

Case Number:	CM15-0053093		
Date Assigned:	03/26/2015	Date of Injury:	08/16/2000
Decision Date:	05/01/2015	UR Denial Date:	02/20/2015
Priority:	Standard	Application Received:	03/20/2015

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: California

Certification(s)/Specialty: Physical Medicine & Rehabilitation, Pain Management

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 60 year old female who sustained a work related injury on August 16, 2000, incurring neck and shoulder pain from repetitive motion and trauma. She was diagnosed with a cervical sprain with radiculopathy, repetitive strain injury, myofascial pain syndrome, rotator cuff tear, and peripheral neuropathy. Treatment included rotator cuff surgery, pain medications, muscle relaxants, acupuncture sessions, sleep aides and home exercise program. Currently, the injured worker complained of persistent right shoulder and neck pain. The treatment plan that was requested for authorization included prescriptions for Norco with a retrospective date of service of February 9, 2015 and Tramadol with a retrospective date of service of February 9, 2015.

IMR ISSUES, DECISIONS AND RATIONALES

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

Retrospective DOS 2/9/15 Norco 10/325mg qty: 30: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Opioids for chronic pain Page(s): 80-81.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines 8 C.C.R. 9792.20 - 9792.26 MTUS (Effective July 18, 2009) Page(s): 44, 47, 75-79, 120 of 127.

Decision rationale: Regarding the request for Norco (hydrocodone/acetaminophen), California Pain Medical Treatment Guidelines state that this is an opiate pain medication. Due to high abuse potential, close follow-up is recommended with documentation of analgesic effect, objective functional improvement, side effects, and discussion regarding any aberrant use. Guidelines go on to recommend discontinuing opioids if there is no documentation of improved function and pain. Within the documentation available for review, there is no indication that the medication is improving the patient's function or pain (in terms of specific examples of functional improvement and percent reduction in pain or reduced NRS) and no discussion regarding aberrant use. As such, there is no clear indication for ongoing use of the medication. Opioids should not be abruptly discontinued, but unfortunately, there is no provision to modify the current request to allow tapering. In light of the above issues, the currently requested Norco (hydrocodone/acetaminophen) is not medically necessary.

Retrospective DOS 2/9/15 Tramadol 50mg qty: 60: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Opioids for chronic pain Page(s): 80-81.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines 8 C.C.R. 9792.20 - 9792.26 MTUS (Effective July 18, 2009) Page(s): 44, 47, 75-79, 120 of 127.

Decision rationale: Regarding the request for tramadol, California Pain Medical Treatment Guidelines state that this is an opiate pain medication. Due to high abuse potential, close follow-up is recommended with documentation of analgesic effect, objective functional improvement, side effects, and discussion regarding any aberrant use. Guidelines go on to recommend discontinuing opioids if there is no documentation of improved function and pain. Within the documentation available for review, there is no indication that the medication is improving the patient's function or pain (in terms of specific examples of functional improvement and percent reduction in pain or reduced NRS) and no discussion regarding aberrant use. As such, there is no clear indication for ongoing use of the medication. Opioids should not be abruptly discontinued, but unfortunately, there is no provision to modify the current request to allow tapering. In light of the above issues, the currently requested tramadol is not medically necessary.