

Case Number:	CM15-0111396		
Date Assigned:	06/18/2015	Date of Injury:	04/01/2001
Decision Date:	07/16/2015	UR Denial Date:	06/08/2015
Priority:	Standard	Application Received:	06/09/2015

HOW THE IMR FINAL DETERMINATION WAS MADE

MAXIMUS Federal Services sent the complete case file to an expert reviewer. He/she has no affiliation with the employer, employee, providers or the claims administrator. He/she has been in active clinical practice for more than five years and is currently working at least 24 hours a week in active practice. The expert reviewer was selected based on his/her clinical experience, education, background, and expertise in the same or similar specialties that evaluate and/or treat the medical condition and disputed items/Service. He/she is familiar with governing laws and regulations, including the strength of evidence hierarchy that applies to Independent Medical Review determinations.

The Expert Reviewer has the following credentials:
 State(s) of Licensure: New Jersey, Alabama, California
 Certification(s)/Specialty: Neurology, Neuromuscular Medicine

CLINICAL CASE SUMMARY

The expert reviewer developed the following clinical case summary based on a review of the case file, including all medical records:

This 49-year-old woman sustained an industrial injury on 4/1/2001. The mechanism of injury is not detailed. Diagnoses included shoulder impingement, tendinitis, pain in soft tissues of limb, and enthesopathy. Treatment has included oral medications and physical therapy. Physician notes dated 5/14/2015 show complaints of bilateral shoulder pain. Recommendations include self-directed therapy, Theracane, foam roll, Prilosec, tramadol, Naprosyn, psychological sessions, and chiropractic treatment.

IMR ISSUES, DECISIONS AND RATIONALES

The Final Determination was based on decisions for the disputed items/services set forth below:

Theracane: Upheld

Claims Administrator guideline: The Claims Administrator did not cite any medical evidence for its decision.

MAXIMUS guideline: The Expert Reviewer did not base their decision on the MTUS. Decision based on Non-MTUS Citation Official Disability Guidelines (ODG) Durable medical equipment (DME) Durable medical equipment (DME).

Decision rationale: According to ODG guidelines, DME "Recommended generally if there is a medical need and if the device or system meets Medicare's definition of durable medical equipment (DME) below. Most bathroom and toilet supplies do not customarily serve a medical purpose and are primarily used for convenience in the home. Medical conditions that result in physical limitations for patients may require patient education and modifications to the home environment for prevention of injury, but environmental modifications are considered not primarily medical in nature. Certain DME toilet items (commodes, bed pans, etc.) are medically necessary if the patient is bed- or room-confined, and devices such as raised toilet seats, commode chairs, sitz baths and portable whirlpools may be medically necessary when prescribed as part of a medical treatment plan for injury, infection, or conditions that result in physical limitations. Many assistive devices, such as electric garage door openers, microwave ovens, and golf carts, were designed for the fully mobile, independent adult, and Medicare does not cover most of these items. See also specific recommendations here: Aquatic therapy; Bathtub seats; BioniCare knee device; Bone growth stimulators; Braces; Canes; Cold/heat packs; Compression cryotherapy; Continuous-flow cryotherapy; Continuous passive motion (CPM); Crutches; Cryocuff; Cryotherapy; Dynamic splinting systems; Dynasplint; Electrical stimulators (E-stim); Electromyographic biofeedback treatment; ERMI knee Flexionater/ Extensionater; Flexionators (extensionators); Exercise equipment; Game Ready accelerated recovery system; Home exercise kits; Joint active systems (JAS) splints; Knee brace; Lymphedema pumps; Mechanical stretching devices (for contracture & joint stiffness); Motorized scooters; Neuromuscular electrical stimulation (NMES devices); Orthoses; Post-op ambulatory infusion pumps (local anesthetic); Power mobility devices (PMDs); RS-4i sequential stimulator; Scooters; Shower grab bars; TENS (transcutaneous electrical nerve stimulation); Therapeutic knee splint; Treadmill exerciser; Unloader braces for the knee; Vacuum-assisted closure wound-healing; Vasopneumatic devices (wound healing); Walkers; Walking aids (canes, crutches, braces, orthoses, & walkers); Wheelchair; Whirlpool bath equipment. "The term DME is defined as equipment which: (1) Can withstand repeated use, i.e., could normally be rented, and used by successive patients; (2) Is primarily and customarily used to serve a medical purpose; (3) Generally is not useful to a person in the absence of illness or injury; & (4) Is appropriate for use in a patient's home. (CMS, 2005) The patient suffered from shoulder pain. There is no documentation of motor deficit that requires a Theracane for mobility. Therefore, the request for Theracane is not medically necessary.

Foam Roll: Upheld

Claims Administrator guideline: The Claims Administrator did not cite any medical evidence for its decision.

MAXIMUS guideline: The Expert Reviewer did not base their decision on the MTUS. Decision based on Non-MTUS Citation Official Disability Guidelines (ODG) Durable medical equipment (DME) Durable medical equipment (DME).

Decision rationale: According to ODG guidelines, DME "Recommended generally if there is a medical need and if the device or system meets Medicare's definition of durable medical equipment (DME) below. Most bathroom and toilet supplies do not customarily serve a medical purpose and are primarily used for convenience in the home. Medical conditions that result in physical limitations for patients may require patient education and modifications to the home environment for prevention of injury, but environmental modifications are considered not primarily medical in nature. Certain DME toilet items (commodes, bed pans, etc.) are medically necessary if the patient is bed- or room-confined, and devices such as raised toilet seats, commode chairs, sitz baths and portable whirlpools may be medically necessary when prescribed

as part of a medical treatment plan for injury, infection, or conditions that result in physical limitations. Many assistive devices, such as electric garage door openers, microwave ovens, and golf carts, were designed for the fully mobile, independent adult, and Medicare does not cover most of these items. See also specific recommendations here: Aquatic therapy; Bathtub seats; BioniCare knee device; Bone growth stimulators; Braces; Canes; Cold/heat packs; Compression cryotherapy; Continuous-flow cryotherapy; Continuous passive motion (CPM); Crutches; Cryocuff; Cryotherapy; Dynamic splinting systems; Dynasplint; Electrical stimulators (E-stim); Electromyographic biofeedback treatment; ERMI knee Flexionater/ Extensionater; Flexionators (extensionators); Exercise equipment; Game Ready accelerated recovery system; Home exercise kits; Joint active systems (JAS) splints; Knee brace; Lymphedema pumps; Mechanical stretching devices (for contracture & joint stiffness); Motorized scooters; Neuromuscular electrical stimulation (NMES devices); Orthoses; Post-op ambulatory infusion pumps (local anesthetic); Power mobility devices (PMDs); RS-4i sequential stimulator; Scooters; Shower grab bars; TENS (transcutaneous electrical nerve stimulation); Therapeutic knee splint; Treadmill exerciser; Unloader braces for the knee; Vacuum-assisted closure wound-healing; Vasopneumatic devices (wound healing); Walkers; Walking aids (canes, crutches, braces, orthoses, & walkers); Wheelchair; Whirlpool bath equipment. "The term DME is defined as equipment which: (1) Can withstand repeated use, i.e., could normally be rented, and used by successive patients; (2) Is primarily and customarily used to serve a medical purpose; (3) Generally is not useful to a person in the absence of illness or injury; & (4) Is appropriate for use in a patient's home. (CMS, 2005)

The patient suffered from shoulder pain. There is no documentation of motor deficit that requires a Foam Roll for mobility. Therefore, the request for Foam Roll is not medically necessary.