to prevent the progression of Dupuytren’s disease.

Strength of Evidence – No Recommendation, Insufficient Evidence (I)
Level of Confidence – Low

Rationale for Recommendation
One moderate-quality trial of radiotherapy found no differences between two types of radiotherapy treatment regimens.(1604) However, the trial had no placebo group and there was no comparison between treatments. In addition, results suggested regression over 1 year. Radiotherapy is non-invasive and has moderate adverse effects, but it is moderately costly and there is no quality evidence of its efficacy. Therefore, there is no recommendation for or against the use of radiotherapy to prevent the progression of Dupuytren’s disease.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: radiotherapy, dupuytren contracture, dupuytrend disease, hand; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 4 articles in PubMed, 32 in Scopus, 1 in CINAHL, 0 in Cochrane Library, 2784 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, CINAHL, Cochrane Library, Google Scholar, and 0 from other sources. Of the 1 article considered for inclusion, 1 randomized trial and 0 systematic studies met the inclusion criteria.

COLLAGENASE INJECTIONS
Collagenase injections have been utilized for treatment of Dupuytren’s disease to lyse and rupture the finger cords that are causing the joint contracture.(1605-1607) (Recommendation: Collagenase Injections for Treatment of Dupuytren’s Disease
Collagenase injections are moderately recommended for treatment of Dupuytren’s disease.

Indications – Dupuytren’s contractures sufficient to result in impairment, nearing impairment, or sufficient to result in significant cosmetic deformity.

Frequency/Dose – Clostridial collagenase 10,000 U injection; repeat injection(s) at 4 to 6 week intervals.
Discontinuation – Resolution of contracture, sufficient reduction for patient to decline additional injection, adverse effects, or failure to respond to 3 injections.

Strength of Evidence – Moderately Recommended, Evidence (B)

Level of Confidence – Moderate

Rationale for Recommendation
Quality studies evaluating the efficacy of clostridial collagenase show considerable benefits.(1605, 1606, 1608, 1609) These injections are minimally invasive, have relatively few reported adverse effects (skin tears if prolonged contracture), but are costly. Therefore, collagenase injections are moderately recommended for treatment of Dupuytren's disease. One trial recommended post-operative manipulation but had no placebo or sham group.(1610)

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: collagenase injections, dupuytren contracture, dupuytren disease, hand; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 5 articles in PubMed, 68 in Scopus, 0 in CINAHL, 2 in Cochrane Library, 1126 in Google Scholar, and 0 from other sources. We considered for inclusion 2 from PubMed, 9 from Scopus, 0 from CINAHL, Cochrane Library, Google Scholar, and 0 from other sources. Of the 11 articles considered for inclusion, 7 randomized trials and 3 systematic studies met the inclusion criteria.

INTRA-OPERATIVE 5-FLUOROURACIL
5-Flourouracil (5-FU) is a chemotherapy drug that has been used for many years to treat cancer, principally as a thymidylate synthase inhibitor. It is administered intravenously or as a topical cream. 5-FU is also used in ophthalmic surgery as an anti-scarring agent, and topically to treat actinic (solar) keratoses and some types of basal cell skin carcinomas. 5-FU has also been used topically to attempt to slow or prevent recurrence of Dupuytren’s disease after surgery by reducing proliferation rates of fibroblasts.(1601)

Recommendation: 5-Fluorouracil for Recurrence of Dupuytren’s Disease in Surgical Patients
5-Fluorouracil is not recommended to prevent the recurrence of Dupuytren’s disease in surgical patients.

Strength of Evidence – Not Recommended, Evidence (C)

Level of Confidence – Low

Rationale for Recommendation
There is one moderate-quality trial of 5-fluorouracil administered intraoperatively which showed no difference when compared with placebo.(1601) 5-Fluorouracil is not invasive, but has adverse effects and is moderately costly. Therefore, 5-Fluorouracil is not recommended to prevent recurrence of Dupuytren’s disease.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: fluorouracil, 5 fluorouracil, dupuytren contracture, dupuytren disease, hand; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 0 articles in PubMed, 7 in Scopus, 0 in CINAHL, 0 in Cochrane Library, 1522 in Google Scholar, and 0 from other sources. We considered for inclusion 0 from PubMed, Scopus, CINAHL, Cochrane Library, 1 from Google Scholar, and 0 from other sources. Of the 1 article considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.
POST-OPERATIVE USE OF NSAIDS AND ACETAMINOPHEN
NSAIDs have been used to treat post-operative swelling from surgery for Dupuytren’s disease and appear to be superior to acetaminophen (paracetamol). (639) Naproxen may also be useful as an analgesic during the immediate post-operative phase. (639)

1. **Recommendation: NSAIDs to Treat Post-operative Swelling from Surgery for Dupuytren’s Disease**
   NSAIDs are moderately recommended to treat post-operative swelling from surgery for Dupuytren’s disease.
   
   **Indications** – Dupuytren’s disease surgical patients.
   
   **Frequency/Dose** – Naproxen 500mg BID. (639)
   
   **Duration** – Trial utilized 3 days of treatment.
   
   **Strength of Evidence** – Moderately Recommended, Evidence (B)
   
   **Level of Confidence** – High

2. **Recommendation: Acetaminophen for Dupuytren’s Surgery**
   Acetaminophen is recommended for Dupuytren’s surgery.
   
   **Frequency/Dose** – Paracetamol 1g QID trialed for 3 days. (Note: an FDA advisory committee recommended a maximum dose of 650mg and there is a suggestion of toxicity at 1g QID especially over a few days and particularly in patients consuming excess alcohol or who have liver disease.) (1611)
   
   **Strength of Evidence** – Recommended, Evidence (C)
   
   **Level of Confidence** – Moderate

**Rationale for Recommendations**
There is one high-quality study evaluating the effect of drugs on acute post-operative swelling after surgery for Dupuytren’s; it documents the efficacy and superiority of naproxen to paracetamol, which in turn was superior to placebo. (639) However, there is no quality evidence that other NSAIDs are inferior to naproxen. NSAIDs and acetaminophen are non-invasive, have low adverse effects (particularly over 3 days), and are low cost. Therefore, they are recommended to treat post-operative swelling and pain resulting from surgery for Dupuytren’s disease.

A comprehensive literature search was conducted using PubMed, Scopus, CINAHL, Cochrane Library, and Google Scholar without date limits using the following terms: anti-inflammatory agents, non-steroidal, dupuytren contracture, dupuytren disease, hand; controlled clinical trial, controlled trials, randomized controlled trial, randomized controlled trials, random allocation, random*, randomized, randomization, randomly; systematic, systematic review, retrospective, and prospective studies. We found and reviewed 1 articles in PubMed, 2 in Scopus, 0 in CINAHL, 1 in Cochrane Library, 440 in Google Scholar, and 0 from other sources. We considered for inclusion 1 from PubMed, 0 from Scopus, CINAHL, Cochrane Library, Google Scholar, and 0 from other sources. Of the 1 article considered for inclusion, 1 randomized trials and 0 systematic studies met the inclusion criteria.

**SURGERY**
Surgical procedures have long been used to attempt to improve range of motion in patients with contracture from Dupuytren’s disease. (1600) The goal of surgical care is to excise or incise the diseased fascia. This treatment does not cure the disease, but is meant to improve severe debilitating joint contractures. Several types of surgery have been used to treat Dupuytren’s disease, depending on the contracture. **Extensive fasciectomy** involves removing as much fascia as possible, including that which is grossly normal. Today, this procedure is not commonly performed because of increased morbidity which often included hematoma, edema, and prolonged post-operative stiffness. **Dermofasciectomy** removes the diseased fascia and the overlying skin. This requires resurfacing (covering) the wound with a full-thickness skin graft. Recurrence rates are quite low with this approach. Because of the radical nature of this procedure, it is usually reserved for patients with recurrent or severe disease. **Regional or selective fasciectomy** involves excising only grossly involved fascia. Although the disease process
clearly extends into clinically normal palmar fascia, this approach has proven successful in correcting MCP joint contractures and some PIP joint contractures; this procedure carries an acceptably low morbidity rate. Some surgeons prefer to leave the skin wound open to heal by secondary intention as a means of decreasing hematoma risk. This approach is commonly used today.

1. **Recommendation: Surgery for Treatment of Dupuytren’s Contracture**

   Surgery using the technique of regional or selective fasciectomy is recommended for contracture due to Dupuytren’s disease.

   **Strength of Evidence – Recommended, Insufficient Evidence (I)**
   **Level of Confidence – Low**

2. **Recommendation: Percutaneous Needle Fasciotomy (aka Needle Aponeurotomy) for Treatment of Dupuytren’s Contracture**

   Percutaneous needle fasciotomy (aka needle aponeurotomy) is not recommended for patients with contractures due to Dupuytren’s disease due to the high recurrence rates common with this technique.

   **Strength of Evidence – Not Recommended, Insufficient Evidence (I)**
   **Level of Confidence – Low**

3. **Recommendation: “Firebreak” Full-thickness Skin Graft for Dupuytren’s Contracture, Extensive Fasciectomy, or Dermofasciectomy for Treatment of Dupuytren’s Contracture**

   “Firebreak” full-thickness skin graft, extensive fasciectomy, or dermofasciectomy for Dupuytren’s contracture is not recommended for routine Dupuytren’s contracture surgery.

   **Strength of Evidence – Not Recommended, Evidence (C)**
   **Level of Confidence – Low**

**Rationale for Recommendations**

There are no quality studies comparing surgical results with non-surgical treatments or with no treatment. Considering the high propensity for Dupuytren’s contracture to progress or reoccur (estimated at 27 to 80% after surgery), surgical studies with sufficient follow-up to determine long-term benefits are needed. However, as some patients with functional limitations appear to improve at least in the short to intermediate term lasting many months to years, regional or selective fasciectomy is recommended. Surgery is invasive, has known adverse consequences including recurrences, and is costly. However, it also appears effective for at least a limited period of time and for some patients it may be the only treatment option available; thus, surgery is recommended particularly for patients with functional limitations. Full-thickness skin graft has been found to be ineffective and thus is not recommended.

**Evidence for Dupuytren’s Disease**

There are 2 high-quality and 15 moderate-quality RCTs incorporated in this analysis. There is also one other study included.
<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seegenschmiedt 2001 RCT</td>
<td>6.0</td>
<td>N = 129 (67 male and 62 female) with clinically evident and progressive early-stage DC. Mean age for Group A and B: 65 ± 11 / 61 ± 14 years.</td>
<td>Group A, radiotherapy 10 x 3 Gy (total dose, 30 Gy) in 2 series (5 x 3 Gy) separated by 8 weeks (N = 63) vs Group B, 7 x 3 Gy (total dose, 21 Gy) in 1 series within 2 weeks (N = 66). Follow-up 3 and 12 months.</td>
<td>At 12 months, reduction of symptoms, nodules and cord observed in both treatment groups (p &lt; 0.01). For subjective responses, 76 (59%) patients (Group A, 41; Group B, 35) stated &quot;regression of DC symptoms&quot; in 120 (61%) sites (A, 60; B, 60); range of regression equal for both groups: &lt;25% regression for 74 of 120 (62%) sites (A, 35; B, 39), 25-50% regression for 37 (31%) sites (A, 35; B, 19), 51-75% regression in 7 (6%) sites (A, 5; B, 2), and &gt;75% regression in 2 (2%) sites (all in group A); 46 (36%) patients (A, 19; B, 27) had &quot;stable condition&quot; in 65 (33%) sites (A, 30; B, 35), whereas 7 (5%) patients (A, 3; B, 4) suffered &quot;progression of DC symptoms&quot; in 13 (7%) sites (A, 5; B, 8).</td>
<td>&quot;Both tested RT regimens have been well accepted and tolerated by patients. Acute toxicity was slightly more enhanced in the low-dose group (21 Gy) than in the mediumdose group (30 Gy), probably due to the dose-time factor.&quot;</td>
<td>No placebo group. RT therapy individualized. Data suggest RT may be effective due to reported regression, but that cannot be proved.</td>
</tr>
<tr>
<td>Badalamente 2002 2 RCTs</td>
<td>8.0</td>
<td>First trial or IIA: N = 36 (31 male and 5 female) with MP joint contractures. Mean age 65 years. Second trial or IIB: IIA trial: Single dose of Collagenase injection of 10,000 U (N = 18 ) vs Placebo consisted of sterile normal saline containing</td>
<td>1 month after injection, 14/18 (77.8%) collagenase group had contracture correction to 0-5º vs. 2/18 (11.1%) placebo. Retreatment of 16 placebo patients who did not respond to 1st blinded injection had flexion contracture correction to 0-5º in 10 after a 1st open-label</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 1a. Quality Studies for the Treatment of Dupuytren’s Disease
<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Description</th>
<th>Intervention</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hurst 2009</td>
<td>308 (245 male and 63 female) with joint contractures of 20 degrees or more; mean age 62.7±9.5 years.</td>
<td>Treatment Group 0.58mg collagenase clostridium was injected into affected cords via 0.25ml of sterile diluent (MCP joints) or 0.20ml sterile diluent (PIP joints). Maximum of 3 injections every 30 days. Treatment cycle included injection, finger extension, and 30 day follow-up (N=204) vs Collagenase injected cords compared to placebo injections meeting primary endpoint (64.0% vs. 6.8%, P&lt;0.001). Collagenase joint ROM compared to placebo, 43.9 to 80.7 degrees vs. 45.3 to 49.5 degrees, (p &lt;0.001).</td>
<td>&quot;Collagenase clostridium histolyticum significantly reduced contractures and improved the range of motion in joints affected by advanced Dupuytren's disease.&quot;</td>
<td></td>
</tr>
<tr>
<td></td>
<td>10,000-U injection; in 2 after 2nd injection; in 1 after 3rd.</td>
<td></td>
<td>Cord I study. Data suggest that compared to placebo collagenase clostridium histolyticum significantly reduced contractures and increased ROM of joints in patients with Dupuytren's disease. Adverse effects (treatment related outcomes) significantly higher in collagenase group.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Age</td>
<td>Intervention</td>
<td>Follow-up</td>
</tr>
<tr>
<td>-------</td>
<td>---</td>
<td>-----</td>
<td>-------------</td>
<td>-----------</td>
</tr>
<tr>
<td>Badalamente 2007 RCT</td>
<td>7.5</td>
<td>18 years</td>
<td>Collagenase injection (10,000 U)</td>
<td>1, 7, 14, and 30 days</td>
</tr>
<tr>
<td>Gilpin 2007 RCT</td>
<td>7.0</td>
<td>66 (56 male and 10 female)</td>
<td>Treatment group: 0.58mg collagenase clostridium histolyticum per injection. Injected directly into Dupuytren’s affected cords. Maximum of 3 injections every 30 days. Treatment cycle included</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>Sponsorship</td>
<td>n/</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>--------</td>
<td>-------------</td>
<td>---</td>
</tr>
<tr>
<td>Auxilium, J.K. is on advisory board of Auxilium. N.J. is an employee of and owns stock options in Auxilium Pharmaceuticals.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mickelson 2014 RCT</td>
<td></td>
<td></td>
<td>No sponsorship or COI.</td>
<td></td>
</tr>
<tr>
<td>Witthaut 2011 Post Hoc RCT</td>
<td></td>
<td></td>
<td>Sponsored by Pfizer Inc. Cord study</td>
<td></td>
</tr>
</tbody>
</table>
sponsored by
Auxilium
Pharmaceuticals, Inc.
Jorg Witthaut is an
investigator for the
collagenase
Clostridium
histolyticum clinical
trial programme. The
remaining authors are
employees of Pfizer
Inc. Groton, CT, USA.

NEW
McGrouther
2014

| 4.0 | N = 58 with
Dupuytren's
contracture or
DC. Mean age
61.4 (8.89), 40
male and 18
female. | Collagenase
clostridium
histolyticum or CCH
injection treatment,
one joint
(N = 49)
vs
CCH Treatment
Primary 2 Joints
(N = 9).
Follow-up for 90
days. | Mean number of injections
per patient for up to 2
affected joints was 1.84,
mean injections per joint was
1.62.
Of the 56, 66% reported that
they were ‘very satisfied’ and
27% ‘quiet satisfied’, 4%
‘neither’, and 0% ‘very
dissatisfied’.
Commonly reported adverse
events; edema peripheral
reported by 79%, contusion
by 55%, pain in extremity by
41%, injection site
hemorrhage by 29% and
injection site pain by 29% of
patients.
| “Collagenase clostridium
histolyticum injection is a
minimally invasive procedure
that can be performed on
an outpatient basis.” |

Data from open label trial.
Data suggest CDH has some
efficacy for management of
DC.

## 5-Fluorouracil vs. Placebo Intraoperative

| Bulstrode
2004
| 5.0 | N = 15 male
with two-digit
disease. Mean
age 61 years. | Treatment rays, 5-
Fluorouracil a 1 cm
section of the
Dupuytren's tissue
was marked and
excised, plus
excision either 0.5
ml of 5-fluorouracil
(25 mg/ml) or 0.5 ml
vs
Metacarpophalangeal joint
movement improved from 68º
(range, 20-109º) to 85º
(range, 32-133º) for control
rays and 69º (range, 29-100º)
to 79º (range, 64-113º) for 5-
fluorouracil treated rays at 3
months. MCP joint range of
motion did not differ at 18
months.
| “The follow-up data have not
demonstrated a significant
difference between the control 5-
fluorouracil treated rays for either
total active motion, or
metacarpophalangeal or proximal
interphalangeal joint movement or
loss of extension.” |

Small sample size. Data
suggest 5-FU ineffective.
<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Details</th>
<th>Outcome</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jerosch-Herold 2011 RCT</td>
<td>5.5</td>
<td>N = 154 (120 male and 34 female) undergoing fasciectomy of dermofasciectomy for Dupuytren’s disease. Mean age hand therapy only 67.5±9.2 years, splint 67.2±10.0 years.</td>
<td>Hand therapy only within 2 weeks after surgery plus removal of sutures (N = 77) vs Hand therapy with night splinting worn for 6 months (N = 77). Follow-up for 12 months after surgery.</td>
<td>There were no statistically significant differences at 12 months between the two groups in DASH score (0.66, -2.79 to 4.11, p = 0.703), degrees of total active flexion of operated digits (-2.02, -7.89 to 3.85, p = 0.493), degrees of total active extension deficit of operated digits (5.11, 2.33 to -12.55, p = 0.172. The mean number of therapy sessions was 5.1 in the splint group and 5.6 in no-splint group. “No differences were observed in self-reported upper limb disability or active range of motion between a group of patients who were all routinely splinted after surgery and a group of patients receiving hand therapy and only splinted if and when contractures occurred.”</td>
</tr>
<tr>
<td>Kemler 2012 RCT</td>
<td>4.5</td>
<td>N = 54 with proximal interphalangeal (PIP) joint flexion contractures of at least 30º. Mean age 63 (9) and 64 (11) for hand therapy alone group, 8 female and 46 male.</td>
<td>Splint plus hand therapy (N = 28) vs Hand therapy alone (N = 26). Follow-up for 3 months.</td>
<td>After 1 year, the splint-plus-hand therapy had mean reduction of 21º in flexion contracture vs 29º in the group receiving hand therapy alone, (p = 0.1). 18 or 64% reported not less than “much improve” vs 19 or 73% of the 26 hand therapy alone, (p = 0.5). At 6 months pain did not differ significantly between group, VAS score 1.9 (2.0) vs 2.1 (2.4), (p = 0.7). “After operative release of a Dupuytren’s contracture, a postoperative protocol using a splint and hand therapy was no better than hand therapy alone in minimizing postoperative flexion contractures.”</td>
</tr>
<tr>
<td>Husby 2001 RCT</td>
<td>8.0</td>
<td>N = 35 (33 male and 2 female) Dupuytren’s contracture</td>
<td>Paracetamol1000mg 4 dimes daily (N = 12) vs Postoperative Dupuytren’s swelling as a percentage of preoperative volume: 5.6±3.8 vs. 6.9±3.7 vs. 8.2±5.1.</td>
<td>“Naproxen might have a clinical relevant effect on swelling when used on minor surgery in the hand, unlike paracetamol. Results suggest a beneficial effect of naproxen over paracetamol, which is superior to placebo, which the</td>
</tr>
<tr>
<td>Study</td>
<td>n</td>
<td>Procedure</td>
<td>Comparator</td>
<td>Outcomes</td>
</tr>
<tr>
<td>-------</td>
<td>---</td>
<td>-----------</td>
<td>------------</td>
<td>----------</td>
</tr>
<tr>
<td>No mention of sponsorship or COI.</td>
<td>6.0</td>
<td>N = 121 (94 male and 19 female) or 125 hands, with Dupuytren’s disease. Mean age 63 years.</td>
<td>Percutaneous needle fasciotomy or PNF (N = 57) vs Limited fasciotomy under either regional anesthesia or general anesthetist using tourniquet in all cases (N = 56).</td>
<td>Follow-up for at 1 and 6 weeks for the primary outcome perimeters.</td>
</tr>
<tr>
<td>van Rijssen 2006 RCT</td>
<td>5.5</td>
<td>N = 111 (17 female and 76 male) with affected hands and minimal passive extension deficit of 30 degrees. Mean age for LF and PNF groups; 63.1 and 62.8.</td>
<td>Limited fasciotomy or LF (N = 41) vs Percutaneous needle fasciotomy or PNF (N = 52).</td>
<td>Follow-up at 1 and 6 weeks, 6 months, and 1, 2, 3, 4 and 5 years.</td>
</tr>
</tbody>
</table>

**Surgical Procedures**

- **PNF:** Largest mean TPED per ray contractures 1 week after PNF 30° (58% reduction), p = 0.001. Follow-up at 6 weeks, results better. Limited fasciotomy: mean TPED at 1 week 15° (73% reduction), p = 0.001. Largest reduction for PNF at MCP, but DIP for LF.

- **LP:** Largest mean TPED per ray contractures 1 week after LP 30° (58% reduction), p = 0.001. Follow-up at 6 weeks, results better. Limited fasciotomy: mean TPED at 1 week 15° (73% reduction), p = 0.001. Largest reduction for LP at MCP, but DIP for LF.

- **In the short term and in cases with a TPED of 90° or less PNF is a good treatment alternative to LP for treatment of Dupuytren’s disease.."**

- **Data suggest that at 5 years, the recurrence rate in the needle fasciotomy group was (84.9%) compared to the limited fasciotomy group (20.9%) and recurrence occurred earlier the needle fasciotomy group.**
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Participants</th>
<th>Procedure</th>
<th>Follow-up</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>NEW Kan</td>
<td>2016</td>
<td>RCT</td>
<td>N = 80 (62 male and 14 female) with primary Dupuytren’s contracture. Mean age 63 ± 9 for PALF and 63 ± 8 for LF group.</td>
<td>Procedure consisting of extensive percutaneous aponeurotomy and lipofilling or PALF (N = 40) vs Limited fasciectomy (N = 40). Follow-up at 2 weeks, 3 weeks, 6 months and 1 year.</td>
<td>At 1 year, 15/85 PALF treated joints or 18%, had some recurrence vs 5/58 limited fasciectomy treated joints or 9%, (p = 0.107). The overall complication rate not significantly different between the groups, (p = 0.402).</td>
<td>“PALF demonstrates a significantly shorter convalescence, similar operative contracture correction, lower incidence of long-term complications, and no significant difference regarding 1-year postoperative results compared with limited fasciectomy.” Data suggest PALF showed shorter recovery times, fewer complications, comparable results to standard fasciotomy group. However, at one year post procedure the PALF group had more recurrence (18% vs.9%).</td>
</tr>
<tr>
<td>Ullah</td>
<td>2009</td>
<td>RCT</td>
<td>N = 79 (65 male and 14 female) with Dupuytren’s contracture of the proximal interphalangeal joint. Mean age of 62.9 years.</td>
<td>Dermofasciectomy or “Firebreak” skin graft performed on one finger (N = 39) vs Fasciectomy was performed on second finger (N = 40). Follow up at 12, 24 and 36 months.</td>
<td>Mean range of movement of PIP 34.6º (1-80º) preoperatively, improved to 85º (2-98º) at 3 years. Progressive recurrence of PIP contracture over 3 years in 11 (12.2%); 5 had fasciotomy with Z-plasty; contracture recurred in 5.4 months vs. 8months for full-thickness skin graft (p = 0.6).</td>
<td>“[N]o difference in recurrence rates between the two methods of treatment at three years and we were surprised at the low recurrence rate after fasciectomy and Z-lasty alone.” Suggests full thickness graft not more effective.</td>
</tr>
<tr>
<td>Citron</td>
<td>2005</td>
<td>RCT</td>
<td>N = 100 63 male and 16 female) with Dupuytren’s disease in one ray only and any degree of resultant</td>
<td>Modified Brunner incision closed with multiple Y-V plasties (N = 62) vs Z-plasty group had longitudinal incision,</td>
<td>Mean post-op deformity on final review or at recurrence 25º in modified Brunner group vs. 24º in Z-plasty group (NS). Recurrence rate 33% modified Brunner vs. 18% Z-plasty group (NS).</td>
<td>“There is no evidence to suggest that the type of incision influences the time distribution of recurrent disease but this possibility cannot be discounted.” Data suggest no differences between the 2 procedures.</td>
</tr>
<tr>
<td>Study</td>
<td>Participants</td>
<td>Procedures</td>
<td>Follow-up</td>
<td>Results</td>
<td>Conclusion</td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>--------------</td>
<td>------------</td>
<td>-----------</td>
<td>---------</td>
<td>------------</td>
<td></td>
</tr>
<tr>
<td>Bhatia 2002 Prospective RCT</td>
<td>N = 31 (28 male and 3 female) undergoing surgery for Dupuytren's disease; age range 38-79 years.</td>
<td>Staple group: staples via an automatic stapling device. Time spent closing recorded. Pain levels recorded during staple removal at 1 week follow-up (N = 13) vs Suture group: received 4-0 monofilament polybaster sutures. Time spent closing recorded. Pain levels recorded during suture removal at 1 week follow-up (N = 18).</td>
<td>Mean skin closure time with sutures 51 seconds per cm and 25 seconds per cm with staples (p &lt;0.001). The mean pain score for removal 2.4 for suture removal and 5.2 for staple removal, (p = 0.008).</td>
<td>&quot;As staples can be inserted in half the time of conventional sutures we recommend their use for closure of extensive palmar wounds following long operative procedures.&quot;</td>
<td>Data suggest patient pain was higher for staple removal over suture removal but staples took less time to insert and no significant differences in wounds once staples or sutures removed.</td>
<td></td>
</tr>
</tbody>
</table>
Appendix 2: Excluded Studies
(Low-quality Randomized Controlled Trials and Non-randomized Studies)
The following low-quality randomized controlled studies (RCTs) and other studies were reviewed by the Evidence-based Practice Hand, Wrist, and Forearm Panel to be all inclusive, but were not relied upon for purposes of the development of this document’s guidance on treatments because they were not of high quality due to one or more errors (e.g., lack of defined methodology, incomplete database searches, selective use of the studies and inadequate or incorrect interpretation of the studies’ results, etc.), which may render the conclusions invalid. ACOEM’s Methodology requires that only moderate- to high-quality literature be used in making recommendations.(1617)

ERGONOMIC INTERVENTIONS

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ripat 2006</td>
<td>RCT</td>
<td>Sponsored by Manitoba Hydro. No mention of COI.</td>
<td>3.0</td>
<td>N = 68 with two or more symptoms of WRUED (Work Related Upper Extremity Disorders). Mean age 42.2 years.</td>
<td>Adapted Group- Microsoft Natural MultiMedia Keyboard adapted to reduce activation force required to depress keys (light touch) (n = 43) vs. Unadapted Group- Standard keyboard with no adaptations made (n = 25). Follow-up for 6 months.</td>
<td>No significant differences between two groups for Symptom Severity (SSS) and Functional Status Scales (FSS) between groups (p &lt;0.05). When data from groups combined, SSS and FSS-typing measures significant at both 12 and 24 week (p &lt;0.0001) at both time points.</td>
<td>&quot;Positive results in reduction of symptom severity and improvement in functional status were identified for participants in both keyboard study groups, providing further evidence to support the use of ergonomic keyboards for individuals with WRUED. The vast majority of participants were satisfied with their study keyboard.&quot;</td>
<td>Both keyboard groups improved over time, however, there were no differences between groups. Some randomized to experimental group were &quot;forced&quot; to use the LT keyboard.</td>
</tr>
<tr>
<td>Hedge 1999</td>
<td>RCT</td>
<td>Sponsored by Honeywell, Inc., Proformix, Inc., Global, Global Contrac and Teknion. No mention of COI.</td>
<td>2.5</td>
<td>N = 38 professionals who used a computer work average of 5.4 hours per day. Mean age 37.4.</td>
<td>DT Group- DT keyboard tray. User measurements taken to put keyboard at comfortable height. (n = 23) Vs. Control Group- conventional adjustable keyboard with or without a padded wrist rest (n = 15). Measurements taken immediately following intervention.</td>
<td>No significant differences between pre- and post-test measurements in the control group for wrist extension and ulnar deviation (p &gt;0.05). Significant difference between pre and post wrist extension in DT group; 17.6 vs. 12.1 (p &lt;0.05). No significant difference for ulnar deviation. (p&gt;0.05). In post-test upper posture index (UPI) there was a significant difference in favor of the DT group vs. control for number of subjects reporting a UPI &lt; 4; 60% vs. 90% (p = 0.044).</td>
<td>&quot;Overall, the wrist movement data, the RULA data and the self-reported musculoskeletal discomfort data all point to improvements within a short time after using the DT system.&quot;</td>
<td>Methodological details sparse.</td>
</tr>
</tbody>
</table>
Nurse case manager training in ICN vs. no ICM training.

"Trained nurses were more likely to recommend accommodations addressing workstation layout, computer-related improvements, furnishings, accessories, and lifting/carrying aids, whereas the untrained nurses were more likely to suggest light duty and lifting restrictions. This study indicates that the training was associated with a change in the practice behavior of case managers regarding the workplace accommodation process."

"More research is needed to identify barriers to implementation and develop more effective approaches to facilitate worksite accommodations in disabled workers with carpal tunnel syndrome and other persistent upper extremity disorders."

N = 101

Mean rate of data entry under supplementary rest break schedule significantly faster than rate under conventional rest break schedule (p <0.0002). No significant effects of stretching on discomfort or performance observed. Discomfort and eyestrain significantly lower with supplementary breaks; supplementary breaks attenuated accumulation of discomfort and eyestrain during work sessions.

"These results provide further converging evidence that supplementary breaks reliably minimize discomfort and eyestrain without impairing productivity."

"Positive results in reduction of symptom severity and improvement in functional status were identified for participants in both keyboard study groups, providing further evidence to support the use of ergonomic keyboards for individuals with WRUED. The vast majority of Both keyboard groups improved over time, however, there were no differences between groups. Some randomized to experimental group were "forced" to use the LT keyboard."
keyboard with no adaptations made. (n = 25) Follow-up for 6 months.

participants were satisfied with their study keyboard.”

Hedge 1999
RCT
Sponsored by Honeywell, Inc., Proformix, Inc., Global, Global
Contrac and Teknion. No mention of COI.

N = 38 professional workers who used a computer at work for an average of 5.4
hours per day. Mean age 37.4.

DT Group- DT keyboard tray. User
measurements taken to put keyboard at
comfortable height (n = 23) vs.
Control Group conventional adjustable
keyboard with/without padded wrist rest.
(n = 15). Measurements immediately
following intervention.

No significant differences between
pre- and post-test measurements
in control group for wrist extension
and ulnar deviation (p >0.05). Significant
difference between pre-
and post-wrist extension in DT
group; 17.6 vs. 12.1 (p <0.05). No
significant difference for ulnar
deviation. (p >0.05). In post-test
upper posture index (UPI) there
was a significant difference in
favor of DT group vs. control for
number of subjects reporting UPI
<4; 60% vs. 90% (p = 0.044)

“Overall, the wrist movement data, the RULA
data and the self-reported
musculoskeletal discomfort
data all point to
improvements within a
short time after using the
DT system.”

Methodological details
sparse.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feuerstein 1993</td>
<td>Non-randomized comparative study</td>
<td>N/A</td>
<td>N = 49</td>
<td>Eligible for multi-component rehab program (n = 34) vs. not eligible (n = 15)</td>
<td>Findings indicated “74% of the treatment group returned to work or were involved in state-supported vocational training in contrast to 40% of the control group (p &lt;0.05).”</td>
<td>“These findings suggest the need to modify treatment components to facilitate an increased return-to-work rate. Areas that may prove useful include a greater emphasis ergonomic modifications at the workplace to reduce the risks of repetitiveness, force, awkward posture, and insufficient work/rest cycles, as well as efforts to modify work</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
CARPAL TUNNEL SYNDROME – DIAGNOSTICS
Electrodiagnostic Studies

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Study Type</th>
<th>Conflict of Interest (COI)</th>
<th>Population/ Case Definition</th>
<th>Investigative Test</th>
<th>Gold Standard / Comparative Test</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jackson 1989</td>
<td>3.5</td>
<td>Diagnostic</td>
<td>N =162 divided into groups: Group 1 (n = 38) healthy volunteers. Group 2 (n = 40) with positive clinical testing but negative Electromyography (EMG) and Nerve Conduction Studies. Group 3 (n = 53) clinical confirmed CTS, positive NCS, Negative EMG. Group 4 (n = 30) clinically confirmed CTS, positive NCS and EMG.</td>
<td>Electrodiagnosti c studies including Palm median nerve latency (Palm (m)). Distal Sensory latency difference between median and radial nerve (DSL (m-r)). Palmar latency difference between median and ulnar nerve (Palm (m-u)). Distal Sensory latency difference (DSL (m-u)). Amplitude of sensory action potential ratios and the 2nd and 5th digit. (Amp).</td>
<td>Screening history as well as physical diagnostic testing was used to ensure the existence of Carpal Tunnel Syndrome, as well as eliminate patients with peripheral neuropathy.</td>
<td>Abnormality percentages of different tests Group 1, 2, 3, 4: Palm (m): 2.6%, 16%, 96%, 94%. DSL (m-r): 0%, 44%, 89%, 100%. DSL (m-u): 5.3%, 44%, 100%, 100%. Palm (m-u): 5.3%, 30%, 98%, 94%. Amp: 0%, 2.3%, 33%, 61%. Group 2 abnormalities using a combination of tests: DSL (m-u) and DSL (m-r): 51%. DSL (m-u) DSL (m-r) and Palm (m-u) 51%.</td>
<td>“Certainly supplemental studies can serve as a discriminating instrument, distinguishing between individuals on a dimension of interest (NCS) when no gold standard is available for validating these measures.”</td>
<td>Study suggests use of comparing median and radial distal sensory latencies in digit 1 and comparing median and ulnar distal sensory latencies in digit 4 when CTS referrals have normal nerve conduction studies.</td>
<td></td>
</tr>
<tr>
<td>Zaher 2012</td>
<td>3.0</td>
<td>Diagnosti c</td>
<td>N=52 with CTS. Follow-up at 12 weeks.</td>
<td>Electrodiagnosti c Studies (n = 20)</td>
<td>MRI (n = 10) vs. Ultrasound (n = 22).</td>
<td>17/20 (85%) had electrodiagnostic findings of prolonged motor and sensory latencies of the median nerve, reduced sensory and motor conduction velocities, and median-ulnar sensory latency difference. 10/10 (100%) underwent MRI showed swelling of media nerve, increased signal intensity, and palmar bowing of transverse carpal ligament. 19/22 (86.3%) with ultrasounds showed enlargement of median nerve at proximal carpal tunnel with increased cross-sectional area over 12 mm2, and palmar bowing and thickening of flexor reticulum.</td>
<td>“Ultrasound is superior to other investigation tools as it provides accurate and rapid diagnosis of CTS with the least cost.”</td>
<td>Study enrolled only subjects with mild CTS. Study suggests ultrasound is superior to other diagnostic techniques for mild CTS due to its relatively low cost and raped results MRI and electrodiagnostic studies did have better diagnostic outcomes.</td>
<td></td>
</tr>
<tr>
<td>Homann 1999</td>
<td>3.0</td>
<td>Diagnostic</td>
<td>N = 824 workers recruited from 6 different companies, with a mean job tenure 8.9±9.1 years</td>
<td>Electrodagnostic testing of median-Ulnar sensory peak latency difference &gt;0.5 ms, more severe was a difference of &gt;0.8 ms.</td>
<td>Self-administered surveys and hand diagrams. Questionnaire asked about symptom severity, and persistence. Workers indicated pain, numbness, and areas of tingling on hand diagram.</td>
<td>Electrodiagnostic (EDX) positive results (n = 139, 16.9%), Physical Examination (PE) positive (n = 165, 20.1%), Wrist, Hand, and Finger Symptoms (WHF Sx) positive (n = 305, 37.0%). Correlation between PE and EDX (n = 36), between WHF Sx and PE (n = 90), EDX and WHF Sx (n = 55). Between all 3 tests (n = 23).</td>
<td>“The combination of results from electrodiagnostic testing and symptom survey procedures appears to provide the best criterion for defining CTS for epidemiologic investigations in which the intent is to evaluate either the impact of intervention or the exposure-response relationship.”</td>
<td>Study reports poor correlation between electrodiagnostic findings, symptom surveys and symptom presentation from physical exams in diagnosing CTS.</td>
<td></td>
</tr>
<tr>
<td>Author/Year</td>
<td>Score</td>
<td>Conflict of Interest (COI)</td>
<td>N</td>
<td>Area of Upper Extremity</td>
<td>Diagnoses</td>
<td>Type of Ultrasound</td>
<td>CT used</td>
<td>MRI Used</td>
<td>More than on rater</td>
</tr>
<tr>
<td>------------</td>
<td>-------</td>
<td>---------------------------</td>
<td>---</td>
<td>-------------------------</td>
<td>-----------</td>
<td>-------------------</td>
<td>---------</td>
<td>----------</td>
<td>------------------</td>
</tr>
<tr>
<td>Uncini 1989</td>
<td>2.5</td>
<td></td>
<td>43 with symptoms and signs of CTS and 33 controls. Group 1: (26 hands) mild abnormalities DPSNL &gt;2.9 msec onset, 3.5 msec peak; group 2: (16 hands) normal or borderline median nerve results D2SNL &lt;2.9 msec, onset or 3.5 msec peak.</td>
<td>Electrodiagnostic studies: median DML, wrist to abductor pollicis brevis (APB); ulnar DML, wrist to abductor digiti minimi (ADM); median sensory nerve latency (SNL) D2 to wrist and D2 to palm; ulnar SNL D5 to wrist; median and ulnar SNL D4 to wrist.</td>
<td>N/A</td>
<td>Both groups had longer median latencies from digit 4 to wrist than digit 2 to wrist. D4 latencies more significant in group 1 (D4 latency onset: 3.7±0.5 and D2: 3.3±0.2) and group 2 (3.0±0.4 and 2.6±0.2) than D2 latencies, suggesting D4 more sensitive than D2-Wr. Significant differences in paired nerves (adjusted for controls) of median D4 SNL - ulnar IV DSL vs. median DML - ulnar DML (group 1: p &lt;0.05 and group 2: p &lt;0.05), and median D4 SNL - ulnar D4 SNL vs. median D2 SNL - ulnar D4 SNL (group 1: p &lt;0.05 and group 2: p &lt;0.05). Meant D4 technique most sensitive for disease detection.</td>
<td>“In conclusion, stimulating digit 4 and comparing latencies to median and ulnar nerves is a simple method that is more sensitive than other techniques in detecting CTS. Detection of the double peak potential recorded over the median nerve allows immediate diagnosis of CTS. Even when the double peak is not recognized, a median and ulnar D4 latency difference greater than 0.5 msec suggests CTS.”</td>
<td>Study suggests stimulation of digit 4 is useful in identification of CTS. D4 latency is longer in CTS patients compared to other digits.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Study Year</td>
<td>N</td>
<td>Wrist Location</td>
<td>Patient Population</td>
<td>Duration of Symptoms</td>
<td>Exam Findings</td>
<td>Nerve Conduction Study Findings</td>
<td>Sensitivity</td>
<td>Specificity</td>
</tr>
<tr>
<td>---------------</td>
<td>------------</td>
<td>----</td>
<td>----------------</td>
<td>-------------------</td>
<td>----------------------</td>
<td>--------------</td>
<td>---------------------------------</td>
<td>-------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Wiesler 2006</td>
<td>Diagnosti c</td>
<td>3.0</td>
<td>Wrist</td>
<td>Patients with symptoms, clinical exam findings, and nerve conduction study findings for CTS. Mean duration of symptoms 12 months (range 1.5-72 months). Mean age 56 years CTS, 36 years controls.</td>
<td>-</td>
<td>-</td>
<td>Philips HDI 5000 with 12.5-MHz linear-array transducer</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Pearson correlation coefficient ultrasound vs. nerve conduction study (NCS): 0.37 (p = 0.013). Sensitivity and specificity: cutoff point of 11+ mm² = sensitivity 91%, specificity 84%; PPV 74%; NPV 95%.</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yesilda g 2004</td>
<td>Diagnosti c</td>
<td>3.0</td>
<td>Wrist</td>
<td>CTS symptoms. Mean age CTS 49.8±8.7 years, controls 42.7±11.3 years.</td>
<td>-</td>
<td>-</td>
<td>12 MHz linear array transducer (ATL 1500 HDI)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Mean±SD cross-sectional area by tracing method: CTS 14.9±4.7 vs. control 7.8±1.6 (p &lt;0.001). Mean±SD cross-sectional area by ellipsoid formula: CTS 14.2±4.5 vs. control 7.5±1.8 (p &lt;0.001). Cutoff for sensitivity and specificity: 10.5mm² for mean cross-sectional area; using tracing method – sensitivity (95% CI) 89.9 (85-94.8), specificity 94.7 (89.7-99.7), PPV 97 (94.3-99.9), NPV 82.7 (74.8-90.6); using indirect method – sensitivity 86.5 (81-92), specificity 93.4 (97.88-99), PPV 96.2 (92.9-99.4), NPV 78.1 (69.5-86.6).</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

"[H]igh-resolution ultrasound is informative in the evaluation of CTS and shows enlargement of the median nerve at the distal wrist crease in symptomatic patients."

A 1:2 (CTS vs. normal). Suggests HRUS may be used to diagnose CTS and enlargement of the median nerve at the wrist crease in symptomatic patients is usually predictive for CTS.

2:1 matched study suggesting ultrasonographic measurement of the median nerve cross-sectional area is a sensitive, specific and useful non-invasive method for the diagnosis of carpal tunnel syndrome."
17/20 (85%) had electrodiagnostic findings of prolonged motor and sensory latencies of median nerve, reduced sensory and motor conduction velocities, and median-ulnar sensory latency difference. 10/10 (100%) underwent MRI showed swelling of median nerve, increased signal intensity, and palmar bowing of transverse carpal ligament. 19/22 (86.3%) with ultrasounds showed an enlargement of median nerve at proximal carpal tunnel with increased cross-sectional area over 12 mm², and palmar bowing and thickening of flexor reticulum.

"Ultrasound is superior to other investigation tools as it provides accurate and rapid diagnosis of CTS with the least cost."

Study enrolled only subjects with mild CTS. Study suggests ultrasound superior to other diagnostic techniques for mild CTS due to its relatively low cost and rapid results MRI and electrodiagnostic studies did have better diagnostic outcomes.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Age</th>
<th>Gender</th>
<th>MRI Equipment</th>
<th>FWHM</th>
<th>Diffusion Metrics</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Guggenberg 2012</td>
<td>Diagnostic</td>
<td>35</td>
<td>35</td>
<td>M</td>
<td>3.0 T MR imager</td>
<td>-</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Horng 2012</td>
<td>Diagnostic</td>
<td>35</td>
<td>35</td>
<td>M</td>
<td>GE 1.5 T Signa Excite MRI system</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Bak 1997</td>
<td>Diagnostic</td>
<td>35</td>
<td>35</td>
<td>M</td>
<td>1.5 T Philips ACS-NT superconductive MR unit</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
</tbody>
</table>

"Normative diffusion values for MR neurography of the median nerve with DTI depend on the anatomic location and age but not on sex." Study suggests ultrasonography comparable to MRI in diagnosing CTS only if both rest and grasp position are combined. Sample size small.
<table>
<thead>
<tr>
<th>Year</th>
<th>Diagnostic</th>
<th>Study Design</th>
<th>N</th>
<th>Condition</th>
<th>MRI</th>
<th>EMG</th>
<th>Wrist</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
<th>Forearm</th>
</tr>
</thead>
<tbody>
<tr>
<td>2003</td>
<td>Deryani</td>
<td>Small sample size. Study suggests MRI in tandem with electrophysiological evaluation to make CTS diagnosis.</td>
<td>55</td>
<td>CTS, of those with CTS and healthy subjects</td>
<td>(-)</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>(-)</td>
<td>(-)</td>
<td>+</td>
<td>(-)</td>
<td>Statistically significant differences between median nerve diameters (at pisiform bone level: 8.47 ± 1.41 mm; and distal radio ulnar joint level: 4.04 ± 1.06 mm and 2.42 ± 0.95 mm), the diameter rations and flexor retinaculum bulging rations (26.21 ± 5.98% and 7.27 ± 4.53%), (p &lt; 0.001). Hyperintensity was fond in 4 of 25 controls and isointensity in 21, (p &lt; 0.001).</td>
<td>[MRI] examination of structural changes that occur in the carpal tunnel, neighboring structures and the median nerve would be useful in the diagnosis of CTS, especially in case with suspected clinical and electrophysiological diagnosis.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
17/20 (85%) had electrodiagnostic findings of prolonged motor and sensory latencies of median nerve, reduced sensory and motor conduction velocities and median-ulnar sensory latency difference. 10/10 (100%) underwent MRI showed swelling of median nerve, increased signal intensity, and palmar bowing of transverse carpal ligament. 19/22 (86.3%) with ultrasounds showed an enlargement of the median nerve at proximal carpal tunnel with an increased cross-sectional area over 12 mm², and palmar bowing and thickening of flexor reticulum.

“Ultrasound is superior to other investigation tools as it provides accurate and rapid diagnosis of CTS with the least cost.”

Study enrolled only subjects with mild CTS. Study suggests ultrasound superior to other diagnostic techniques for mild CTS due to its relatively low cost and rapid results MRI and electrodiagnostic studies did have better diagnostic outcomes.

### CARPAL TUNNEL SYNDROME – TREATMENT

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Exercise</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Horng 2011</td>
<td>RCT</td>
<td>3.5</td>
<td>N = 60 with CTS. Mean age 50.5+9.4 years.</td>
<td>Difference between before and after treatment: Symptom severity – Group 1: -0.7+0.8; Group 2: -0.3+0.6; Group 3: -0.6+0.6; p = 0.56; Functional status – Group 1: -0.4+0.5; Group 2: 0.1+0.5; Group 3: -0.2+0.7 p = 0.04; Pain scale – Group 1: -19.7+24.6; Group 2: -10.5+18.0; Group 3: -17.2+26.2; p = 0.44</td>
<td>“To improve the functional status and quality-of-life of CTS patients, the combination of tendon gliding exercises, paraffin therapy, and splinting might be more effective than the combination of nerve gliding exercises, paraffin therapy, and splinting.”</td>
<td>Baseline comparability differences in functional status scores of the three groups.</td>
</tr>
<tr>
<td>Heebner 2008</td>
<td>RCT</td>
<td>2.0</td>
<td>N = 60 diagnosed with CTS by physician. Mean age 52</td>
<td>No statistical difference reported between groups. P-values not provided. Compared to baseline, follow-up scores for median nerve</td>
<td>“The results of this study suggest that persons with CTS in a community hospital do not benefit from a one-time nonsurgical methodological details sparse.”</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Duration</td>
<td>N</td>
<td>Age</td>
<td>Intervention</td>
<td>Outcome</td>
<td>Comments</td>
</tr>
<tr>
<td>-------</td>
<td>----------</td>
<td>---</td>
<td>-----</td>
<td>-------------</td>
<td>---------</td>
<td>----------</td>
</tr>
<tr>
<td>Tal-Akabi 2000</td>
<td>2.0</td>
<td>N = 21 with CTS mean duration of 2.3±2.5 years from surgery waiting list. Mean age: 47.1±14.8 years.</td>
<td>N = 21 with CTS mean duration of 2.3±2.5 years from surgery waiting list. Mean age: 47.1±14.8 years.</td>
<td>Neurodynamic mobilization (ULTT2a) (n = 7) vs. Carpal bone mobilization (n = 7) vs. No treatment (n = 7). Follow-up and intervention length are uncertain.</td>
<td>Only the post-intervention Pain Relief Scale (PRS) demonstrated significant difference between the three groups (p&lt;0.01). Mean PRS – Neurodynamic: 3.14; Carpal Bone: 3.71; Control: 0.</td>
<td>&quot;The study has failed to show significant differences in the effectiveness between mobilization of the median nerve and carpal bone mobilization in the treatment of patients presenting with carpal tunnel syndrome.&quot;</td>
</tr>
<tr>
<td>Bardak 2009</td>
<td>1.5</td>
<td>N = 111 (111 hands) with CTS. Mean age 49.14±9.6 years.</td>
<td>N = 111 (111 hands) with CTS. Mean age 49.14±9.6 years.</td>
<td>Group 1 standard conservative treatment (SCT) (n = 41) vs. Group 2 SCT plus tendon and median nerve gliding exercises (n = 35) vs. Group 3 tendon and median nerve gliding exercises (n = 35). Follow-up 2 and 11 months.</td>
<td>Symptom total point change – Group1: -7.4; Group 2: -10.5; Group 3: -2.9. Significant difference between Groups 1 and 3 and Groups 2 and 3 (p&lt;0.001). Functional status scale change – Group 1: -6.7; Group 2: -6.7; Group 3: -3.8. Significant difference between Groups 1 and 3 and Groups 2 and 3 (p &lt;0.001).</td>
<td>&quot;In conclusion, in cases of idiopathic CTS, conservative treatment is clinically effective. Adding tendon and nerve gliding exercises is also beneficial to the management of long-term CTS. Tendon and nerve gliding exercises alone are inferior to other modalities.&quot;</td>
</tr>
<tr>
<td>Gurcay 2009</td>
<td>3.5</td>
<td>N =32 female, housewife patients with clinically and EDS confirmed mild or moderate CTS. Mean age 40.8±11.2 years.</td>
<td>N =32 female, housewife patients with clinically and EDS confirmed mild or moderate CTS. Mean age 40.8±11.2 years.</td>
<td>Group A: local injection of 6mg betamethasone through 25-guage needle near distal wrist-flexion crease (n = 18) vs. Group B: meloxicam 15 mg/day, PO, for 3 weeks (n = 14). Both groups advised to wear wrist splints in neutral position at night for 3 weeks. Follow-up at 3 months.</td>
<td>No significant difference found between the groups for Functional Status Scale (FSS) scores, Jebsen Taylor Test (JTT) scores, or electrophysiological findings at 3 months (p&lt;0.05).</td>
<td>&quot;[T]he two treatment methods resulted in some functional gains in hand dexterity and improvement in electrophysiological data, but with neither of the methods demonstrating superiority.&quot;</td>
</tr>
<tr>
<td>Stransky 1989</td>
<td>3.5</td>
<td>N = 15 EDS confirmed</td>
<td>N = 15 EDS confirmed</td>
<td>200mg of Vitamin B6 vs. placebo</td>
<td>&quot;Significant changes in nerve conductions and EMGs did not occur when initial and follow-up data were compared. Clinical findings</td>
<td>&quot;Vitamin B6 seems to have no advantage over conservative therapy for carpal tunnel syndrome.&quot;</td>
</tr>
</tbody>
</table>

**NSAIDs**

<table>
<thead>
<tr>
<th>Study</th>
<th>Duration</th>
<th>N</th>
<th>Age</th>
<th>Intervention</th>
<th>Outcome</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tal-Akabi 2000</td>
<td>2.0</td>
<td>N = 21 with CTS mean duration of 2.3±2.5 years from surgery waiting list. Mean age: 47.1±14.8 years.</td>
<td>N = 21 with CTS mean duration of 2.3±2.5 years from surgery waiting list. Mean age: 47.1±14.8 years.</td>
<td>Neurodynamic mobilization (ULTT2a) (n = 7) vs. Carpal bone mobilization (n = 7) vs. No treatment (n = 7). Follow-up and intervention length are uncertain.</td>
<td>Only the post-intervention Pain Relief Scale (PRS) demonstrated significant difference between the three groups (p&lt;0.01). Mean PRS – Neurodynamic: 3.14; Carpal Bone: 3.71; Control: 0.</td>
<td>&quot;The study has failed to show significant differences in the effectiveness between mobilization of the median nerve and carpal bone mobilization in the treatment of patients presenting with carpal tunnel syndrome.&quot;</td>
</tr>
<tr>
<td>Bardak 2009</td>
<td>1.5</td>
<td>N = 111 (111 hands) with CTS. Mean age 49.14±9.6 years.</td>
<td>N = 111 (111 hands) with CTS. Mean age 49.14±9.6 years.</td>
<td>Group 1 standard conservative treatment (SCT) (n = 41) vs. Group 2 SCT plus tendon and median nerve gliding exercises (n = 35) vs. Group 3 tendon and median nerve gliding exercises (n = 35). Follow-up 2 and 11 months.</td>
<td>Symptom total point change – Group1: -7.4; Group 2: -10.5; Group 3: -2.9. Significant difference between Groups 1 and 3 and Groups 2 and 3 (p&lt;0.001). Functional status scale change – Group 1: -6.7; Group 2: -6.7; Group 3: -3.8. Significant difference between Groups 1 and 3 and Groups 2 and 3 (p &lt;0.001).</td>
<td>&quot;In conclusion, in cases of idiopathic CTS, conservative treatment is clinically effective. Adding tendon and nerve gliding exercises is also beneficial to the management of long-term CTS. Tendon and nerve gliding exercises alone are inferior to other modalities.&quot;</td>
</tr>
<tr>
<td>Gurcay 2009</td>
<td>3.5</td>
<td>N =32 female, housewife patients with clinically and EDS confirmed mild or moderate CTS. Mean age 40.8±11.2 years.</td>
<td>N =32 female, housewife patients with clinically and EDS confirmed mild or moderate CTS. Mean age 40.8±11.2 years.</td>
<td>Group A: local injection of 6mg betamethasone through 25-guage needle near distal wrist-flexion crease (n = 18) vs. Group B: meloxicam 15 mg/day, PO, for 3 weeks (n = 14). Both groups advised to wear wrist splints in neutral position at night for 3 weeks. Follow-up at 3 months.</td>
<td>No significant difference found between the groups for Functional Status Scale (FSS) scores, Jebsen Taylor Test (JTT) scores, or electrophysiological findings at 3 months (p&lt;0.05).</td>
<td>&quot;[T]he two treatment methods resulted in some functional gains in hand dexterity and improvement in electrophysiological data, but with neither of the methods demonstrating superiority.&quot;</td>
</tr>
<tr>
<td>Stransky 1989</td>
<td>3.5</td>
<td>N = 15 EDS confirmed</td>
<td>N = 15 EDS confirmed</td>
<td>200mg of Vitamin B6 vs. placebo</td>
<td>&quot;Significant changes in nerve conductions and EMGs did not occur when initial and follow-up data were compared. Clinical findings</td>
<td>&quot;Vitamin B6 seems to have no advantage over conservative therapy for carpal tunnel syndrome.&quot;</td>
</tr>
<tr>
<td><strong>No mention of sponsorship or COI.</strong></td>
<td><strong>did not correlate with electrodiagnostic findings.</strong></td>
<td><strong>Lidocaine Patches</strong></td>
<td><strong>Methocted. details sparse.</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>--------------------------------------------</td>
<td>---------------------------------</td>
<td>----------------------------------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Moghtaderi 2009</td>
<td>2.5</td>
<td>N = 65 with clinical and electrodiagnostic evidence of CTS. Aged 18-75 years.</td>
<td>Group 1 received ELMA cream (n = 30) vs. Group 2 received one injection of methylprednisolone acetate 40 mg at wrist (n = 35). Follow-up for 4 weeks.</td>
<td>Significant changes reported in pain in both groups, (p &lt; 0.001). Treatment-related adverse events (AEs) reported in 2 patients in group 1 (5.7%) and 10 patients in group 2 (28.5%).</td>
<td>&quot;ELMA cream was effective in reducing pain associated with CTS and well tolerated and it may offer patients with CTS an effective, noninvasive symptomatic treatment.&quot;</td>
<td></td>
</tr>
<tr>
<td>RCT</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No mention of sponsorship or COI.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jensen 2006</td>
<td>1.5</td>
<td>N = 40 with CTS. Age 18-75.</td>
<td>Lidocaine patch 5% daily (n = 20) vs. Lidocaine 1% single injection of 0.5mL plus methylprednisolone acetate 40mg at start of study (n = 20). Follow-up for 4 weeks.</td>
<td>Statistically significant decreases in 10 of 20 PQAS pain descriptor ratings occurred with both treatments, (p &lt;0.0025); 8 ratings showed no significant trends for decreasing before treatment to after treatment. No significant differences found between treatment conditions on any of the PQAS items.</td>
<td>&quot;The results support the validity of the PQAS items for assessing the effects of pain treatment on pain qualities of carpal tunnel syndrome.&quot;</td>
<td></td>
</tr>
<tr>
<td>RCT/Parallel-group/Open label</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sponsored by grant from Endo Pharmaceuticals, Inc (M.P.J.). M.P.J. and S.R.N. received research support and/or consulting fees. A.R.G., N.O. and B.S.G. hold stock options in Endo Phar.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Magnets**

<table>
<thead>
<tr>
<th><strong>Combination Magnetic Field Therapy</strong></th>
<th><strong>Methodological details sparse.</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Weintraub 2008</td>
<td>3.5</td>
</tr>
<tr>
<td>RCT</td>
<td></td>
</tr>
<tr>
<td>Sponsored by Nikken, Inc. No COI.</td>
<td></td>
</tr>
</tbody>
</table>

**Pulsed Magnetic Field Therapy**
<table>
<thead>
<tr>
<th>Study</th>
<th>Total N</th>
<th>Description and Treatment</th>
<th>Baseline comparison</th>
<th>Baseline comparability data</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arikan 2011 RCT</td>
<td>3.5</td>
<td>N = 57 hands from 38 patients with idiopathic CTS. Mean age: 48.8 years. Pulsed Magnetic Field Therapy 30 minutes/day for 3 weeks using BTL-09 device (n = 28 hands/19 patients) vs. Sham therapy same procedure without running device (n = 29 hands/19 patients). Assessment at baseline and end of treatment. Follow-up 1 month post-treatment.</td>
<td>No significant change was observed between groups for either clinical parameters or electrophyslogic studies (p &gt; 0.05).</td>
<td>&quot;We conclude that magnetic field and placebo magnetic field treatments in the patients with idiopathic carpal tunnel syndrome are effective to both clinical and electrophysiological endpoints in short term, but not superior to each other.&quot;</td>
<td>Baseline comparability data suggest randomization failure and possible quasi randomization &quot;every other&quot;.</td>
</tr>
<tr>
<td>Dakowicz 2011 RCT</td>
<td>2.5</td>
<td>N = 38 with diagnosed idiopathic CTS confirmed by ENG. Mean age 50.8±10.3 years. Low-level laser therapy (LLLT) using Ga-As Physioter D-50 for 5 minutes and 33 seconds (N = 18) vs. Pulsed magnetic therapy (PMF) with Magnetronic MF-10 for 15 minutes (N = 20). Two series of 10 sessions, with 2 week break between. Assessment after each series and at 6 months post-treatment.</td>
<td>No between-groups comparisons were made. In both groups, VAS improved after each series and at 6-months post-treatment (p &lt;0.05).</td>
<td>&quot;The presented study demonstrated that a clinical improvement in CTS patients was observed after LLL as well as PMF.&quot;</td>
<td>&quot;(W)e conclude that magnetic field and placebo magnetic field treatments in the patients with idiopathic carpal tunnel syndrome are effective to both clinical and electrophysiological endpoints in short term, but not superior to each other.&quot;</td>
</tr>
<tr>
<td>Bhatia 2000 RCT</td>
<td>3.5</td>
<td>N = 102 Plaster splint vs wool and crepe bandage. &quot;There were no reported problems with wound breakdown or other symptoms at the 2 week follow-up. Using the Mann-Whitney U test, there were no significant statistical differences in the [pain scores or number of tablets ingested up top 3 days postoperatively between the two groups.]&quot;</td>
<td>No between-groups comparisons were made. In both groups, VAS improved after each series and at 6-months post-treatment (p &lt;0.05).</td>
<td>&quot;The presented study demonstrated that a clinical improvement in CTS patients was observed after LLL as well as PMF.&quot;</td>
<td>&quot;This prospective randomized study has not supported the use of plaster. We believe that patients undergoing carpal tunnel release should be treated postoperatively with a bulky wool and crepe bandage.&quot; States single blinded, but unclear how blinding was done.</td>
</tr>
<tr>
<td>Horng 2011 RCT</td>
<td>3.5</td>
<td>N=60 patients with symptoms (pain, numbness within median nerve distribution, nocturnal pain), positive Phalen Group 1: paraffin therapy (in hospital 2x a week, administered by none-dip method at 55ºC) plus splint (custom made neutral volar wrist splint to be worn at night for at least 8 weeks) plus tendon gliding exercise Functional status difference before/after treatment (mean±SD): Group 1; 0.4±0.5 vs. Group 2; 0.1±0.5 vs. Group 3; -0.2±0.7 (p = 0.04). NS between groups for symptom severity score (p = 0.56), pain scale (p = 0.44), Disability of the Arm</td>
<td>No between-groups comparisons were made. In both groups, VAS improved after each series and at 6-months post-treatment (p &lt;0.05).</td>
<td>&quot;The presented study demonstrated that a clinical improvement in CTS patients was observed after LLL as well as PMF.&quot;</td>
<td>&quot;This prospective randomized study has not supported the use of plaster. We believe that patients undergoing carpal tunnel release should be treated postoperatively with a bulky wool and crepe bandage.&quot; States single blinded, but unclear how blinding was done.</td>
</tr>
</tbody>
</table>

**Splinting**

<table>
<thead>
<tr>
<th>Study</th>
<th>Total N</th>
<th>Description and Treatment</th>
<th>Baseline comparability data</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bhatia 2000 RCT</td>
<td>3.5</td>
<td>N = 102 Plaster splint vs wool and crepe bandage. &quot;There were no reported problems with wound breakdown or other symptoms at the 2 week follow-up. Using the Mann-Whitney U test, there were no significant statistical differences in the [pain scores or number of tablets ingested up top 3 days postoperatively between the two groups.]&quot;</td>
<td>No between-groups comparisons were made. In both groups, VAS improved after each series and at 6-months post-treatment (p &lt;0.05).</td>
<td>&quot;The presented study demonstrated that a clinical improvement in CTS patients was observed after LLL as well as PMF.&quot;</td>
</tr>
</tbody>
</table>

---

**Notes:**
- **RCT** indicates randomized controlled trials.
- **COI** indicates conflicts of interest.
- **N** represents the number of participants.

---

**Baseline comparability data:**
- Sparse baseline comparability data. At 6 months, both groups showed comparable (in)efficacy.
- States single blinded, but unclear how blinding was done.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>Participants</th>
<th>Interventions</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Koca 2014</td>
<td>RCT</td>
<td>N=75 patients with idiopathic CTS; presence of paresthesia, pain, and/or vasomotor symptoms of hand through distribution of median nerve for longer than 6 weeks; positive Phalen’s maneuver and/or Tinel’s sign and/or carpal compression test. Mean age Group I – 35.4±4.2 years, Group II – 34.2±5.2, Group III 34.9±4.8 years.</td>
<td>Group I: splint therapy, neutral position wrist splint with aluminum bar at night for 3 weeks (n = 25) vs. Group II: transcutaneous electrical stimulation, TENS on the carpal ligament and palmar area of hand at pulse rate of 100 Hz frequency and stimulation period of 80 ms, 20 minute sessions for 15 total sessions (n = 25) vs. interventional current, IFC therapy at base frequency of 4,000 Hz with a modulation frequency range of 20 Hz, electrodes placed on 1/3 mid portion of volar area of forearm, palmar area of hand, and thenar area of hand, 20 minute sessions for 15 sessions (n = 25). Assessments at baseline and 3 weeks after completion of treatment.</td>
<td>NS between TENS and splint therapy for improvement in clinical scores (p &gt;0.05). VAS (mean±SD) at 6 weeks: IFC 4.80±1.18 vs. splint 6.37±1.18 (p = 0.001); IFC vs. TENS 6.68±1.42 (p &lt;0.001). Median nerve motor distal latency (mMDL) mean±SD at 6 weeks: IFC 3.89±0.88 vs. splint 4.06±0.61 (p = 0.001); IFC vs. TENS 4.06±0.88 (p = 0.003). Median sensory nerve conduction velocity (mSNCV) mean±SD at 6 weeks: IFC 41.80±1.76 vs. splint 40.75±1.48 (p = 0.010); IFC vs. TENS 41.38±1.78 (p = 0.021). Symptom severity (mean±SD) at 6 weeks: IFC 1.90±1.21 vs. TENS 2.50±0.78 (p = 0.039).</td>
</tr>
</tbody>
</table>

**Sign or positive Tinel sign, and electrophysiologic evidence of CTS. Mean age 50.5±9.4 years.** three times daily holding each position for 7 seconds and then repeating the exercises 5 times per session (N=20) vs. Group 2: paraffin therapy plus splint plus nerve gliding exercise (N=20) vs. Group 3: paraffin therapy plus splint (N=20). Follow up 2 months after treatment. | Shoulder, and Hand (DASH) questionnaire (p = 0.29), World Health Organization Quality of Life Questionnaire Brief Version (WHOQOL-BREF) physical domain (p = 0.31), WHOQOL-BREF psychologic domain (p = 0.53), WHOQOL-BREF social domain (p = 0.88), and WHOQOL-BREF environmental domain (p = 0.45). | gliding exercises, paraffin therapy, and splinting. | 

**Koca 2014** RCT No mention of sponsorship or COI. |

Gurcay 2012 | RCT | N=54 female housewives with mild-to-moderate CTS diagnosed with clinical and | Group I: phonophoresis with 0.1% betamethasone applied over carpal tunnel at frequency 1 MHz and intensity 1W/cm² for 10 minutes. Group II: control in favor of phonophoresis (p = 0.012). NS between groups for grip strength. | Boston Symptom Severity Scale (BSSS): significant at 3 months, phonophoresis vs. control in favor of phonophoresis (p = 0.012). NS between groups for grip strength. | 

"We observed no added benefit or increased motor skills or hand dexterity in the groups after treatments." | Sparse baseline data and comparable efficacy. | 

**Gurcay 2012** RCT No mention of sponsorship or COI. |
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Intervention</th>
<th>Outcome</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sevim 2004</td>
<td>Prospective, randomized, blinded trial</td>
<td>120</td>
<td>Betamethasone injections 4cm proximal to the carpal tunnel vs. injections distal to carpal tunnel vs. just splinting vs. control</td>
<td>“Splinting provided symptomatic relief and improved sensory and motor nerve conduction velocities at the long-term follow-up when the splints were worn almost every night. Proximal and distal injections of steroids were ineffective on the basis of both clinical symptoms and electrophysiologic findings.”</td>
<td>Steroid injections may be beneficial short-term in mild and moderate CTS. However, splinting provided long term symptomatic relief and improved sensory and motor nerve conduction.</td>
</tr>
<tr>
<td>Straika 1998</td>
<td>RCT</td>
<td>120</td>
<td>Splint vs. Splint with energized high voltage pulse unit</td>
<td>“In the energized group, post-treatment evaluation showed statistically significant decreases in the amount of stimulation required to stimulate the median nerve and the amount of hand edema and pain. The energized group also had improved repetitive task times. None of these improvements occurred in the non-energized group.”</td>
<td>“HVPC appears to be an effective method for minimizing the severity of repetitive stress injuries of the wrist.”</td>
</tr>
<tr>
<td>Madjdinasab 2008</td>
<td>RCT</td>
<td>48</td>
<td>Splint group: neutral position splint at night and during the day if possible for 6 weeks (n = 24) vs. steroid group: oral prednisolone 20mg/day for 2 weeks (n = 24). Assessments at</td>
<td>No significant differences between groups for median nerve sensory, motor distal latency, and conduction velocity (p &gt;0.05).</td>
<td>“Both treatment methods (splint and oral steroids) are effective but they don’t have any significant difference between two methods after six weeks follow up.”</td>
</tr>
</tbody>
</table>

Electrophysiologic evidence. Mean age 43.7±8.4 years. Minute sessions, 3 days a week for 3 weeks (n = 18) vs. Group II: iontophoresis with 0.1% betamethasone, 2 mA for 10 minutes a day, 3 days a week for 3 weeks (n = 16) vs. Group III: wrist splint only, custom-made volar thermoplastic splint in neutral position worn at night only for 3 weeks (n = 18). Assessments at baseline, 3 months after treatment.

Strength (p = 0.280) and 9-hole peg test, NHPT (p = 0.811).

Steroid injections may be beneficial short-term in mild and moderate CTS. However, splinting provided long term symptomatic relief and improved sensory and motor nerve conduction.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Sample Size</th>
<th>Intervention</th>
<th>Comparator</th>
<th>Follow-up</th>
<th>Results</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dincer 2009</td>
<td>RCT</td>
<td>No mention of sponsorship. No COI.</td>
<td>2.5</td>
<td>N = 60 females with bilateral mild to moderate CTS diagnosis made by electromyography and clinical examination. Mean age: 34 years for Sp, 30 years for SpUS, 36 years for SpLLL group.</td>
<td>Splinting only (Sp), (N=40) vs. Splinting + Ultrasound therapy. A total of 10 US treatment sessions were performed once a day, 5x a week for 2 weeks (SpUS), (N=40) vs. Splinting plus low level laser therapy (SpLLL), (N=40). Follow up visits: in first month, and third month, after treatment. Patients were instructed to wear the splints at night for 3 mo. Ultrasound therapy was administered to each other for 3 min per session, on the area over carpal tunnel at a frequency of 3 MHz and an intensity of 1.0W/cm² in continuous mode with a transducer 5 cm² in size with gel.</td>
<td>After profile analysis (MANOVA), results showed that improvements in SpUS and SpLLL groups statistically significantly better than those seen in Sp group (p = 0.0429 and p = 0.0001). Also, difference between SpUS and SpLLL groups significant (p =0.03). Both SpUS and SpLLL groups had statistically significantly better improvement than Sp group at 3 months (p &lt;0.0001 for both groups) On the other hand, no significant differences between SpUS and SpLLL group profiles. VAS pain scores improved in all groups at 1 and 3 month vs. baseline. Both SpUS and SpLLL groups improvements significantly better than Sp group improvement over time (p = 0.0001 for both). SpLLL group showed significantly better improvement than SpUS group.</td>
<td>&quot;In conclusion, the results of this study demonstrate the effectiveness of conservative treatments for mild to moderate CTS. Combining US or LLL therapy with splinting appeared to be more effective than splinting alone in our study. However, the combination of LLL therapy with splinting appeared to be superior to splinting plus US, especially for improvements in symptom severity, pain alleviation, and patient satisfaction. Further research with larger patient samples and longer follow-up periods are required to independently confirm our findings, and to determine the most effective doses and protocols for LLL and US therapies.&quot;</td>
<td></td>
</tr>
<tr>
<td>Burke 1994</td>
<td>RCT</td>
<td>1.5</td>
<td>N = 59</td>
<td>Splints vs. optimal angle</td>
<td>Randomization unclear, study states blinded but that seems unlikely.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pinar 2005</td>
<td>RCT</td>
<td>No mention of sponsorship or COI.</td>
<td>1.5</td>
<td>N = 26 females with NCS positive CTS</td>
<td>Grip strength TGE vs. splint plus reduced use (pre/post): 17.8±6.1/22.0±6.8 vs. 20.4±4.7/21.7±4.3 (p &lt;0.05) between groups. Most results negative.</td>
<td>&quot;Significant progress was detected in both control and experimental groups during the posttreatment phase compared with the initial phase (P&lt;0.05). However, when the 2 groups were compared, the experimental group in which nerve gliding exercises were added to conservative therapy</td>
<td>Low sample size. Blinding unclear. Diagnostic criteria unclear, including NCS and 9 other criteria that seem unlikely fulfilled for all. No non-treatment comparison. No between group differences. Conclusion for ultrasound not clearly supported. If bilateral</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Country</td>
<td>Method</td>
<td>Outcome</td>
<td>Findings</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>---</td>
<td>---------</td>
<td>--------</td>
<td>---------</td>
<td>----------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Khosrawi 2012</td>
<td>72</td>
<td>Iran</td>
<td>RCT</td>
<td>Acupuncture vs Control</td>
<td>Approaches demonstrated more rapid pain reduction; these patients also showed greater functional improvement, especially in grip strength (P&lt;0.05). CTS (12/30) treated the same and double-counted in results, weakening conclusions.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ho 2014</td>
<td>26</td>
<td>Taiwan</td>
<td>RCT</td>
<td>Electroacupuncture vs Acupuncture</td>
<td>Symptom severity scores baseline vs. week follow-up. Electroacupuncture decreased significantly (p = 0.02). Shortened median sensory latency, baseline to 4 weeks. Electro-Acu not statistically significant. Median Nerve F wave mean latency, baseline to 4 weeks. Grip strength baseline to 4 weeks post treatment. Acu Grip strength to 4 weeks post treatment, Acu 23.3±9.85 to 27.88±12.44 (p = 0.01) vs Electro-Acu no significant difference. “Despite the limitations in this study, we found that safety depth acupuncture and electroacupuncture could exert different positive therapeutic effects for patients with CTS. As evidenced by the improvement of Symptomology using electroacupuncture and improvements of grip strength, electrophysiological findings, and physical provocation sign of using acupuncture, the findings of this study provide references in clinical decision making when selecting proper treatment programs for symptomatic CTS patients.”</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Acupuncture Group underwent treatment in eight sessions of 60 minute duration over 4 weeks and also night splinting (n = 32). Control group was given vitamins B1, B6 and sham acupuncture. Also had night splinting (n = 32). Follow-Up at baseline 2 weeks and 4 weeks. Global Symptom Score (GSS) acupuncture group vs, control group at Week 4: 14.6±5.4 (p <0.001) vs 22.5±8.9 (p = 0.17). Nerve Conduction Velocity at 4 weeks, Acupuncture vs control: 37.6±8.3 vs 33.2±5.9 (p <0.02) |

Methodological details sparse.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>N</th>
<th>Design</th>
<th>Description</th>
<th>Outcomes</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cai 2009</td>
<td>RCT</td>
<td>0.5</td>
<td>98 cases of CTS all history of strain or traumatic injury of wrist joint. Mean age: Warm Needling Group Range 32-67 years; Control Group 35-71 Years old</td>
<td>Acupuncture group: warm needling techniques and Tuina relaxing manipulations. 10 30 minute sessions (n = 60) vs. Control Group Given block therapy with 10mg Triamcinolone A and lidocaine once every 3-5 days, Dibazol and Vitamin B1 were taken orally 3 times daily until trial over. (n = 38). Follow-up only mentioned only after 1 course of treatment. (no specific time frame)</td>
<td>Clinically cured (clinical symptoms disappeared, movement restored, negative in Carpal canal irritating test. Acu group 49 (81.67%) vs control 18 (47.37%) (p&lt;0.01).</td>
<td><em>Acupuncture plus Tuina manipulation is a simple therapy for carpal tunnel syndrome, but with remarkable therapeutic effects.</em></td>
</tr>
<tr>
<td>Stasinopoulos 2005</td>
<td>RCT</td>
<td>3.0</td>
<td>25 with unilateral idiopathic carpal tunnel syndrome, mild to moderate nocturnal pain, and paraesthesia lasting &gt;3 months. The mean age was 47.4 years.</td>
<td>Polarized polychromatic noncoherent light or Bioptron light administered perpendicular to carpal tunnel area for 6 minutes at operating distance 5-10cm from carpal tunnel area, 3x weekly for 4 and 6 weeks (n = 25). Outcome measures used were participants’ global assessments of nocturnal pain and paraesthesia, respectively, at 4 weeks and 6 months. Follow-up at 4 weeks, and 6 months.</td>
<td>At 4 weeks, 2 (8%) had no change in nocturnal pain, 6 (24%) were in slightly less nocturnal pain, 12 (48%) were much better in regard to nocturnal pain and 5 (20%) were pain-free. At 6 months, 3 patients (12%) were slightly better in regard to nocturnal pain, 13 (52%) were much better regarding nocturnal pain, and 9 patients (36%) were pain-free.</td>
<td><em>Nocturnal pain and paraesthesia associated with idiopathic carpal tunnel syndrome improved during polarized polychromatic noncoherent light (Bioptron light) treatment.</em></td>
</tr>
<tr>
<td>Pratelli 2015</td>
<td>RCT</td>
<td>2.5</td>
<td>70 symptomatic hands clinically diagnosed and electromyographically proven CTS. Mean age 54.2 (38-74)</td>
<td>Group 1 (n = 35) treated with Fascial manipulation (FM) 45 minute session 3x a week for 3 weeks vs. Group 2 (n = 35) Low Level Laser Therapy 5x a day for 10 minute sessions. Follow-up 10 days before treatment, BTCQ symptomatic and functional and Visual Analogue Scale scores baseline – first follow up, group 1: 3.027, 3.097, and 6.00 vs 1.362, 1.40, 0.80 (p &lt;0.0001). Baseline vs second follow up 3.027, 3.097, and 6.00 vs 1.27, 1.31, 0.714 (p &lt;0.0001). Group 2 BTCQ</td>
<td>*FM appears to be an appropriate treatment not only for musculoskeletal dysfunction but also for common nerve entrapments as in carpal tunnel syndrome. The method is effective and non-invasive. It gives excellent results for the</td>
<td></td>
</tr>
</tbody>
</table>

**Low-level Laser Therapy**

**Manipulation and Mobilization**
<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>Participants</th>
<th>Interventions</th>
<th>Outcome Measures</th>
<th>Results/Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heebner 2008 RCT</td>
<td>2.0</td>
<td>N = 61 with CTS confirmed using Nerve Conduction Velocity testing; mean age 52 (32-75)</td>
<td>Group 1 (n = 28) standard care provided by hospital (night splinting, tendon gliding exercises) vs. Group 2 (n = 32) same as group 1 but addition of neurodynamic mobilization exercise median nerve bias. Follow-up baseline, 1 and 6 months after initial treatment.</td>
<td>Symptomatic and functional as well as Visual Analogue Scale baseline vs follow up 1: 3.52, 2.90, 5.51 vs 2.66, 2.58, 5.00 (p &lt;0.001). Worsening of symptoms in group 2 from follow up 1 to 2. Relief of local symptoms and for restoring functionality with benefits that remain at three month follow up.</td>
<td>The results of this study suggest that persons with CTS in a community hospital do not benefit from a one-time nonsurgical intervention that includes splinting instruction and standard tendon-gliding exercises alone or splinting and tendon gliding along with neural mobilization exercises.</td>
</tr>
<tr>
<td>Bialosky 2011 RCT</td>
<td>1.5</td>
<td>N = 40 females; mean age for individuals with CTS: 40.75±10.38, 38.25 ± 12.32 for healthy individuals.</td>
<td>Group 1 (n = 20) with clinically diagnosed CTS (Tinel’s, Phalen’s, Carpal Compression Tests) vs. Group 2 (n = 20) age matched and no sign of CTS. Follow-up 2x a week for 3 weeks.</td>
<td>No statistically significant change in outcome measures associated with Neurodynamic Intervention. Baseline relationship between clinical pain and pain sensitivity w/ signs and symptoms of CTS: MP flexor retinaculum after sensation 0.88 (p &lt;0.01). Change in usual pain over 3 weeks in MP flexor retinaculum after temporal summation -0.57 (p = 0.05) and after sensation -0.55 (p = 0.01).</td>
<td>Participants with signs and symptoms of CTS differed from healthy age- and sex-matched controls in suprathreshold measures of pain sensitivity suggesting a central mechanism of pain. Immediate change in mechanical pain sensitivity and after sensation and 3-week change in temporal summation were associated with improvements in clinical pain intensity suggesting prognostic factors and a potential mechanism for improvement respectively.</td>
</tr>
<tr>
<td>Moraska 2008 RCT</td>
<td>3.5</td>
<td>N = 27 with CTS for at least 6 months.</td>
<td>General massage (GM, n = 13) focused on reducing muscular tension and enhancing circulation to back, neck, and both upper extremities v. targeted massage (TM, n = 14)</td>
<td>Grip strength: TM showed significantly greater strength increase compared to GM, p=0.04.; improvement for TM first seen after 7th massage and maintained following 11th massage and for at least4 weeks.</td>
<td>The results from this study suggest that massage therapy may be a useful part of a conservative care treatment regimen, although additional research support is needed.</td>
</tr>
</tbody>
</table>

**Methodological details** sparse. High dropout rate.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>Sponsorship</th>
<th>COI</th>
<th>n</th>
<th>Age</th>
<th>Intervention</th>
<th>Outcome Measures</th>
<th>Key Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moraska 2010 RCT</td>
<td>3.5</td>
<td>Same as Moraska 2008</td>
<td>Same as above (Moraska 2008)</td>
<td>Same as above (Moraska 2008)</td>
<td>30 minute structured massage treatments over 6 weeks.</td>
<td>weeks after last treatment, p&lt;0.01 for all time points.</td>
<td>Therapeutic Touch</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blankfield 2001 RCT</td>
<td>1.5</td>
<td>N = 21 with electro diagnostically confirmed CTS. Mean age 57.4 for therapeutic touch treatment group, 55.2 for sham treatment.</td>
<td>Therapeutic touch (TT) group (n = 11) vs. sham (n = 10), 1x a week for 6 weeks. Follow-up period not mentioned.</td>
<td>Mean motor distal latencies (baseline/follow-up): TT (5.4±0.9/5.2±1.1ms) vs. sham (6.1±1.8/5.9±1.0msec), p &gt;0.15. Pain/relaxation scores NS.</td>
<td>“[TT] was no better than placebo in influencing median motor nerve distal latencies, pain scores, and relaxation scores.”</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blackfield 2001 RCT</td>
<td>3.5</td>
<td>N = 46 with CTS. Mean age: group 1: 45.20 years, group 2: 43.31 years; group 3: 44.53 years. First group received 0 W/cm² ultrasound treatment (placebo) (n = 15) Vs. second received 1.0 W/cm² continuous ultrasound treatment (n = 16) Vs. third received 1.0 W/cm² 1:4 pulsed ultrasound treatment (n = 15). Administered for 5 days a week for a total of 15 sessions. All patients also wore night splints during treatment period. Follow up: not mentioned.</td>
<td>Significant improvements in all groups as per post-treatment Functional Status Scale score (p&lt;0.05 all groups), Symptom Severity Scale score (first group: p&lt;0.05, second group: p&lt;0.01, third group: p&lt;0.001) and Visual Analogue Scale score (first and third groups: P&lt;0.01, second group: p&lt;0.001). Sensory conduction velocities improved in 2nd and 3rd groups (p&lt;0.01). Distal latency in 2nd finger showed improvement only in 3rd group (p&lt;0.01) and action potential latency in palm improved only in 2nd group (p&lt;0.05).</td>
<td>“The results of this study suggest that splinting therapy combined with placebo and pulsed or continuous ultrasound have similar effects on clinical improvement. Patients treated with continuous and pulsed ultrasound showed electrophysiological improvement; however, the results were not superior to those of the placebo.”</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oztas 1998 RCT</td>
<td>2.5</td>
<td>N = 18 females with CTS in 30 hands. Mean age: Group A: 53.2 years; Group B: 51.3 years. Group A: continuous ultrasound therapy with intensity of 1.5 W/cm² (n = 10) Vs. Group B: US therapy with intensity of 0.8 w/cm² (n = 10) Vs. Group C: US therapy</td>
<td>Night pain/paresthesia before treatment/after treatment: Group A: 2.30±.68/1.40±.52; Group B: 2.60±.70/1.70±.82; Group C: 2.60±.69/1.40±.97. Mean distal latency: Group A: 5.85±1.87/6.00±1.95; Group B: 6.40±1.70/6.10±1.82.</td>
<td>“Ultrasound therapy in CTS was comparable to placebo ultrasound in providing symptomatic relief, and the probability of a negative effect on motor Single blind (patient). Suggests ultrasound not effective. Small sample size of 18 women. Methodological details sparse.”</td>
<td>Ultrasound vs. Placebo</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Copyright © 2016 Reed Group, Ltd.
<table>
<thead>
<tr>
<th>Study</th>
<th>Methodology</th>
<th>Sponsorship</th>
<th>Participants</th>
<th>Intervention</th>
<th>Follow-up</th>
<th>Outcomes</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duymaz 2012 RCT</td>
<td>RCT</td>
<td>No sponsorship or COI</td>
<td>3.5</td>
<td>N = 58 unemployed with CTS confirmed by provocation tests and EMG and symptoms of numbness, tingling, weakness, and pain in hands for at least 3 months but not more than 1 year. Mean age 51.85±7.29 years.</td>
<td>Group I: iontophoresis with dexamethasone 0.4% at a current 2 mA for 20 minutes (n = 20) vs. Group S: iontophoresis sham using water at current 2 mA for 20 minutes (n = 18) vs. Group U: underwater ultrasound 5 minutes per session using direct current at an intensity of 0.8 W/cm², 3 applications once a day 5x a week for 3 weeks (n = 20). All received: training on performing tendon and nerve gliding exercises to be completed for 3 sets of 10 everyday; ergonomic training for daily living activities; and neutral wrist splinting at night. Follow-up after 3 months.</td>
<td>Mean±SD VAS on movement difference between pre and post treatment values Group I vs. Group S vs. Group U: 2.75±1.71 vs. 0.66±1.13 vs. 1.30±1.83 (p&lt;0.001). Mean±SD VAS at rest difference between pre and post treatment values Group I vs. Group S vs. Group U: 2.55±1.76 vs. 0.50±0.78 vs. 1.20±1.73 (p &lt;0.001).</td>
<td>&quot;Our study results suggest that dexamethasone iontophoresis administration combined with tendon gliding exercises, splint and activity modification is reliable and effective in the treatment of patients with mild CTS.&quot;</td>
</tr>
<tr>
<td>Dincer 2009 RCT</td>
<td>RCT</td>
<td>No mention of sponsorship. No COI</td>
<td>2.5</td>
<td>N = 60 females with bilateral mild to moderate CTS diagnosis made by electromyography and clinical examination. Mean age 34 years for Sp, 30 years for SpUS, 36 years for SpLLL group.</td>
<td>Splinting only (Sp), (n = 40) vs. splinting + Ultrasound therapy, Total 10 US treatment sessions performed once a day, 5x a week, for 2 weeks (SpUS), (n = 40) vs. Splinting plus low level laser therapy (SpLLL), (n = 40) Follow up: 1st and 3rd month after treatment. Patients to wear splints at night for 3 months. Ultrasound therapy administered to each other for 3 minutes per session on area over</td>
<td>After profile analysis (MANOVA), results showed improvements in SpUS and SpLLL groups were statistically significantly better than those in Sp group (p = 0.0429 and p = 0.0001, respectively). Also, difference between SpUS and SpLLL groups significant (p = 0.03). Both SpUS and SpLLL groups had statistically significantly better improvement than Sp group at 3 months (p &lt;0.0001 for both groups) On other hand, no significant differences between SpUS and SpLLL.</td>
<td>In conclusion, the results of this study demonstrate the effectiveness of conservative treatments for mild to moderate CTS. Combining US or LLL therapy with splinting appeared to be more effective than splinting alone in our study. However, the combination of LLL therapy with splinting appeared to be superior to splinting plus US, especially for improvements in symptom severity, pain alleviation,</td>
</tr>
</tbody>
</table>

Methodological details sparse.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N (Hands)</th>
<th>Intervention</th>
<th>Primary Outcomes</th>
<th>Comparator</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aygul 2005 RCT</td>
<td>3.5</td>
<td>N = 31 (56 hands)</td>
<td>Local steroid injection 1 ml dexamethasone sodium phosphate vs. iontophoresis treatment with 1-4 mA galvanic current and mixture of 0.1% dexamethasone sodium phosphate vs. phonophoresis frequency of 3 MHz and intensity of 1.0 W/cm², with transducer of 5 cm², including mixture of 0.1% dexamethasone sodium phosphate</td>
<td>Injection group had a steady significant improvement for all parameters except SNAP&lt;sub&gt;a&lt;/sub&gt;, mTLI, and mMNCV at the first follow-up visit. Iontophoresis had significant improvements in the D4M-D4U and mTLI. Phonophoresis group had significant improvement of D4D-D4U and mMDL found 2 months after treatment.</td>
<td>“Steroid injection in CTS is more effective than iontophoresis and phonophoresis treatment in the short- to medium-term in patients with mild to moderate idiopathic CTS, and that the most sensitive neurophysiologic parameters at follow-up were D4D-D4U and D2M-DSU, which are objective parameters indicating the outcome of CTS treatment.”</td>
<td>Random in abstract, but nowhere in methods.</td>
</tr>
<tr>
<td>Gurcay 2012 RCT</td>
<td>3.5</td>
<td>N = 52 with CTS analyzed based on clinical and electrophysiological criteria. Mean age 43.7 ± 8.4 (range 24-57) years.</td>
<td>Group I, phonophoresis, 0.1% betamethasone applied over area of CT at frequency of 1 MHz and an intensity of 1 W/cm², plus wrist splint (n = 18) vs. Group II, 0.1% betamethasone iontophoresis from positive electrode at dosage of 2 mA for 10 minutes/day, plus wrist splint (n = 18) vs. Group III or control, instructed to use wrist splint alone (n = 18). Follow-up at 3 months.</td>
<td>At 3 months (T1), Boston Symptom Severity Scale (BSSS) improved in group I (p &lt; 0.001), group II (p = 0.001), and group III (p &lt; 0.001) compared to baseline (T0). Grip strength, and nine-hole peg test (NHPT) in all groups; I, II and III at 3 month vs baseline improved, (p &gt; 0.05). A significant difference between groups for BSSS, F = 4.599, (p = 0.015). No statistical difference between groups for grip strength at T0 and T1, X² = 2.546, (p = 0.280).</td>
<td>“Symptom severity improved in all groups after treatment, but no superiority was determined among the treatment groups with respect to motor skills and hand dexterity.”</td>
<td>Sparse baseline data and comparable efficacy.</td>
</tr>
</tbody>
</table>

### Carpal Tunnel Injections

<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N (Hands)</th>
<th>Intervention</th>
<th>Primary Outcomes</th>
<th>Comparator</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aygul 2005</td>
<td>3.5</td>
<td>N = 31 (56 hands)</td>
<td>Local steroid injection 1 ml dexamethasone</td>
<td>Injection group had a steady significant improvement for all</td>
<td>“Steroid injection in CTS is more effective than”</td>
<td>Random in abstract, but nowhere in methods.</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Inclusion Criteria</td>
<td>Interventions</td>
<td>Outcomes/Findings</td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------</td>
<td>-------</td>
<td>------------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gurcay 2012</td>
<td>52</td>
<td>Clinical and electrophysiological criteria. Mean age 43.7 ± 8.4 (range 24–57) years.</td>
<td>Group I: phonophoresis, 0.1% betamethasone applied over CT at frequency 1 MHz and intensity 1 W/cm², plus wrist splint (n = 18) vs. Group II: 0.1% betamethasone iontophoresis from positive electrode at 2 mA for 10 minutes/day, plus wrist splint (n = 18) vs. Group III or control, instructed to use wrist splint alone (n = 18). Follow-up at 3 months.</td>
<td>At 3 months (T1), Boston Symptom Severity Scale (BSSS) improved in group I (p &lt;0.001), group II (p = 0.001), group III (p &lt;0.001) vs. baseline (T0). Grip strength, and 9-hole peg test (NHPT) in groups; I, II and III at 3 month vs. baseline improved (p &gt;0.05). Significant difference between groups for BSSS, F = 4.599, (p = 0.015). No statistical difference between groups for grip strength at T0 and T1, X² = 2.546, (p = 0.280). <em>Symptom severity improved in all groups after treatment, but no superiority was determined among the treatment groups with respect to motor skills and hand dexterity.</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Seok 2013</td>
<td>36</td>
<td>CTS with positive Tinel sign or Phalen test, and numbness and tingling at least two of first, second and third digit. At least 19 years of age.</td>
<td>Extracorporeal shock wave therapy or ESWT group one session with 1000 shocks at a frequency of 360 shocks per minute (n = 15) vs. Local corticosteroid or CS injection group received 1 milliliter of triamcinolone acetonide 40mg (n = 16). Follow-up at 3 months.</td>
<td>VAS score of 7.06±1.89 in ESWT group vs 6.87±1.26 in CS injection group. VAS score improvement at 1 month/3 months; 4.56±0.81 in ESWT vs 4.13±1.50 CS group/4.18±1.05 vs 3.31±1.82. Symptom severity score at 1 month; 20.13±6.24, (p &lt;0.05) and at 3 months; 19.73±4.48 vs 18.25±3.71 CS group, (p &lt;0.05). Significant difference between ESWT and CS groups found only at median sensory distal latency 1 months after treatment. <em>ESWT can be as useful as CS injection for relieving symptoms of carpal tunnel syndrome.</em></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**RCT**

No mention of sponsorship or COI.

**Seok 2013**

No mention of sponsorship. No COI.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>N</th>
<th>Title</th>
<th>Table</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sevim 2004</td>
<td>3.0</td>
<td>Prospective randomized and blinded trials</td>
<td>N = 120 EDX confirmed</td>
<td>Betamethasone injections 4cm proximal to carpal tunnel vs. injections distal to carpal tunnel vs. just splinting vs control</td>
<td>“Splinting provided symptomatic relief and improved sensory and motor nerve conduction velocities at the long-term follow-up when the splints were worn almost every night. Proximal and distal injections of steroids were ineffective on the basis of both clinical symptoms and electrophysiologic findings.”</td>
<td>Steroid injections may be beneficial short-term in mild and moderate CTS. However, splinting provided long term symptomatic relief and improved sensory and motor nerve conduction.</td>
</tr>
<tr>
<td>Kamanli 2011</td>
<td>2.5</td>
<td>RCT</td>
<td>N = 19 with bilateral CTS. Mean age for groups PIG and DIG: 42±10 and 52±13.</td>
<td>Proximal approach steroid injection group (PIG), with triamcinolone aetonide 20 mg (n = 10) vs. Distal approach steroid injection group (DIG), with triamcinolone aetonide 20mg (n = 9). Follow-up for 3 months.</td>
<td>BCTS / VAS-pain (0-10) / and HAQ at baseline and 3 months: 66.7 ± 12 and 31.6 ± 8.2 at 3 months vs 51.7 ± 14.7 and 34.9 ± 16 in DIG group/8.5 ±1.1 and 3.3 ± 2 vs 8.3 ± 1.7 and 3.9 ± 22 / and 0.97 ± 0.38 and 0.43 ± 0.22 vs 0.63 ± 0.53 and 0.29 ± 0.14.</td>
<td>“[S]teroid injection from distal approach (palmar) into the carpal tunnel on patients with CTS is very comfortable, easy, effective and alternative.”</td>
</tr>
<tr>
<td>Stepić 2008</td>
<td>1.5</td>
<td>RCT</td>
<td>N = 40 with CTS. The mean age of 51.6 years.</td>
<td>Group 1: surgical decompression of median nerve by open release of carpal tunnel (n = 20) vs. Group 1: perineural injection 1ml betamethason immediately after surgical decompression (n = 20). Follow-up 7, 30, and 90 days.</td>
<td>90 days after surgical procedure, both groups showed statistically significant better results in second group (t = -2.116; p = 0.043). Final measurements did not show statistically important difference between treatment methods applied, SCS1 = 45.347 msec in first group vs SCS2 = 47.673 msec in second group.</td>
<td>“Intraoperative application of the corticosteroid injection during the surgical decompression results in faster regaining of conduction speed of the median nerve.”</td>
</tr>
<tr>
<td>Worség 1996</td>
<td>4.0</td>
<td>Two consecutive case series</td>
<td>N = 126 EDS confirmed</td>
<td>64 surgeries treated endoscopically vs. 62 surgeries by open release of carpal ligament.</td>
<td>“No significant differences between the groups were obtained regarding postoperative symptom severity.”</td>
<td>“The new device provides a reliable tool for single portal carpal tunnel release, although the risk of inadvertent damage to the neurovascular structures always remains a possibility with the endoscopic carpal tunnel technique.”</td>
</tr>
<tr>
<td>Study</td>
<td>Method</td>
<td>N</td>
<td>Treatment</td>
<td>Outcome</td>
<td>Comment</td>
<td></td>
</tr>
<tr>
<td>--------------</td>
<td>-------------</td>
<td>------</td>
<td>--------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Demirci 2002</td>
<td>RCT</td>
<td>4.0</td>
<td>Intracarpal betamethasone 6.4mg injections at Weeks 0 and 2 vs. open CTR</td>
<td>Boston questionnaire symptoms scale (0/3/6 months): open (3.4±0.7/1.3±0.3/1.3±0.3) vs. steroid (3.3±0.7/1.5±0.5/1.7±0.8). NS months 0-3 and p = 0.003 at 6 months.</td>
<td>Conservative steroid treatment provides short term improvement compared to surgery. It is relatively easy to apply, has a lower cost and comparable results short term and may be considered before surgery is done.</td>
<td></td>
</tr>
<tr>
<td>Nitz 1989RCT</td>
<td>3.5</td>
<td>N = 60</td>
<td>Open surgery vs. surgery with tourniquet</td>
<td>“Three weeks after the operation 77% of the patients in the tourniquet group had denervation in other than thenar muscles. Only one patient in the control (no tourniquet) group had similar electromyographic abnormalities after surgery. Tourniquet time and pressure did not vary significantly between those patients with or without postoperative forearm denervation. Mean operative time for the tourniquet and control groups was nearly identical.”</td>
<td>“These findings indicate that upper extremity tourniquet application results in subclinical, temporary changes in the muscles of the forearm, probably on the basis of nerve changes and denervation.”</td>
<td></td>
</tr>
<tr>
<td>Brüser 1999</td>
<td>RCT</td>
<td>3.5</td>
<td>N = 80 with CTS</td>
<td>Short (2.5cm) vs. long (4.5cm) incision</td>
<td>“The long incision resulted in a significant 10% loss of strength only at week three, otherwise no significant difference was found between the results of the two groups.”</td>
<td></td>
</tr>
<tr>
<td>Mackenzie 2000</td>
<td>RCT</td>
<td>3.5</td>
<td>Open surgery vs. endoscopic methods</td>
<td>Grip strengths (baseline/weeks 1/2/4): endoscopic (43/29/42/44) vs. open (39/21/29/30) (p &lt;0.01 at 2 and 4 weeks).</td>
<td>“Endoscopic carpal tunnel release provides faster recovery of strength than short-incision open carpal tunnel release and improves early postoperative comfort and function to a small degree.”</td>
<td></td>
</tr>
<tr>
<td>Borisch 2003</td>
<td>RCT</td>
<td>3.5</td>
<td>N = 273 EDS confirmed</td>
<td>Open CTR with vs. without epineurotomy.</td>
<td>“Study showed no significant difference in the recovery of sensory conduction velocity and distal motor latency after open decompression of the DROP OUT RATES WERE HIGH.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Methodology</td>
<td>Score</td>
<td>Description</td>
<td>Findings</td>
<td>Comments</td>
<td></td>
</tr>
<tr>
<td>------------</td>
<td>-------------</td>
<td>-------</td>
<td>------------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Finsen 1999</td>
<td>RCT</td>
<td>3.5</td>
<td>N = 74 (82 wrists) on whom open carpal tunnel release performed. Mean age 48 in mobilized group; 51 in immobilized group.</td>
<td>Mobilized Group: light dressing and told to move wrist and fingers as comfort allowed. (n = 45) vs. Immobilized Group: well-padded plaster of Paris splint with wrist in slight dorsiflexion. (n = 37). Follow-up at 2 and 6 weeks, and 6 months. Post-op VAS pain scores indicated patients in both groups benefited from post-op treatments. But no significant differences in mean VAS pain score at any time point for mobilized vs. immobilized; Pre-op 56mm vs. 51mm; 2 weeks 6mm vs. 5mm; 6 weeks 6mm vs. 2mm; 6 months 3mm vs. 2mm.</td>
<td>“Thus immobilization confers no advantage with regard to regress of the original complaints postoperatively. Nor did immobilization reduce the frequency of common complications, such as scar or pillar pain.” No advantage to splinting after carpal tunnel release surgery.</td>
<td></td>
</tr>
<tr>
<td>Hansen 2009</td>
<td>RCT</td>
<td>3.5</td>
<td>N= 47 (54 hands) diagnosed with idiopathic CTS who required release of carpal tunnel. Mean age 48 years.</td>
<td>Novafil Group: Interrupted non-absorbable sutures. 5/0 monofilament polybutester (n = 26 hands) vs. Caprosyn Group-Continuous absorbable subcuticular 4/0 monofilament polyglytone sutures. (n = 28 hands). Follow-up assessed daily in patient’s journal until sutures removed 10-14 days after surgery. Pain monitored through this period. Cosmetic appearance measured at 3 months. VAS pain score significantly lower in Caprosyn group vs. Novafil group at post-op day 1 (p = 0.04) and post-op day 2 (p = 0.02). However, difference in VAS pain score not significant at any other time point. Caprosyn group showed better cosmetic result with 25/28 hands showing nice appearance when being evaluated by surgeon vs. 18/26 in Novafil group. However, this difference not significant (p = 0.14).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cellocco 2005</td>
<td>RCT</td>
<td>3.5</td>
<td>N = 185 affected by mild to moderate median nerve compression. 222 carpal tunnel release procedures</td>
<td>Group A: Mini-open blind technique using Knifelight (n = 82, 99 procedures) Vs. Group B- limited open technique (n = 103, 123 procedures). Follow-up at 19 and 30 months following surgery. Group A returned to work significantly quicker in mean days than group B; 16.6 days vs. 25.4 days (p &lt;0.001). Mean score for first section of Boston Carpal Tunnel Questionnaire (BCTi) significant at 19 months for group A vs. B: 1.46 vs. 2.04.</td>
<td>“Our study suggests that the mini-open blind CT release can be a safe procedure, even when performed using a small transverse wrist incision.” Short follow up period (19 months) favored transverse procedure, but at 30 months the differences between groups decreased. Group A experienced shorter recovery rate and less pain &amp; numbness.</td>
<td></td>
</tr>
</tbody>
</table>

Copyright© 2016 Reed Group, Ltd.
<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>RCT</th>
<th>COI</th>
<th>Patients</th>
<th>Methodology</th>
<th>Results</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heidarian 2013</td>
<td>3.5</td>
<td>No sponsorship or COI.</td>
<td>N = 59 with indication for carpal tunnel release. Mean age 47.6 years.</td>
<td>Open Group: Open carpal tunnel release surgery (n = 30) vs. Knifelight Group: (n = 29). Follow-up immediately after surgery and 3 weeks and 6 months.</td>
<td>Knifelight group vs. open group showed significantly shorter operation time: 8.5 min vs. 21 min (p &lt;0.001), significantly shorter mean scar length (mm): 14.8mm vs. 40.7mm (p &lt;0.001). Knifelight also significantly quicker return to daily activity vs. Open; 34.4 days vs. 51.9 days (p = 0.015). VAS pain score at 3 weeks not significant between groups (p = 0.24).</td>
<td>“In conclusion according to the results of this study, compared to the open release method, Knifelight technique could significantly decrease the mean duration of surgery, incision length and time to return to work.”</td>
<td>Sparse methodological details. Short follow-up time (3 weeks). Knifelight “appears” to decrease surgical time, scar length and time to resume normal activities, but pain ratings for both groups were comparable.</td>
</tr>
<tr>
<td>Ucar 2012</td>
<td>3.0</td>
<td>No mention of sponsorship or COI.</td>
<td>N = 90 with CTS syndrome. Mean age 46.75 years.</td>
<td>G1 Group- Distal approach. A 2 cm vertical incision on the ulnar side of the thenar crease beginning at the distal wrist crease. (n = 45) Vs. G2 Group- Proximal approach. A 2 cm vertical incision made on the ulnar side of the palmaris longus tendon, beginning proximal to wrist crease. (n = 45) Follow-up at one month. Final follow up mean 30.4 months in G1 and 31.0 months in G2.</td>
<td>Boston Carpal Tunnel questionnaire scores used for assessment. Both groups increased significantly in Symptom and Functional scales from baseline from pre-op to 1 month follow up (p &lt;0.001) and from 1 month follow-up to final follow-up (p &lt;0.001). Functional and Symptom scores not significant between groups at any follow-up period (p &gt;0.05). G2 showed significantly shorter mean operation time vs. G1; 10.7 min vs. 18.6 min (p &lt;0.001). G2 also significantly less scar tissue pain vs. G1; 6.7% vs. 24.4% (p = 0.02).</td>
<td>“Finally, the absence of relapse and good clinical results make both surgical techniques used in this study suitable. For this reason, we consider that the selection of the mini-surgical technique used should depend on the experience and skill of the surgeon.”</td>
<td>Sparse methodology and short follow-up time (1 month). Mean surgical time and scar tissue pain were less in group 2 (the 2 cm proximal incision group).</td>
</tr>
<tr>
<td>Kang 2008</td>
<td>3.0</td>
<td>No sponsorship or COI.</td>
<td>N = 72 with diagnosed CTS. Mean age 34.8 years.</td>
<td>Arthroscopic Excision Group- 2 stab incisions at standard 3-4 and 4-5 portal sites (n = 41) vs. Open Excision Group- Transverse skin incision 2 to 3cm in length (n = 31). First follow-up 5-7</td>
<td>Main outcome ganglion recurrence. At 2nd follow-up, arthroscopic group 1 ganglion recurrence vs. 0 in open group (p = 0.381). Not significant at final follow up. One post-op complication in arthroscopic group vs. open</td>
<td>“Although other patient-preferred benefits such as improved earlier return of motion may still exist, the results of our study suggest that the technique of arthroscopic surgery does</td>
<td>High drop out rate. At 12 months, recurrence rates between these two procedures are comparable and arthroscopy is not superior to open procedure.</td>
</tr>
<tr>
<td>Study</td>
<td>Methodology</td>
<td>Participants</td>
<td>Design</td>
<td>Setting</td>
<td>Interventions</td>
<td>Outcomes</td>
<td>Comments</td>
</tr>
<tr>
<td>---------------</td>
<td>-------------</td>
<td>--------------</td>
<td>--------</td>
<td>---------</td>
<td>---------------</td>
<td>----------</td>
<td>----------</td>
</tr>
<tr>
<td>Tian 2007</td>
<td>RCT</td>
<td>N = 62 (70 hands) with CTS. Mean age 52 years.</td>
<td>Endoscopic vs. Open carpal tunnel release.</td>
<td>Follow-up assessments taken at 3 months and final follow-up ranged from 18 to 48 months.</td>
<td>No significant difference between endoscopic and open groups for 2-point discrimination score at 3 months: 5.3 vs. 5.9 (p &gt;0.05). Rate of scar tenderness significantly lower in Endoscopic group vs. Open Group: 36.0% vs. 65.0% (p &lt;0.05). Mean operation time significantly lower in Endoscopic Group vs. Open Group; 12 vs. 38 minutes (p &lt;0.01).</td>
<td>“The endoscopic carpal tunnel release is a reliable method in the treatment of idiopathic carpal tunnel syndrome. It has the advantages of slight scar tenderness, less operation time, less in-hospital stay, early functional recovery, safety and high satisfaction compared with open methods.”</td>
<td></td>
</tr>
<tr>
<td>Sorensen 2013</td>
<td>RCT</td>
<td>N = 38 requiring endoscopic carpal tunnel release verified using neurophysiological testing; Mean (range) age 49 (31-76) for LA group and 52 (36-69) for IVRA group.</td>
<td>Local anesthesia group receiving 10ml (4mL given in proximal direction under subcutaneous fascia, 4mL subcutaneously in palm and 2mL subcutaneously in the distal wrist crease) Ropivacaine (n = 19) vs. Intravenous regional anesthesia group receiving 1% Mepivacaine (n = 19). Assess at baseline, during surgery, immediately after surgery, 2 hours and 24 hours post-op.</td>
<td>Immediately after surgery and 2 hours post-op, significant differences in mean (SD) VAS arm pain reported between LA and IVRA group: End of surgery: 0.2 (0.6) vs. 1.4 (1.8), (p &lt;0.05), 2 hours post-op: 0.2 (0.5) vs. 1.4 (1.8), (p &lt;0.05). During drug administration and immediately after surgery, significant differences in mean (SD) VAS arm pain reported between LA and IVRA groups: During administration: 2.1 (2.6) vs. 4.3 (1.7), (p &lt;0.05), End of surgery: 0.6 (0.9) vs. 2.4 (2.3), (p &lt;0.05).</td>
<td>“[L]A is generally a safe and effective method for ECTR after installing the LA in the subcutaneous tissue and under the subcutaneous fascia (in a proximal direction) alone, without installation of LA into the carpal tunnel...LA was more effective than IVRA at reducing patient-experienced overall pain at the end of the operation and pain in the hand 2 hours later. Furthermore, patients required less additional analgesia after surgery with LA than those treated under IVRA.”</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reed Group, Ltd. 2007</td>
<td>Double-blind</td>
<td>N = 25 (50 hands) with bilateral carpal tunnel syndrome; Mean (±SD) age 57 (±10) for all participants.</td>
<td>Right handed injection (n = 25 hands) vs. Left handed injection (n = 25 hands). All participants received allocated hand treatment upon randomization, followed by treatment on opposite hand 6-12 weeks later.</td>
<td>In comparison of mean (±SD) unadjusted VAS scores and adjusted VAS scores for buffered and non-buffered lidocaine, there were significant differences: Buffered lidocaine unadjusted- 4.60 (±1.50), adjusted- 4.63 (±1.32), vs. Buffered lidocaine unadjusted- 4.60 (±1.50), adjusted- 4.63 (±1.32), vs.</td>
<td>“[T]he results proved the buffered lidocaine could reduce the pain experienced during local anesthetic injection before carpal tunnel release.”</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Anesthesia during Surgery**

**Lee 2013**

RCT Double-blind
Sponsored by the Seoul National University Hospital

- **Participants**: N = 25 (50 hands) with bilateral carpal tunnel syndrome; Mean (±SD) age 57 (±10) for all participants.
- **Interventions**: Right handed injection (n = 25 hands) vs. Left handed injection (n = 25 hands). All participants received allocated hand treatment upon randomization, followed by treatment on opposite hand 6-12 weeks later.
- **Outcomes**: In comparison of mean (±SD) unadjusted VAS scores and adjusted VAS scores for buffered and non-buffered lidocaine, there were significant differences: Buffered lidocaine unadjusted- 4.60 (±1.50), adjusted- 4.63 (±1.32), vs.
- **Comments**: “[T]he results proved the buffered lidocaine could reduce the pain experienced during local anesthetic injection before carpal tunnel release.”

**Methodological details**: Sparse methodology.
<table>
<thead>
<tr>
<th>Study</th>
<th>Methodological details</th>
<th>N</th>
<th>Case details</th>
<th>Intervention</th>
<th>Follow up</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Braithwaite 1993</td>
<td>Randomized trial (?)</td>
<td>2.5</td>
<td>N = 23 requiring carpal tunnel release; Participant ages not reported. Assessment at baseline and after each injection.</td>
<td>0.5% Bupivacaine injection alongside 1:200,000 adrenaline without tourniquet (n = 23 arms) vs. 0.5% Bupivacaine alone and pneumatic tourniquet (n = 23 arms). All received both treatments, but on randomized arms. Assessments at baseline, post-op and 14 days.</td>
<td>During procedure, participants demonstrated higher mean (SD) VAS pain scores with tourniquet compared to adrenaline limb: 4.7 (2.8) vs. 2.3 (1.7), (p &lt;0.01). Participants’ symptom diaries had no difference in paresthesia, post-op pain or bruising when comparing adrenaline and tourniquet limbs 14 days post-op.</td>
<td>“The use of adrenaline-containing local anaesthetic provides a satisfactory operative field, avoids the discomfort of a tourniquet and allows bilateral simultaneous carpal tunnel release to be accomplished without the need for general anaesthesia.”</td>
</tr>
<tr>
<td>Ozer 2005</td>
<td>RCT</td>
<td>2.5</td>
<td>N = 40 requiring surgical decompression of carpal tunnel. Mean age 48.2 (30-64) alkalised group: 52.8 (42-67) non-alkalised group.</td>
<td>Alkalised group received 10ml prilocaine hydrochloride 2% buffered with 1ml sodium bicarbonate 8.4% (n = 20) vs. non-alkalised group receiving 10ml prilocaine hydrochloride 2% (n = 20). Assessment baseline, hourly for 6 hours post-op and 12 hours.</td>
<td>At 1, 3, 6, 12 hours post-op, alkalised group exhibited significantly lower mean (SD) VAS scores vs. non-alkalinised: 1 hour- 0 vs 0.5 (0.52), (p = 0.02), 3 hours- 0.12 (0.35) vs. 1.75 (1.05), (p = 0.001), 6 hours- 1.12 (0.35) vs. 2.16 (1.33), (p = 0.036), and 12 hours- 2.12 (0.83) vs. 2.75 (0.75), (p = 0.06).</td>
<td>“Buffered prilocaine provides a longer pain-free period for patients following surgical decompression of the median nerve. It is easy, safe, and cost-effective and it appears that the routine use of alkalised prilocaine solution in patients undergoing carpal tunnel surgery may improve the comfort and prolong the duration of analgesia.”</td>
</tr>
<tr>
<td>Watts 2004</td>
<td>RCT</td>
<td>1.5</td>
<td>N = 64 undergoing local anesthesia for open carpal tunnel decompression: Mean (range) age 57 (28-89) years for both groups.</td>
<td>Buffered lidocaine group receiving 5ml of 2% plain lidocaine plus 0.5ml sodium bicarbonate 8.6% (n = 32) Vs. Plain lidocaine group receiving 5ml of 2% plain lidocaine plus 0.5ml of sodium chloride 0.9% (n = 32). Assessments at baseline and post-op.</td>
<td>Although both groups reported pain improvement, here were no statistically significant results reported between groups for mean VAS pain scores, verbal pain scores or anxiety scores.</td>
<td>“[T]he pain of injection is not actually a major problem for most patients undergoing carpal tunnel decompression and there is no benefit in injecting buffered lidocaine. The pain scores for both groups were low and most patients reported that they were “not at all” anxious about having a similar injection again in the future. We did however note a correlation between increased pain score and increased”</td>
</tr>
</tbody>
</table>

**Methodological details sparse.**
Watts 2005

RCT

No mention of sponsorship or COI.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Watts 2005</td>
<td>1.5</td>
<td>N = 86</td>
<td>27-gauge dental needle group (n = 46) vs. 23-gauge needle group (n = 40). Both groups received 4.4ml of 2% xylocaine with adrenaline 1:80,000 with pre-filled 2.2ml vials. Assessments at baseline and post-op.</td>
<td>Participants receiving injection via 27-gauge dental needle had significantly lower mean (SEM) VAS pain scores vs. standard 23-gauge needle: 22 (2.4) vs 33 (3.8), (p &lt;0.02). Not significant when analyzing verbal response scale. Participants also self-reported less mean (SEM) anxiety with 27-gauge needle vs. 23-gauge: 7 (1.4) vs. 15 (3.4), (p &lt;0.05).</td>
<td>&quot;[P]atients reported less anxiety about future injections when the pain of the injection was reduced.&quot;</td>
<td>Methodological details sparse.</td>
</tr>
</tbody>
</table>

Yiannakopoulos 2004

RCT

No sponsorship or COI.

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yiannakopoulos 2004</td>
<td>1.5</td>
<td>N = 64 requiring carpal tunnel decompression verified by electrodiagnostic and clinical evidence alongside local anesthesia; Mean (SD) age 61 (8) years for all participants.</td>
<td>Group A; Lidocaine 1% mixed with normal saline group (n = 20) Vs. Group B; 10ml alkalized lidocaine 1% at room temperature (22ºC) (n = 22) Vs. Group C; 10 ml alkalized lidocaine warmed in 40ºC water bath for 30 minutes (n = 22) All groups received allocated treatment into palmar skin. Assessments at baseline and post-op.</td>
<td>Mean (SD) infiltration pain scores were significantly lower in Groups B&amp; C compared to Group A: A- 21(11) &amp; 42 (12), B- 25 (12) &amp; 19 (7) vs. C- 21 (4) &amp; 10 (4), (p&lt;0.001). Group C also had significantly lower values compared to Group B, (p&lt;0.001).</td>
<td>&quot;We have found that buffering lidocaine with bicarbonate and warming the anesthetic solution helps to reduce pain on infiltration in patients undergoing carpal tunnel decompression.&quot;</td>
<td>Methodological details sparse.</td>
</tr>
</tbody>
</table>

MALLET FINGER

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gruber 2014</td>
<td>3.5</td>
<td>N = 51 with fractured or unfractured mallet finger. Mean (±SD) age 49 (±14) for splint group and 51</td>
<td>Full-time custom-made thermoplastic splint group (n = 25) vs. No splint control group (n = 26). Follow up at 4 weeks.</td>
<td>No significant differences reported between splint and control groups for average final extensor lag, disability or treatment satisfaction.</td>
<td>&quot;[T]here is not much benefit to additional night splinting after completing the standard splinting protocol for mallet finger. The extra cost and time associated with obtaining a custom-made removable splint should be balanced with the patient’s preferences. It is possible that a subset of Data suggests night splinting did not improve mallet finger outcomes in terms of extensor lag, disability or treatment satisfaction.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>N</td>
<td>Splint Type</td>
<td>Duration</td>
<td>Success Rate</td>
<td>Comment</td>
</tr>
<tr>
<td>-----------------------</td>
<td>--------</td>
<td>---</td>
<td>-------------</td>
<td>----------</td>
<td>--------------</td>
<td>-------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Garberman 1994</td>
<td>RCT</td>
<td>75</td>
<td>Stack vs. Dorsally placed aluminum-foam splint. Splinted continuously for 6-10 weeks, then nightly for 4 weeks.</td>
<td>Splint treatment success (with no more than (10^\circ) extensor lag) in 17 of 21 (81.0%) in early group and 15 of 19 (78.9%) in delayed splint group. Fractures and type of splint immaterial.</td>
<td>“Splinting was as effective in the delayed treatment population as it was in the early treatment population.”</td>
<td>Study design unclear as described as both retrospective and randomized. Dropout rate also unclear.</td>
</tr>
<tr>
<td>Kinninmonth 1986</td>
<td>RCT</td>
<td>44</td>
<td>Perforated vs. Stack splint. Splinted at least 6 weeks.</td>
<td>Successes were 79% Stack vs. 84% perforated splint.</td>
<td>“The perforated mallet finger splint can produce consistently good results even in those patients who would not tolerate a conventional splint. The fact that it is unnecessary to remove it for hygiene purposes is to its advantage.”</td>
<td>High success rates, but methods sparse.</td>
</tr>
</tbody>
</table>
### EXTENSOR COMPARTMENT TENOSYNOVITIS

#### MRI

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest</th>
<th>Score</th>
<th>Number</th>
<th>Area</th>
<th>Diagnoses</th>
<th>Type of MRI used</th>
<th>Type of CT used</th>
<th>T1 weighted</th>
<th>T2 weighted</th>
<th>More than one rater</th>
<th>Surgery Performed</th>
<th>Clinical outcomes</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hadidy 2009</td>
<td>Diagnosti  c</td>
<td>No mention of Sponsors hip of COI</td>
<td>2.5</td>
<td>N=29 patients (21 Females, and 8 males) clinically diagnosed with De Quervain’s Tenosynovitis; Mean Age 49.4 ± 17.5.</td>
<td>Wrist</td>
<td>De Quervain’s Tenosynovitis</td>
<td>1.5 T</td>
<td>N/A</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>Ultra-Sonography positive findings frequency (percentage) based upon following parameters; Tendon Thickening, Peritendinous edema, septation, subcutaneous edema, synovial thickening: 100%, 97.5%, 70%, 35%, 50%. MRI positive findings (Percentage) based upon the same parameters, respectively: 95%, 85%, 60%, 55%, 45%.</td>
<td>&quot;Our statistical results indicate that ultrasound is, in general a more useful technique than MRI in identifying abnormalites in patients with De Quervain disease.&quot;</td>
</tr>
<tr>
<td>Author/Year</td>
<td>Study Type</td>
<td>Scoring</td>
<td>Sample Size</td>
<td>Comparison Group</td>
<td>Results</td>
<td>Conclusion</td>
<td>Comments</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------------</td>
<td>---------------------</td>
<td>---------</td>
<td>-------------</td>
<td>------------------------------------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sawaizumi 2007</td>
<td>RCT</td>
<td>3.0</td>
<td>N = 36</td>
<td>Intra-sheath triamcinolone injection (1ml TC and 1ml of 1% lidocaine hydrochloride)</td>
<td>The 1-point injection excellent in 9 hands (50%), and 2-point injection excellent in 15 hands (75%); p &lt;0.001.</td>
<td>Accurate injection of triamcinolone into sheath of both extensor pollicis brevis and abductor pollicis longus tendon considered very effective for deQuervain's disease.</td>
<td>Selection for treatment based on consecutive cases rather than randomization.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Avci 2002</td>
<td>RCT with pseudo-randomization</td>
<td>3.0</td>
<td>N = 19 wrists (18 females) with de Quervain's positive Finkelstein's. All pregnant or lactating.</td>
<td>Glucocorticosteroid injection (methylprednisolone 10mg plus 0.5mL 0.5% bupivacaine vs. thumb spica splint. Follow-up until asymptomatic and had stopped nursing (mean 12 months).</td>
<td>Complete relief in 100% of injection group vs. 0% splint group, though pain reportedly relieved while wearing splint. Recurrences in one injected patient.</td>
<td>&quot;Splinting does not provide satisfactory pain relief.&quot;</td>
<td>Population was pregnant or lactating. Small sample size. Sparse details. Randomization was every other. Data suggest injection superior to splinting.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kosuwon 1996</td>
<td>RCT</td>
<td>2.0</td>
<td>N = 140 with de Quervain's. Duration of symptoms unstated.</td>
<td>Steroid injection (dose and medication not specified) with vs. without wrist immobilization</td>
<td>Satisfactory results in 74% splinted vs. 75% unsplinted (NS). Lost days in splinted group mean 28 vs. 11, p&lt;0.05.</td>
<td>&quot;There is no difference in the results of treatment in this condition whether the patients were immobilized in a splint or not. However, the days lost from work in the group of non immobilization is less than the group of immobilization.&quot;</td>
<td>Abstract only. Sparse details. Results suggest no difference in outcomes whether wrist immobilized after injection or not.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Witt 1991</td>
<td>Prospective Cohort/Case Series</td>
<td>1.5</td>
<td>N = 95 (99 wrists) with de Quervain's. Symptom duration unstated.</td>
<td>One mL injection of 1% lidocaine plus methylprednisolone acetate 40mg. Minimum 12 months follow-up.</td>
<td>54% satisfactory. 30 wrists required surgical release. 22/30 (73%) of operated wrists had separate EPB compartment.</td>
<td>&quot;Injection of one milliliter of a 1 per cent lidocaine solution and one milliliter of a suspension containing forty milligrams of methylprednisolone acetate. Twelve patients (twelve wrists) were lost to follow-up. Of the remaining eighty-seven wrists, fifty-four (62 per-cent) had a satisfactory outcome at a mean of eighteen months (minimum follow-up, twelve months). The duration of symptoms before treatment did not affect the outcome.&quot;</td>
<td>Not randomized trial. Primary purpose was to assess steroid flare. 73% of treatment failures had separate compartment for extensor pollicis brevis. Baseline symptom duration not predictive.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## NON-SPECIFIC HAND, WRIST, AND FOREARM PAIN

### Electrodiagnostic Studies

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Study Type</th>
<th>Conflict of Interest</th>
<th>Population/Case Definition</th>
<th>Investigative Test</th>
<th>Gold Standard / Comparative Test</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calder 2009</td>
<td>3.5</td>
<td>Diagnostic</td>
<td></td>
<td>N = 46 (22 asymptomatic control subjects, 8 at-risk subjects, and 16 subjects with non-specific arm pain. Mean age 38.1 years.</td>
<td>Surface electromyographic (SEMG) activity</td>
<td>Comparing controls with patients and patients at risk.</td>
<td>Age significantly different among groups; control subjects significantly younger (p &lt;0.05). Mean spike amplitude (MSA) significantly increased by 039 mV across all levels of % maximum contraction in patients with NSAP, showing 325% increase (p &lt;0.05). At-risk group showed significant increase of 0.43 mV (430%) from 10% to 70% of MVC (p &lt;0.05). In healthy controls MSA increased 1.1 mV (550%) from 10 to 70% of MVC (p &lt;0.05).</td>
<td>“The NSAP group presented with differences in how the spike shape measures change with increasing contraction level that may be indicative of myogenic changes, a result that is consistent with previous quantitative EMG findings.”</td>
<td>Controls significantly younger than study group which may influence results. Spike shape differences in EMG testing may provide valuable information in evaluating neuromuscular disorders.</td>
</tr>
</tbody>
</table>

## RADIAL NERVE ENTRAPMENT

### Electrodiagnostic Studies

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Study Type</th>
<th>Conflict of Interest</th>
<th>Population/Case Definition</th>
<th>Investigative Test</th>
<th>Gold Standard / Comparative Test</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Verhaar 1991</td>
<td>3.5</td>
<td>Diagnostic</td>
<td>No sponsorship or COL.</td>
<td>N = 16 (11 males, 5 females) with chronic pain of the lateral elbow who</td>
<td>Force measurements, nerve conduction studies and concentric-needle</td>
<td>Werne criteria</td>
<td>No statistically significant differences reported in the motor latencies</td>
<td>“Our data do not support the hypothesis that the signs and symptoms in most patients who have radial tunnel syndrome do not have evidence of compression in the posterior interosseous nerve.”</td>
<td>Data suggest patients with radial tunnel syndrome do not have evidence of compression in the posterior interosseous nerve.</td>
</tr>
<tr>
<td>Spindler 1990</td>
<td>Diagnos tic</td>
<td>N = 3 (0 males, 3 females) with no evidence of neurological disease from superficial branch of radial nerve (SBRN) and lateral antebrachial cutaneous nerve (LACN) in the forearm. Mean age 39 years.</td>
<td>Conduction study of SBRN and LACN</td>
<td>Not stated</td>
<td>LACN: stimulation of radial nerve at elbow evoked response over LACN in forearm. Normal radial sensory conduction in forearm: 61.4±3.1 m/sec. Difference between LACN and SBRN conduction in same arm: 1.9±1.6 m/sec. We therefore recommend that during all examinations for possible entrapment of the radial nerve in the forearm, the musculocutaneous nerve at the elbow also be stimulated. If the response is recordable from the radial sensory pickup electrodes while stimulating the musculocutaneous nerve, radial sensory conduction studies should be performed.</td>
<td>Small sample size (N=30) Data suggest value in stimulating the musculocutaneous nerve at the elbow when evaluating RNE.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### SCAPHOID FRACTURES

#### Bone Scans

| Author/Year | Study Type | Conflict of Interest (COI) | Score | N | Area of Body | Diagnoses | Type of Bone Scans | CT used | MRI Used | More than one rater | Blinding of rater | Myelography | Surgery Performed | Clinical outcomes assessed | Long term follow up (mean when noted) | Results | Conclusion | Comments |
|-------------|------------|---------------------------|-------|---|-------------|-----------|-------------------|---------|----------|-------------------|----------------|------------|------------------|---------------------------|-------------------|------------|-----------|
| O’Carroll 1982 | Diagnostic | 3.5 | 30 | Wrist | Scaphoid fracture. Mean age 32 years. | 99m Tc Methylene Diphosphonate and large field of view Gamma camera. | - | - | - | - | - | - | - | 6 weeks | 6 of 30 patients had scaphoid fractures. All 6 with scaphoid fractures gave positive bone scan but 5 additional patients without fractures gave positive bone scans. | Bone scanning, however, detected all scaphoid fractures but had a relatively high false positive rate. | Data suggest scintigraphy accurately diagnosed all true fractures and accurately detected those without fracture. |

#### MRI

| Author/Year | Study Type | Conflict of Interest (COI) | Score | Number | Area of Body | Diagnoses | Type of MRI used | Type of CT used | T1 weighted images | T2 weighted images | X-ray | Myelography | Surgery Performed | Clinical outcomes assessed | Long term follow up (mean when noted) | Results | Conclusion | Comments |
|-------------|------------|---------------------------|-------|--------|-------------|-----------|-------------------|----------------|-------------------|-------------------|-------|------------|------------------|---------------------------|-------------------|------------|-----------|
| Kumar 2005 | Diagnostic | 3.5 | N=22 patients (17 male, 5 females) with suspected scaphoid fractures from an HWF | Scaphoid fractures | 1.5 T MRI (Siemens Vision) | N/A | + | - | - | - | - | - | - | - | Scaphoid fractures detected in 6 of 22 participants based on low signal intensity on T1W. Using MRI within 24 hours of trauma had 100% sensitivity (CI: 72.3-100) and | Our study demonstrates that MRI can be used for early assessment of occult scaphoid fractures within 24 hours of injury. In a tertiary level hospital containing a MRI facility | Small sample size. Data suggest MRI may be effective to detect occult scaphoid fractures. |
| Imaeda 1992 Diagnostic | 3. 5 | N=28 patients (25 males, 3 females) Mean Age: 26 | WRI   | Scaphoid Fracture | General electric signa system operated at 1.5 tesla with a 3 inch circular surfaces coil two-dimensional Fourier transform multisecti on using an 8-10 cm. field of view, and | - | + | + | - | - | - | - A fresh fracture was identified by decreased or iso signal intensity on the T1-weighted image and increased signal intensity on the TZ-weighted image. This increase continued until bony union was apparent on radiographs. On the TZ-weighted image, high signal intensity was characteristic of fresh fractures and suggested “Signal intensity on T1- and T2-weighted images can be used to indicate the prognosis of each scaphoid fracture. A low signal intensity on a T1-weighted image combined with a high signal intensity on a T2-weighted image would serve as a basis for a diagnosis of scaphoid fracture and provide some indication as to the likelihood of | 100% specificity (CI: 75.8-100). operating at least 5 days a week, it is possible to accommodate an MRI of the wrist with a modified shortened protocol. This would lead to early initiation of treatment at the time of presentation and decrease the number lost to follow-up. The proposed clinical pathway would also reduce the over treatment of patients allowing scaphoid fracture to be ruled out at presentation—thus enabling early return to work instead of the usual 10 days in a plaster cast.” | No mention of sponsorship or COI. | Data suggest MRI depicts increased visualization of wrist anatomy which is useful for diagnosing and assessing the extent of union of scaphoid fracture. |
sections 3 mm. thick taken with a 1.5 mm. interspace gap. Spin-echo pulse sequences and T1-weighted (T.R. = 500-600 msec., T.E. = 20-30 msec.), Proton Density weighted (T.R. = 2000 msec., T.E. = 20-30 msec.) and T2-weighted (T.R. = 2000 msec., T.E. = 80 msec.) images that bony union was possible. When bony union was complete, the intensity of the signal for the scaphoid on both T1- and T2-weighted images returned to normal. healing. When bony union is complete, the signal intensity for both proximal and distal fragments on the T1- and T2-weighted images resumes the iso level.”

| Sharifi 2015 | 3. 5 | N=175 patients with suspected scaphoid fracture. 163 | Wrist Scaphoid Fracture Type not mentioned | n/a | - | - | - | - | - | + | Negative MRI exam was found in 159 patients (90.9%) and positive in 16 patients (9.1%). T-test in both groups mean pain score “It seems that although this method, beside other criteria in the diagnosis of fractures, can be useful in future Data suggest pain measurement in combination with MRI for suspected bone fractures is not useful in
males, 12 females. Mean age 28.06±5.952 years.

In negative and positive MRI groups was 7.3±1.75 and 8.75±1.52 respectively that was p<.002 (95% CI). Sensitivity and specificity will be 94% and only 8% for scores 4.5 and higher, 94% and 18% for score of 5.5 and higher, and 94% and 38 for scores of 6.5 and higher respectively. While the sensitivity decreases down to 87% and the specificity increases up to 57% for score 7.5 and the sensitivity and specificity will be 75% and 72% for scores 8.5 and higher, respectively and finally, the sensitivity and specificity will be 31% and 87% for scores 9.5 and higher, respectively.

Gaebler 1996
N=32 patients with clinically suspected scaphoid fracture. Wrist Scaphoid Fracture 1.5 Tesla + + + + - - + + Fracture of the scaphoid was not confirmed in 26 cases. The MRI scans showed no indication of fracture, but in 11 cases, lesions of small sample. Data suggest MRI is useful in diagnosing scaphoid fracture.
21 males, 11 females. Mean age: 29.5 years.

Seneviratne 2013

2. 5

N=110 patients with suspected scaphoid fracture. 72 males, 38 females. Mean age: 35.14 years.

Wrist Scaphoid Fracture 1.5 Tesla - - - - - - - - Twenty-four scans showed occult bone fractures. Three of these patients had scaphoid fractures diagnosed by MRI. Seventy-two patients had other incidental findings with MRI.

the carpus, carpal ligaments, and the radius were found. Scaphoid fracture was found in 6 patients by conventional radiographs after an average time of 16.3 days and for these 6 cases in 3.5 days. Specificity of MRI exams for occult scaphoid fractures was 100% (95% CI 89.12-100) and sensitivity of 100% (95% CI 60.7-100). In 11 cases scaphoid fracture could not be confirmed. In 2 cases an occult fracture of capitatum was diagnosed by MRI. In 3 cases, injuries of the radius were found.

immediately to prevent patients with simple wrist sprains from being immobilized for weeks and thus save money in our health system. The advantages of our method to diagnose occult scaphoid fracture with MRI are several: no radioactive diagnostic means are required, sensitivity and specificity are 100%, the method is readily available at our clinic, and $7200 per 100,000 inhabitants could be saved by routine use of this conventional method.

Data suggest performing MRI 2 weeks after an acute wrist injury is useful in visualization of multiple wrist injuries including soft tissue and many other non-scaphoid wrist fractures.
| Schmitt 2011 | 2.5 | N=10 patients with radial-sided wrist pain. 4 males, 6 females. Mean age 36.9 years. | Wrist Scaphoid Fracture | 1.5 Tesla 16-row scanner | + + - - + + + + | Two cases with congenital thumb hypoplasia, the entire scaphoid was affected by extensive volume loss and osteosclerosis. Six patients presented with altered scaphoid shapes. In remaining 2 patients, shapes of scaphoid were unremarkable. In all patients, increased density of bone substance was evident. A gradient of osteosclerosis was evident in other 8 cases with bone density being most intense at the proximal and decreasing to distal pole. The sclerotic and cystic changes of scaphoid bone structure were better assessed in “Pathoanatomy of Preiser’s disease and the differentiation into three zones of bone marrow viability can be explained with the retrograde blood supply of the scaphoid. In its natural course, three different stages can be depicted with the initial stage seen only in MRI.” | Very small sample. Data suggest MRI beneficial in visualization of anatomy of the three bone marrow zones in Preiser’s disease when compared to radiographs. |
### Surgery

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jeon2009 RCT</td>
<td>3.5</td>
<td>41 (39 males, 2 females) patients with an acute scaphoid fracture. Mean age was 28.5 years.</td>
<td>Volar Approach Group- A transverse stab incision was made at about 1 cm distal to the scaphotrapezial joint (N=19) Vs. Dorsal Approach Group- After stab incision over the center of the circle (usually over the scapholunate joint), the guide wire was driven dorsal to volar so that it exited at the radial base of the thumb. (N=22) Follow-up at 4, 8 and 12 weeks.</td>
<td>There were no significant differences between Dorsal and Volar approach for Pain (p=0.91), Flexion (p=0.989), Extension (p=0.300), Radial Deviation (p=0.380), Ulnar Deviation (p=0.414), and Modified Mayo wrist score (p=0.446).</td>
<td>“This study suggests that screws are placed more parallel to the long axis of the scaphoid and perpendicular to the fracture line via the dorsal approach; however, there was no significant difference with regard to functional outcome and bone union.”</td>
<td>Data suggest comparable efficacy.</td>
</tr>
</tbody>
</table>

### METACARPAL FRACTURES

Functional Therapies vs. Casting or Splinting

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
</table>

---

9 patients assessed with CT imaging.
<table>
<thead>
<tr>
<th><strong>Conflict of Interest (COI)</strong></th>
<th><strong>Hansen 1998 RCT</strong></th>
<th><strong>Immobilization</strong></th>
<th><strong>Sørensen 1993 RCT</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>No mention of sponsorship or COI.</td>
<td>N = 105 with fracture of neck of ring or little metacarpal bone</td>
<td>Dorso-ulnar plaster-of-Paris from proximal interphalangeal joint to elbow (n = 35) vs. functional brace around wrist (n = 35) vs. elastic bandage (n = 35). Study duration 4 weeks. Follow-up at 3 months.</td>
<td>N = 133 with fractures (140) of 2nd-5th metacarpal bones. Age range 10+.</td>
</tr>
<tr>
<td><strong>Immobilization</strong></td>
<td><strong>VAS during 4 weeks treatment: plaster-of-Paris 1.5 vs. functional brace 1.8 vs. elastic bandage 2.7 (p &lt; 0.05). Median restriction of MCPJ movement at 4 weeks: plaster-of-Paris 20º vs. functional brace 0º vs. elastic bandage 10º (p &lt; 0.05). Median restriction of MCPJ movement at 3 months: plaster-of-Paris 0º vs. functional brace 0º vs. elastic bandage 10º (p &lt; 0.05).</strong></td>
<td><strong>Percent reduced mobility 4 weeks after injury after cast removal: metacarpal brace 4% vs. cast 31% (p &lt; 0.01).</strong></td>
<td></td>
</tr>
<tr>
<td><strong>DISTAL FOREARM FRACTURES</strong></td>
<td><strong>“Patients treated with a functional brace mobilized as fast as patients treated with elastic bandage and faster than patients treated with plaster-of-Paris.”</strong></td>
<td><strong>“We found that the benefits did not outweigh the risks of the functional fracture bracing, and we cannot recommend the test version of the Galveston metacarpal brace.”</strong></td>
<td><strong>High drop out in carpal-brace group (58%) compared to plaster-of-Paris (19%). Therefore, comparisons of the two treatment groups not possible. At 3 months the patients completing study reported equal mobility.</strong></td>
</tr>
<tr>
<td>Author/Year</td>
<td>Study Type</td>
<td>Score (0-11)</td>
<td>Sample Size</td>
</tr>
<tr>
<td>-------------</td>
<td>------------</td>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Lagerström</td>
<td>Clinical trial</td>
<td>3.5</td>
<td>N = 33</td>
</tr>
<tr>
<td>Scand J Rehabil Med 1999;31:49-54</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pasila 1974</td>
<td>RCT</td>
<td>3.0</td>
<td>N = 135</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oskarsson 1997</td>
<td>Case series</td>
<td>N/A</td>
<td>N = 110</td>
</tr>
<tr>
<td>Van Der Linden 1981</td>
<td>RCT</td>
<td>3.5</td>
<td>N = 250 (39 male/211 female)</td>
</tr>
</tbody>
</table>
## Immobilization/Fixation

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wik 2009 RCT</td>
<td>3.5</td>
<td>N = 72 females with low-energy trauma, displaced Colles’ fractures initially considered for closed reduction and immobilization in plaster cast. Age &gt;50.</td>
<td>Reduction and a complete plaster cast (n = 34) vs. Reduction and a dorsal plaster splint (n = 38). Immobilization for 5 weeks with follow-up at 1 and 10 days and 5 weeks after reduction.</td>
<td>Mean dorsal angulation 10 days after reduction: slightly better in the dorsal plaster splint group, (p = 0.04). Radial length at 5 weeks was better in the complete plaster group, (p = 0.02).</td>
<td>“Surgeons caring for such cases may choose the immobilization method for the first 10 days following reduction according to their individual preferences and those of the injured person.”</td>
<td>Data suggest comparable efficacy between 2 groups suggesting personal preference for type of immobilization method.</td>
</tr>
<tr>
<td>Gupta 2011 RCT</td>
<td>3.5</td>
<td>(N=40) (9 females/31 males) patients younger than 50 years of age with a fracture distal radius with dorsal angulation more than 20 degrees, severe dorsal comminution, radial shortening more than 10 mm, and wrist joint fracture.</td>
<td>Closed reduction and long arm cast applied under brachial plexus block (Group A, N=20), followed after one week to check radiograph for any displacement; plaster removed at 6 weeks and then physiotherapy was started; reassessed at 3 months v. closed reduction and external fixation with poly axial clamp and distractor rod under brachial plexus block (Group B, N=20); plaster removed at 6 weeks and then physiotherapy was started; reassessed at 3 months.</td>
<td>Expenditure incurred (Rs) (mean±SD): Group A (1894.30±251.11) v. Group B (8156.84±377.33), p=0.00.</td>
<td>“[B]oth the methods of treatment produced no significant difference in outcome, functionally and anatomically, but the extra cost of anesthetic drugs, operation theatre charge, hazards of anesthesia, cost of implant can be avoided in cases of closed reduction and immobilization in plaster cast.”</td>
<td>Data suggest comparable efficacy between groups (unstable distal radius fractures treated either with closed reduction plus cast vs. closed reduction and external fixation lead to same functional and anatomical outcomes.</td>
</tr>
</tbody>
</table>

### DISTAL PHALANX FRACTURES

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antibiotic Prophylaxis</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sloan 1987</td>
<td>3.5</td>
<td>N = 85 (81 males; 4 females) adult</td>
<td>No antibiotics (N = 10) vs.</td>
<td>Infection rate (no antibiotic vs</td>
<td>“Three different antibiotics regimes were</td>
<td>Data showed similar efficacy with all 3</td>
</tr>
</tbody>
</table>
RCT
No mention of industry sponsorship or conflict of interest (COI)

<p>| Patients with open fracture of the distal phalanges of less than 6 hours; mean age 36.9 ± 15.9 | Cephradine 500mg 4 times for 5 days (N=24) vs. Cephradine 1gm injection pre-operatively and Cephradine 500mg 4 times for 5 days (N = 24) vs. Cephradine 1gm injection pre-operatively and 1gm orally post-operatively. (N = 25) Follow up 2 and 5 days and thereafter as appropriate | Antibiotic): 30% vs 2.7% (p=0.02) compared, with no difference in the infection rate: therefore the simplest and surest method, a single pre-operative dose and a single post-operative dose, is recommended.” | Antibiotic groups but the non-antibiotic group had a 30% infection rate. Data suggest a single pre-op and a single post-op dose of antibiotics is effective. |</p>
<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score</th>
<th>Number</th>
<th>Area of Spine</th>
<th>Diagnoses</th>
<th>Type of X-rays</th>
<th>CT used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes assessed</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sakamoto 2013</td>
<td>Clinical Study</td>
<td>3.5</td>
<td>N=17 (7 males, 10 females). Mean age is 48.9 years.</td>
<td>Wrist/Knee area.</td>
<td>Ganglion Cyst</td>
<td>Radiograph</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>Interossseous ganglion appears as a well-defined osteolytic lesion located near a joint.</td>
<td>Radiographs as well as clinical information are important for the accurate diagnosis of intraosseous ganglia.</td>
<td>Data suggest plain radiographs and clinical information are important in making an accurate diagnosis of intraosseous ganglia.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conflict of Interest (COI)</td>
<td>Khan 2011 Randomized Control Trial</td>
<td>Aspirations</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------------------------</td>
<td>----------------------------------</td>
<td>-------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>No sponsorship or COI.</td>
<td>N = 36 with dorsal wrist ganglion; mean Age 31 (17-45)</td>
<td>Group 1 (N=18) Patients treated with an open surgical excision. Vs. Group 2 (N=18) Patients treated using aspiration with 18G needle, followed by injection of triamcinolone acetonide. Follow-up at baseline, 1, 2, 6 weeks, and 6 months. Success Rate, group 1 vs group 2: 17 (94.4%) vs 11 (61.1%) (p = 0.041). Rate of Recurrence, Group 1 vs Group 2: 1 (5.6%) vs 7 (39.9%) (p = 0.041). No complications in any of study groups during study period. “Although the aspirations, triamcinolone acetonide injection plus wrist immobilization is one of the alternative methods, surgery was the most successful form of treatment when considering the cure rate of dorsal wrist ganglion, though we analyzed only a small group; our results can only be an indicator.” Small sample size. Data suggest surgical excision superior to aspiration plus triamcinolone plus wrist immobilization for treatment of dorsal wrist ganglion (94.4% vs 61.1%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Varley 1997 RCT</td>
<td>N = 85 (63 female /22 male) patients that had a ganglion cyst for more than three months Age plus range for aspiration alone (83 (13-71)) vs Aspiration and steroid injection (N = 43 (31 female/12 male)). Follow-up mean and range (46 weeks (26-89))</td>
<td>Both treatment types produced a 33% success rate. “This study demonstrates that 33% of ganglia are successfully treated by aspiration: additional injection of steroid injection is of no benefit and may cause subcutaneous fat atrophy and skin depigmentation.” Data suggest similar efficacy between groups with addition of steroid adding no benefit and may increase skin depigmentation and fat atrophy.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Aspiration (without other intervention)
Aspiration and Surgical Excision and Steroid Injection

Balazs 2015
Randomized Control Trial
No sponsorship or COI.

| N=125 (85 males, 40 females). Mean age is 31 years. | Recurrence (N=11) Vs No Recurrence (N=114) Mean follow up of 45 months. | Recurrence incidence of 9% over 45 months. | Patients whose occupation or activities require forceful wrist extension should be counseled on the considerable risk of residual pain and functional limitations that may occur after open dorsal wrist ganglion excision. | Data suggest persistent pain post open dorsal wrist ganglion excision in active duty military personnel is common and these persons should be counselled on the risk of residual pain post procedure. |

HAVS Diagnostic Testing

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score</th>
<th>N</th>
<th>Area of Body</th>
<th>Diagnoses</th>
<th>Type of Thermography</th>
<th>CT used</th>
<th>MRI Used</th>
<th>More than one rater</th>
<th>Blinding of rater</th>
<th>Myelography</th>
<th>Surgery Performed</th>
<th>Clinical outcomes</th>
<th>Long term follow-up (mean when noted)</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coughlin 2001</td>
<td>Same as OCC MED Case Control</td>
<td>5</td>
<td>31 subjects in two groups. Group A: 10 healthy volunteers. 5 men, 5 women. Median age of 35. Group B: 21 patients. 20 men, 1 woman.</td>
<td>Hand</td>
<td>HAVS with RP</td>
<td>Cold Provocation</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>After cold provocation, the finger temperature and time for the finger temperature to return to pre-cooling levels were able to distinguish the HAVS group and</td>
<td>“CPT has a good sensitivity, specificity, positive predictive value and negative predic”</td>
</tr>
<tr>
<td>Median age of 45</td>
<td>the normal group. The sensitivity of CPT was low after cooling, but reach up to 95% 3 min after rewarming. The accuracy of the test was also the greatest towards the last stages of rewarming. The specificity and PPV were high during precooling stages and remained relatively high during the rewarming stages. NPV was low during the precooling stage and became high (&gt;90) during the rewarming stages.</td>
<td>Positive value; it strongly supports the clinical diagnosis of digital vasospasm.</td>
<td>digital vasospasm.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Author/Year</td>
<td>Score</td>
<td>Study Design</td>
<td>Population/Case Definition</td>
<td>Investigative Test</td>
<td>Gold Standard/Comparative Test</td>
<td>Results</td>
<td>Conclusion</td>
<td>Comments</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------------</td>
<td>-------</td>
<td>--------------</td>
<td>-----------------------------</td>
<td>--------------------</td>
<td>---------------------------------</td>
<td>------------------------------------------------------------------------</td>
<td>------------</td>
<td>----------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bogadi-Šare 1994</td>
<td>3.5</td>
<td>Case Control</td>
<td>N = 45 men 29 chain-saw workers VS 16 healthy controls Mean age = 47</td>
<td>Finger Thermometry</td>
<td>Photoplethysmography</td>
<td>Chain-saw workers had significantly lower temperature compared to the control group. Viewing the pulse amplitude can distinguish the control group from the chain-saw workers, but cannot determine the subgroups. Plethysmography yielded most suitable results at a diagnostic limit of 75% reduction amplitude, which gave a sensitivity of 62% and a specificity of 87%.</td>
<td>“The present study presents a supplementar y diagnostic procedure which reflects the degree of vasoconstriction and vasodilatative ability, can confirm the clinical impression, and can be used for assessment of disability.”</td>
<td>Data suggest there is considerable variation to cold provocation in terms of the vascular response which impedes the defining of normal vs. abnormal reactions. No single test could distinguish cases from controls.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lindsell 1999</td>
<td>3.0</td>
<td>Case Control</td>
<td>N = 107 76 subjects exposed to vibration VS</td>
<td>Vibrotactile threshold measurement, thermal threshold measurement, and finger systolic blood</td>
<td>No gold standard mentioned. All investigational test were compared against each other.</td>
<td>Results showed that as the overall numbness score increased, cold thresholds decreased</td>
<td>“The present study shows that thermal thresholds (hot and cold) and vibrotactile Data suggest some vascular and neurological signs occur independently but some signs like blanching and numbness and</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Design</td>
<td>N</td>
<td>Exposed Group</td>
<td>Controls Group</td>
<td>Exposure</td>
<td>Test</td>
<td>Findings</td>
<td>Notes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------------</td>
<td>------</td>
<td>----------</td>
<td>-------</td>
<td>-----------------------------------</td>
<td>----------------</td>
<td>----------</td>
<td>-------------------------------</td>
<td>---------------------------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kurozawa 1991</td>
<td>2</td>
<td>Case-Control</td>
<td>122</td>
<td>100 men receiving annual exam for vibration syndrome</td>
<td>22 healthy men</td>
<td>VWF</td>
<td>Finger systolic blood pressure after cold provocation</td>
<td>Finger systolic blood pressure was positively correlated with vibrotactile thresholds at 125 Hz (p=0.016). Also, the numbness score was positively correlated with vibrotactile thresholds at 125 Hz (p=0.033) and hot thermal thresholds (p=0.019). Finger systolic blood pressure did not correlate with the other test ran.</td>
<td>Tingling may be related as they are highly correlated.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lawson 1997</td>
<td>2</td>
<td>Case-Control</td>
<td>34</td>
<td>Cold Provocation testing</td>
<td></td>
<td>VWF</td>
<td>Sensorineural tests after initial questionnaire</td>
<td>“Although neither sensorineural data suggests multiple tests are required to make an</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Manually workers not exposed to vibration (Control)

No mention of age or gender

Kurozawa 1991

Case Control

N = 122 men

100 men receiving annual exam for vibration syndrome

Vs 22 healthy men

Mean age = 56.0

Finger systolic blood pressure after cold provocation

Cold-water immersion test

The exposed with severe VWF group had a FSBP significantly different compared to the other three groups. The FSBP resulted in a specificity of 81.7% and sensitivity of 90.3% for VWF.

Cold water immersion test showed no significant differences in the skin temperature and recovery rates of three groups.

“Measurements of skin temperature after cold-water immersion were found not to be feasible for discriminatin among exposed men without VWF, those with mild VWF, and those with severe VWF.”

Data suggest skin temperature measurements pre and post immersion in cold water for 10 minutes cannot be used to estimate the severity of vibration induced white finger.
exposed to hand transmitted vibration
Mean age = 47.1

examination, the participants were split into different sensorineural stages (0, 1, 2, or 3). Cold provocation testing was done after the examination, which resulted in a sensitivity of 75% and a specificity of 61%.
test or cold provocation test results alone are confirmatory, collectively they are very supportive of a diagnosis of the hand-arm vibration syndrome”

accurate diagnosis of HAVS.

---

### Serologic Testing or Connective Tissue Disorders Testing

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Score</th>
<th>Study Design</th>
<th>Population/Case Definition</th>
<th>Investigative Test</th>
<th>Cold Standard / Comparative Test</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kennedy 1999</td>
<td>3.0</td>
<td>Case Control</td>
<td>N = 16 men with RP secondary to vibration white finger disease (Median age of 59.5 yrs) Vs 8 healthy men (Median age of 59.0 yrs)</td>
<td>sICAM-1 and IL-8 enzyme immunoassay</td>
<td>Skin perfusion (erythrocyte flux)</td>
<td>Patients with HAVS have a statistically higher level of sICAM-1 (median 284.5ng/mL, range 168.2-531.8ng/mL) than the healthy patients (median 209.7ng/mL, range 99.6-285.5ng/mL, p=0.02). The healthy patients had higher IL-8 levels (Median “The increase in sICAM-1 that occurs in patients with HAVS suggests that leucocyte adhesion is increased and that adherent neutrophils may contribute to the microvascular damage seen in this disease.” Very small sample (n=11). Data suggesting patients with HAVS had higher S-ICAM-1 levels than controls.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
27.7pg/mL, range 16.1-70.4pg/mL) than the patients with HAVS (median 13.8pg/mL, range 11.5-25.4pg/mL, P<0.01). The skin perfusion was not significantly different between the two groups.

### LACERATION MANAGEMENT

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Conflict of Interest (COI) Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mouzas 1975</td>
<td>RCT</td>
<td>3.5</td>
<td>N = 104</td>
<td>Dexon suture vs. silk suture vs. polyethylene suture vs. nylon suture</td>
<td>One wound in each Dexon, polyethylene, and nylon groups was frankly injected, 4 wounds sutured with silk injected. By 7-10 days 77.3% (17/22) of Dexon wound, 68.2% (15/22) of polyethylene wound and 73.9% (17/23) of nylon wound.</td>
<td>“Dexon was seen to possess certain advantages in that it caused as little tissue reaction as the other sutures but did not have to be removed subsequently.”</td>
<td>Not clear if an RCT as randomization and allocation not described. No blinding.</td>
</tr>
<tr>
<td>Sutton 1985</td>
<td>RCT</td>
<td>3.5</td>
<td>N = 76</td>
<td>4/0 Ethilon interrupted mattress sutures vs. Steristrips applied on tincture of benzoin for closure of wounds.</td>
<td>&quot;Sutures appeared to be associated with increased necrosis of the wound and slower healing than adhesive tapes, particularly when used for flap lacerations…The mean healing time for the 23 patients whose flap lacerations were closed with tapes was 39 days; 20 of these patients were neither admitted to hospital nor received grafts.”</td>
<td>“This study shows that for most pretibial lacerations conservative management on an outpatient basis is all that is necessary, and that adhesive tapes are to be preferred for the primary closure of such wounds.”</td>
<td>Lack of study details. May not be applicable to upper extremity lacerations.</td>
</tr>
<tr>
<td>Bernard 2001</td>
<td>RCT</td>
<td>3.5</td>
<td>N = 42</td>
<td>2-octyl cyanoacrylate vs. standard suture for the closure of excisional wounds</td>
<td>No differences in early complications between groups. Suture group scored higher on VAS (63.3mm for suture vs. 47.8mm for tissue adhesive); difference statistically significant (p = 0.02). Suture group had higher median score on Hollander Wound Scale, but not statistically significant (p = 0.09).</td>
<td>“The cosmetic outcome of cutaneous excisional surgery wounds closed with standard suturing was found to be superior to that of wounds closed with octyl cyanoacrylate.”</td>
<td>Study not random-selection based on patient choice. Study population children and adolescents, but may be appropriate for excision wounds in general, all wounds</td>
</tr>
</tbody>
</table>
Sutures were used in MacGregor's 1989 RCT study, involving 3.5 scores awarded by subjects for ease and satisfaction of closure by doctors at insertion, compared to staples. Significantly more patients awarded staples full marks at insertion for method acceptability, although they were same at removal. The study concluded that the use of staples to close traumatic skin lacerations compares favorably with the traditional method of suturing. A sparse study details and lack of analytical details were also noted.

### HUMAN AND ANIMAL BITES AND ASSOCIATED LACERATIONS

<table>
<thead>
<tr>
<th>Author/Year Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>MacGregor 1989 RCT</td>
<td>3.5</td>
<td>N = 100</td>
<td>Staple vs. suture closure with local anesthetic for patients with lacerations.</td>
<td>Scores awarded for ease and satisfaction of closure by doctor at insertion were similar. Significantly more patients awarded staples full marks at insertion for method acceptability, although they were same at removal.</td>
<td>&quot;The use of staples to close traumatic skin lacerations compares favorably with the traditional method of suturing.&quot;</td>
<td>Sparse study details. Lack of analytical details.</td>
</tr>
<tr>
<td>Elenbaas 1982 RCT</td>
<td>3.5</td>
<td>N = 63</td>
<td>Oxacillin x 5 days vs. placebo.</td>
<td>No significant difference in infection rates between two groups; 2 infections vs. 0 in antibiotic group. Both developed in hand.</td>
<td>&quot;Good wound toilet and attention to adequate follow-up wound care will result in a minimal incidence of infection in dog bite injuries. Antibiotic prophylaxis does not further reduce this incidence.&quot;</td>
<td>High dropout rate (17/63). Study details sparse, including allocation and blinding methods.</td>
</tr>
<tr>
<td>Maimaris 1988 RCT</td>
<td>3.5</td>
<td>N = 169</td>
<td>Sutures vs. no sutures of dog-bite lacerations.</td>
<td>Overall infection rate 7.7%. No significant difference in infection rate between sutured and non-sutured lacerations. Significant difference in infection rate of hand vs. rest of body (p &lt;0.01).</td>
<td>&quot;Dog bite lacerations should receive thorough surgical treatment and can be safely sutured at presentation. However, special care should be given to hand wounds and patients with delayed presentation.&quot;</td>
<td>Sparse study details. No blinding. Randomization and allocation details not provided.</td>
</tr>
<tr>
<td>Author/Year Study Type</td>
<td>Score (0-11)</td>
<td>Sample Size</td>
<td>Comparison Group</td>
<td>Results</td>
<td>Conclusion</td>
<td>Comments</td>
</tr>
<tr>
<td>------------------------</td>
<td>--------------</td>
<td>-------------</td>
<td>------------------</td>
<td>---------</td>
<td>------------</td>
<td>----------</td>
</tr>
<tr>
<td>Berggren 2001 RCT</td>
<td>2.5</td>
<td>N = 33 wait-listed for CMC joint replacement</td>
<td>Three groups: 1) technical accessories, 2) semi-stable textile splint, and 3) non-stabilizing leather splint. All received advice on ADLs.</td>
<td>Patients’ need for operation over 7 years were 3, 4, and 3 respectively over 7 months and 2 additional patients during rest of 7 years (1 each in each splint group).</td>
<td>“We therefore recommend that patients with arthritis of the carpometacarpal joint of the thumb are offered a similar programme in addition to access to accessories and splints preoperatively.”</td>
<td>Methodological details sparse; 7-year follow-up a strength. No differences between the groups results in suggestions of either equal in/efficacy.</td>
</tr>
<tr>
<td>Adams 2014 RCT</td>
<td>3.0</td>
<td>N = 18, 27 thumbs (11 female/3 male), those with symptoms of thumb base osteoarthritis, AUSCAN score &gt;5 for thumb base pain, and hand function score &gt;9</td>
<td>Optimal NHS occupational therapy vs Optimal NHS occupational therapy and biomechanically active splint vs Optimal NHS occupational therapy and biomechanically inactive splint</td>
<td>Differences in AUSCAN pain scores were statistically significant between the three groups at 4 weeks (F (2,18) = 5.892, p = 0.011). Other differences in various measures were also statistically significant: AUSCAN stiffness (F (2,18) = 22.629, p &lt; 0.001), Michigan Hand Outcomes Questionnaire ADL Both Hands (F (2,18) = 15.352, p &lt; 0.001), MHQ Aesthetics Right Hand (F (2,18) = 4.545, p = 0.025), MHQ Aesthetics Left Hand (F (2,18) = 6.018, p = 0.010), and Global Rating of Change Scale (F (2,20) = 3.640, p = 0.047).</td>
<td>“Over a 4-week period patients who received thumb base splints recorded significantly less improvement in pain scores, hand disability and global change than patients who received optimal NHS OT care or optimal NHS OT care plus placebo splints. This study demonstrated that placebo splints were credible and that further work needs to be conducted on the effectiveness and efficacy of splinting for thumb base OA.”</td>
<td>Abstract only. Small sample. Data suggest thumb splints showed not to be used for thumb OA.</td>
</tr>
<tr>
<td>Weiss 2000 RCT Crossover</td>
<td>3.0</td>
<td>N = 26 (21 female/5 male) with several grades of first carpometacarpal osteoarthritis and radiography</td>
<td>Short splint first, splint immobilized carpometacarpal joint only vs Long splint first, splint immobilized metacarpophalangeal joint, carpometacarpal joint, and wrist</td>
<td>Both splints decreased thumb pain significantly when evaluated with a Friedman RM ANOVA rank test (X²F = 6.2, p = 0.001). The effect of either the long or short splint on thumb pain was not statistically significant. There was no significant difference in pinch strength with either splint when measured with a one-factor repeated-measures ANOVA (F = 4.862, p = 0.033, df = 1, 23).</td>
<td>“Splinting is an effective conservative treatment that helps reduce pain in patients with osteoarthritis of the first CMC joint. However, the splints did not increase pinch strength nor did they lessen the pain associated with pinch. Splinting also helped reduce subluxation (under inclusion of 8 participants who had additional problems: carpal tunnel syndrome, scaphotrapezial trapezoid arthritis, and de Quervain tenonitis). Data suggest splinting for first carpometacarpal joint may reduce pain but functional outcomes changes such as</td>
<td></td>
</tr>
</tbody>
</table>

**SPLINTING AND EXERCISE**

<table>
<thead>
<tr>
<th>Splint vs. Splint</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Berggren 2001 RCT</td>
<td>2.5</td>
<td>N = 33 wait-listed for CMC joint replacement</td>
<td>Three groups: 1) technical accessories, 2) semi-stable textile splint, and 3) non-stabilizing leather splint. All received advice on ADLs.</td>
<td>Patients’ need for operation over 7 years were 3, 4, and 3 respectively over 7 months and 2 additional patients during rest of 7 years (1 each in each splint group).</td>
<td>“We therefore recommend that patients with arthritis of the carpometacarpal joint of the thumb are offered a similar programme in addition to access to accessories and splints preoperatively.”</td>
<td>Methodological details sparse; 7-year follow-up a strength. No differences between the groups results in suggestions of either equal in/efficacy.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Splint vs. No Splint</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adams 2014 RCT</td>
<td>3.0</td>
<td>N = 18, 27 thumbs (11 female/3 male), those with symptoms of thumb base osteoarthritis, AUSCAN score &gt;5 for thumb base pain, and hand function score &gt;9</td>
<td>Optimal NHS occupational therapy vs Optimal NHS occupational therapy and biomechanically active splint vs Optimal NHS occupational therapy and biomechanically inactive splint</td>
<td>Differences in AUSCAN pain scores were statistically significant between the three groups at 4 weeks (F (2,18) = 5.892, p = 0.011). Other differences in various measures were also statistically significant: AUSCAN stiffness (F (2,18) = 22.629, p &lt; 0.001), Michigan Hand Outcomes Questionnaire ADL Both Hands (F (2,18) = 15.352, p &lt; 0.001), MHQ Aesthetics Right Hand (F (2,18) = 4.545, p = 0.025), MHQ Aesthetics Left Hand (F (2,18) = 6.018, p = 0.010), and Global Rating of Change Scale (F (2,20) = 3.640, p = 0.047).</td>
<td>“Over a 4-week period patients who received thumb base splints recorded significantly less improvement in pain scores, hand disability and global change than patients who received optimal NHS OT care or optimal NHS OT care plus placebo splints. This study demonstrated that placebo splints were credible and that further work needs to be conducted on the effectiveness and efficacy of splinting for thumb base OA.”</td>
<td>Abstract only. Small sample. Data suggest thumb splints showed not to be used for thumb OA.</td>
</tr>
</tbody>
</table>

| Weiss 2000 RCT Crossover | 3.0 | N = 26 (21 female/5 male) with several grades of first carpometacarpal osteoarthritis and radiography | Short splint first, splint immobilized carpometacarpal joint only vs Long splint first, splint immobilized metacarpophalangeal joint, carpometacarpal joint, and wrist | Both splints decreased thumb pain significantly when evaluated with a Friedman RM ANOVA rank test (X²F = 6.2, p = 0.001). The effect of either the long or short splint on thumb pain was not statistically significant. There was no significant difference in pinch strength with either splint when measured with a one-factor repeated-measures ANOVA (F = 4.862, p = 0.033, df = 1, 23). | “Splinting is an effective conservative treatment that helps reduce pain in patients with osteoarthritis of the first CMC joint. However, the splints did not increase pinch strength nor did they lessen the pain associated with pinch. Splinting also helped reduce subluxation (under inclusion of 8 participants who had additional problems: carpal tunnel syndrome, scaphotrapezial trapezoid arthritis, and de Quervain tenonitis). Data suggest splinting for first carpometacarpal joint may reduce pain but functional outcomes changes such as |
c evidence of condition
Age = Mean of 57

<table>
<thead>
<tr>
<th>Intensive vs. Standard Exercise</th>
<th>3.5</th>
<th>N = 60 hospitalized RA patients</th>
<th>Intensive (daily HEP, greater number of repetitions) vs. standard exercise program for 12 weeks.</th>
<th>At 14 weeks, grip strength favored intensive group (p = 0.04).</th>
<th>Compared with a traditional programme, an intensive hand exercise programme is well tolerated and more effective in improving hand function in patients with RA.</th>
<th>Non-randomized, as first 30 assigned standard treatment and next 30 intensive. Suggests superiority of more intensive exercise regimen for severely affected RA.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rønningen 2008</td>
<td></td>
<td>Controlled clinical trial</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.5</td>
<td>N = 35 women with thumb base osteoarthriti s</td>
<td>Control (N = 16) – Participated in JP programme Vs Splint and Exercise (SE) Group (N = 19) – Splints 24h/day combined with daily home exercise</td>
<td>There was no significant difference between the two groups at baseline and before the intervention. The SE group showed a significant decrease in the pain at night, pain on motion, stiffness and disability in daily activities over the period before/after the intervention and before/at the follow up. The SE group increased their hand grip force by 27% between the intervention and at 1-year follow up.</td>
<td>The splinting and exercise regimen added to a JP programme gives a greater improvement of pain, stiffness, grip force and daily activities than the JP programme alone.</td>
<td>Data suggest combination splinting and exercise program combined with a joint protection program improves pain, stiffness and quality of life compared to a joint protection program alone.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Participants reported that while wearing the short splint 42% of daily activities were easier to complete, 51% were the same, and 7% were more difficult. While wearing the long splint 16% of activities were easier, 28% the same, and 56% were more difficult.

Participants rated the short splint higher on a visual analog scale compared to long splints (t = 2.85, p = 0.01).

stress) in the early stages of osteoarthritis. The shorter splint was preferred by most patients with first CMC osteoarthritis. In the past, conservative treatment for first CMC osteoarthritis has been frequently employed without qualitative or quantitative support, but the results of this study suggest that splinting is a well-tolerated and effective means of managing first CMC osteoarthritis.
### NSAIDs

#### Diclofenac vs. Amtolmetin

<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Treatment</th>
<th>Outcome</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Niccoli 2002 RCT</td>
<td>3.5</td>
<td>N = 90 hand, hip or knee OA</td>
<td>Amtolmetin 600mg BID for 3 days then 600mg a day for 11 days vs. Diclofenac 50mg TID vs. Rofecoxib 25mg QD for 2 weeks total treatment.</td>
<td>Diclofenac reduced creatinine clearance. Rofecoxib gained body weight, systolic blood pressure, diastolic blood pressure and serum sodium with decrease in daily urine volume. No significant changes in parameters with AMG. Diclofenac more efficacious than other 2 drugs (p &lt;0.001).</td>
<td>&quot;Diclofenac mainly impaired blood renal flow and the glomerular filtration rate, while rofecoxib negatively influenced the renal sodium-water exchange. AMG demonstrated a renal sparing effect, although the exact mechanism is unclear.&quot;</td>
</tr>
</tbody>
</table>

### COMPLEMENTARY AND ALTERNATIVE THERAPIES

#### Chondroitin vs. Placebo

<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Treatment</th>
<th>Outcome</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Verbruggen 2002</td>
<td>2 RCTs</td>
<td>3.5</td>
<td>N = 46 Chondroitin polysulphate 50mg IM twice weekly for 8 weeks every 4 months (46) or chondroitin sulphate 500mg TID (34) vs. placebo for 3 years</td>
<td>Baseline differences in destructive IP joint OA with CPS 23.9% vs. placebo 47.8%. CS 35.3% vs. placebo CS 35.9% at baseline. However, data presented compared with aggregate placebo group, precluding analysis of CS study alone. DIPs for CS study at 3 years 2.6 vs. 3.5 placebo (p = 0.155). PIPs CS 2.3 vs. 2.8, p = 0.373. MCPs CS 0.4 vs. 0.5, p = 0.70.</td>
<td>&quot;The data recorded during these pilot studies should help investigators to design future long-term clinical experiments.&quot;</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Treatment</th>
<th>Outcome</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rovetta 2002 RCT</td>
<td>3.0</td>
<td>N = 24 DIP and/or PIP joint OA</td>
<td>Chondroitin sulfate 800mg a day plus naproxen 500mg a day vs. naproxen only for 2 years.</td>
<td>Chondroitin plus naproxen group had increase of 1 joint with erosive OA at 1 year and none at 2 years, vs. naproxen group with 6 patients, 7 joints (p &lt;0.05).</td>
<td>&quot;Chondroitin sulfate failed to stop the usual time-associated progression in the number of finger joints presenting erosions in EOA of the hands. It was, however, associated with a lower increase in the number of finger joints with erosions detected after 2 years of radiological observation.&quot;</td>
</tr>
</tbody>
</table>

#### Yoga vs. No Therapy

<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>N</th>
<th>Treatment</th>
<th>Outcome</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Garfinkel 1994 RCT</td>
<td>3.0</td>
<td>N = 26 DIP or PIP joint OA</td>
<td>Yoga (supervised 1x a week for 8 weeks) vs. no program. After 10 weeks, controls offered to cross over (2 did not) and remaining subjects randomized. Six remained in controls.</td>
<td>Tenderness improved in yoga (2.20±1.32 vs. 0.4±0.94, p = 0.001). Range of motion increased (p = 0.002). Improvements in grip strengths did not differ (yoga 4.21±4.69/control 3.36±5.89, p = 0.69).</td>
<td>&quot;This yoga derived program was effective in providing relief in hand OA.&quot;</td>
</tr>
<tr>
<td>Author/Year</td>
<td>Study Type</td>
<td>Score (0-11)</td>
<td>Sample Size</td>
<td>Comparison Group</td>
<td>Results</td>
</tr>
<tr>
<td>-------------</td>
<td>------------</td>
<td>--------------</td>
<td>-------------</td>
<td>------------------</td>
<td>---------</td>
</tr>
<tr>
<td><strong>Mathieux 2009</strong>&lt;br&gt;RCT</td>
<td>3.0</td>
<td>N = 60 early RA</td>
<td>Multidisciplinary (n = 6) team-led program. Video, “comprehensive OT,” motor training, skill training, joint protection, counseling, advice, assistive devices, splints, education, psychosocial support. Treatment for 3 months.</td>
<td>Health Assessment Questionnaire scores: OT (0.19±0.19) vs. controls (0.35±0.32), p &lt;0.001. Dominant hand grip strengths: OT (53.9±24.2 kPa) vs. controls (37.3±22.9), p = 0.021.</td>
<td>“[A]n early extended information programme improved hand function in patients with early RA.”</td>
</tr>
</tbody>
</table>

**POST-OPERATIVE REHABILITATION AND REHABILITATION OF PATIENTS WITH FUNCTIONAL DEFICITS: CTS AND OTHER DISORDERS**

<table>
<thead>
<tr>
<th>Author/Year</th>
<th>Study Type</th>
<th>Score (0-11)</th>
<th>Sample Size</th>
<th>Comparison Group</th>
<th>Results</th>
<th>Conclusion</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Rocchi 2014</strong>&lt;br&gt;RCT</td>
<td>No mention of sponsorship. No COI.</td>
<td>3.5</td>
<td>N = 30 with acute complete tear of ulnar collateral ligament (UCL) of thumb treated with surgery. Mean age 39 years.</td>
<td>Standard spica splint for 4 weeks with motion limited to IP joints (n =15) vs. Modified spica splint with freedom to move MCP joint for 4 weeks with motion on both the IP and MCP joints (n = 15). All patients provided flexion-extension exercises. Follow-up at 1, 2, 6, and 12 months.</td>
<td>No significant differences between groups (no p-values reported for study outcomes).</td>
<td>“This study suggests that the surgical repair of the skier’s thumb lesion, combined with the immediate restoring of active MCP ROM protected by a modified spica splint is effective and safe and allows a faster return to manual activities compared to traditional method of postoperative splinting.”</td>
<td>Small sample but data suggest the early motion management group had less pain at 2 months compared to controls and all functional measures became similar at 12 months. The lost work time is shortened in the early-motion group by 12 days.</td>
</tr>
<tr>
<td><strong>Finsen 1999</strong>&lt;br&gt;RCT</td>
<td>3.5</td>
<td>N = 74 with NCS undergoing open CTR</td>
<td>All bulky dressing for 2 days, then: 1) very light dressing and move wrist and fingers “as much as comfort allowed, but avoid heavy lifting for the first” 6 post-op weeks vs. plaster of Paris splint for 2 weeks and rigid orthosis for 2 more weeks.</td>
<td>VAS pain and discomfort scores (pre/2 weeks/6 weeks/6 months): Immobilized (56/6/6/3) vs. mobilized (51/5/2/2).</td>
<td>“Physiotherapy was usually not prescribed,” apparently as an uncontrolled confounder.</td>
<td>Authors conclude that “4 weeks of postoperative immobilization confers no detectable benefit.”</td>
<td>Sparse data. Pseudorandomization on Norwegian social security number. NCS not required. Data suggest immobilization not indicated. No advantage to splinting after carpal tunnel release surgery.</td>
</tr>
<tr>
<td><strong>Bury 1995</strong>&lt;br&gt;RCT</td>
<td>3.0</td>
<td>N = 40 open CTR patients with 43 carpal tunnel releases evaluated</td>
<td>2 weeks of post-op wrist splinting vs. a bulky dressing only</td>
<td>No statistically significant differences between two groups using subjective parameters of patient satisfaction with outcome and objective parameters of grip and lateral pinch strength,</td>
<td>“We found no beneficial effect from postoperative splinting after open carpal tunnel release when compared to a bulky dressing alone.”</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
complication rates, and digital and wrist range of motion. No clinical evidence of bowstringing could be noted in either group of patients.

Martins 2006

| RCT | 3.0 | N = 52 EDS con-firmed | Post-op immobilization vs. no immobilization for open CTR patients | Average of SSS was 33.38±7.33 in group A and 31.77±7.56 in group B. Post-op, SSS average 11.38±4.57 in group A, and 12.33±4.77 in group B (p = 0.059). | “Wrist immobilization in the immediate post-operative period have no advantages when compared with no immobilization in the end result of carpal tunnel release.” |

POST-OPERATIVE REHABILITATION

### Follow-up After CTR

| Atherton 1999 | RCT | 4.0 | N = 100 with CTR | Follow-up with general practitioner vs. hand clinic with 2-week follow-up. | More wound infections diagnosed in general practice setting (14% vs. 0%). Authors believe “most were given antibiotics, perhaps unnecessarily.” | “The waiting time for assessment and suture removal was shorter at the GP surgery than in the outpatient department… but significantly more patients were diagnosed as having wound infections.” | Sparse details; 1 page report. Randomization unclear. No data on risks for infection. CTR procedure not described. Limitations result low-quality study despite 4.0 grading. |

### Physiotherapy Post-Op

| Naik 2007 | RCT | 2.5 | N = 30 with Colles’ fracture who underwent external fixation and removed after 2 months. Age not reported. | Maitland mobilization technique: moist heat 15 minutes followed by Maitland manipulations (Grade 1 and 2) for 1st week of treatment then Grade 3 and 4 2nd week. (n = 15) vs. Mulligan mobilization technique: most heat for 15 minutes, Mulligan manipulations in pain free glides (n = 15). No mention of follow-up time. | Mean±SD pain relief (Maitland vs. Mulligan): 3.93±1.09 vs. 4.73±1.03 (p = 0.029). Mean±SD ROM (Maitland vs. Mulligan): active ROM 12.060±6.37 vs. 7.730±2.37 (p = 0.020); passive ROM 14.460±8.67 vs. 9.660±2.89 (p=0.05). Mean±SD scores for functional tasks (Maitland vs. Mulligan): 3.2±0.86 vs. 4.4±1.05 (p = 0.002). | “Mulligan’s mobilization technique could be used effectively when the pain predominates while Maitland’s mobilization technique could be effectively used to restore mobility when pain is not the major concern to patients with colles’ fracture.” | Small sample (N = 30). Sparse methodology. Data suggest Mulligan’s better for pain relief. |

### Physical Therapy/Occupational Therapy

| Rasotto 2015 | RCT | 3.5 | Sponsored by Italian Workers’ Compensation Authority (INAIL). COI, Rassotto | Mean±SD difference in pain rating baseline to end of study (IG vs. CG): neck -1.29±2.72 vs. 0.39±2.51 (p = 0.0164); shoulder -0.94±1.09 vs. 0.17±2.02 (p=0.0224); wrist -1.40±1.87 vs. -0.39±0.93 (p = 0.0007). | “This personalized approach suggests a greater effect than a non-personalized standard protocol; however any potential longer term value of customized exercise program deserves further investigation.” | Very high dropout and non-compliance in exercise arm. Individualized treatment. Data suggest strength training may reduce neck and wrist pain among those relatively few who remained |
received a grant from INAIL.

perform normal daily activities (\(n = 34\)). Follow-up at 5 months and within 2 weeks from end of study.

compliant. Data subject to non-interventional control bias.

| Taylor 1994 RCT | 3.5 | \(N = 30\) following removal of plaster after Colles’ fracture. Mean age 62.6±8.8 years. | Experimental group: 5 minutes Maitland passive joint mobilization; superficial heat; active exercises; home advice to use affected wrist/hand for all daily activities vs. control group: sham mobilization (soft tissue massage), superficial heat, active exercises, home advice treated 2x a week. Included in study until discharged from physiotherapy. | N no significant differences between groups. “This clinical trial found that the inclusion of passive joint mobilisation into a physiotherapy treatment regime was no more effective than soft tissue massage at increasing the range of active wrist extension in Colles’ fracture patients following removal of plaster.” | Pilot study with small sample size. Data suggest comparable efficacy between passive joint mobilization and soft tissue massage. |
References


397. Melhorn J. Carpal tunnel syndrome: three points of view on risk and recovery. J Workers Comp. 2006;1555-64.


Copyright© 2016 Reed Group, Ltd.


Horch RE, Allmann KH, Laubenberger J, Langer M, Stark GB. Median nerve compression can be detected by magnetic resonance imaging of the carpal tunnel. Neurosurgery. 1997;41(1):76-82; discussion -3.


Korthals-de Bos IB, Gerritsen AA, van Tulder MW, et al. Surgery is more cost-effective than splinting for carpal tunnel syndrome. Part II: Results of an economic evaluation alongside a randomized controlled trial. *BMC Musculoskel Disord.* 2006;7:686.


Copyright © 2016 Reed Group, Ltd.


1554. Winter G. Formation of the scab and the rate of epithelialization of superficial wounds in the skin of the young domestic pig. 1962.


Copyright © 2016 Reed Group, Ltd.


Copyright © 2016 Reed Group, Ltd.


