

Case Number:	CM15-0004440		
Date Assigned:	01/26/2015	Date of Injury:	01/21/2008
Decision Date:	04/02/2015	UR Denial Date:	12/11/2014
Priority:	Standard	Application Received:	01/08/2015

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: North Carolina
Certification(s)/Specialty: Family Practice

CLINICAL CASE SUMMARY

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

The injured worker is a 47 year old male, who sustained an industrial injury on January 21, 2008. He has reported left neck pain. The diagnoses have included left shoulder internal derangement, upper extremity repetitive overuse, cervical facet joint pain, cervical disk protrusion and cervical sprain/strain. Treatment to date has included radiographic imaging, diagnostic studies, surgical intervention of the left shoulder, left carpal tunnel release, conservative therapies, medications and lifestyle modifications. Currently, the IW complains of left neck pain. The injured worker reported an industrial injury in 2008, resulting in chronic left neck pain. He was treated conservatively and surgically without resolution of the pain. Evaluation on June 24, 2014, revealed an improvement of pain and the ability to perform activities of daily living with the use of pain medications. Medications were renewed and a urinary drug screen was recommended to monitor for prescription medication compliancy. Evaluation on October 30, 2014, revealed continued pain. Medications were renewed. On December 11, 2014, Utilization Review non-certified a request for Hydrocodone/APAP tab 10/325mg #90 for continuation of a taper with possible discontinuation over 2-3 months noting the MTUS, ACOEM Guidelines, (or ODG) was cited. On December 22, 2014, the injured worker submitted an application for IMR for review of requested Hydrocodone/APAP tab 10/325mg #90 for continuation of a taper with possible discontinuation over 2-3 months.

IMR ISSUES, DECISIONS AND RATIONALES

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

Hydrocodone/APAP tab 10/325mg #90 for continuation of a taper with possible discontinuation over 2-3 months: Overturned

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

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines opioids
Page(s): 76-84.

Decision rationale: The California chronic pain medical treatment guidelines section on opioids states for ongoing management: On-Going Management. Actions Should Include:(a) Prescriptions from a single practitioner taken as directed, and all prescriptions from a single pharmacy.(b) The lowest possible dose should be prescribed to improve pain and function.(c) Office: Ongoing review and documentation of pain relief, functional status, appropriate medication use, and side effects. Pain assessment should include: current pain; the least reported pain over the period since last assessment; average pain; intensity of pain after taking the opioid; how long it takes for pain relief; and how long pain relief lasts. Satisfactory response to treatment may be indicated by the patient's decreased pain, increased level of function, or improved quality of life. Information from family members or other caregivers should be considered in determining the patient's response to treatment. The 4 A's for Ongoing Monitoring: Four domains have been proposed as most relevant for ongoing monitoring of chronic pain patients on opioids: pain relief, side effects, physical and psychosocial functioning, and the occurrence of any potentially aberrant (or non-adherent) drug-related behaviors. These domains have been summarized as the "4 A's" (analgesia, activities of daily living, adverse side effects, and aberrant drug taking behaviors). The monitoring of these outcomes over time should affect therapeutic decisions and provide a framework for documentation of the clinical use of these controlled drugs. (Passik, 2000)(d) Home: To aid in pain and functioning assessment, the patient should be requested to keep a pain diary that includes entries such as pain triggers, and incidence of end-of-dose pain. It should be emphasized that using this diary will help in tailoring the opioid dose. This should not be a requirement for pain management.(e) Use of drug screening or inpatient treatment with issues of abuse, addiction, or poor pain control.(f) Documentation of misuse of medications (doctor shopping, uncontrolled drug escalation, drug diversion).(g) Continuing review of overall situation with regard to nonopioid means of pain control.(h) Consideration of a consultation with a multidisciplinary pain clinic if doses of opioids are required beyond what is usually required for the condition or pain does not improve on opioids in 3 months. Consider a psych consult if there is evidence of depression, anxiety or irritability. Consider an addiction medicine consult if there is evidence of substance misuse. When to Continue Opioids(a) If the patient has returned to work(b) If the patient has improved functioning and pain(Washington, 2002) (Colorado, 2002) (Ontario, 2000) (VA/DoD, 2003) (Maddox-AAPM/APS, 1997) (Wisconsin, 2004) (Warfield, 2004) The long-term use of this medication class is not recommended per the California MTUS unless there documented evidence of benefit with measurable outcome measures and improvement in function. There is a documented 70 % decrease in pain and a 70% increase in function in the clinical documentation provided for review. Therefore, the request is certified.