

Case Number:	CM15-0077730		
Date Assigned:	04/29/2015	Date of Injury:	03/10/2010
Decision Date:	05/29/2015	UR Denial Date:	04/09/2015
Priority:	Standard	Application Received:	04/23/2015

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: Maryland, Texas, Virginia

Certification(s)/Specialty: Internal Medicine, Allergy and Immunology, Rheumatology

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 65 year old male, who sustained an industrial injury on 03/10/2010. The initial complaints or symptoms included low back pain/injury. The initial diagnoses were not mentioned in the clinical notes. Treatment to date has included conservative care, medications, MRIs, conservative therapies, injections, and lumbar surgery (12/21/2011). Currently, the injured worker complains of ongoing low back pain, increase swelling in the feet, and increased muscle spasms. The injured worker reported good relief with the use of a H-wave unit, which allows for decreased medication. However, the treatment plan includes continuation of medications (including Tylenol #3, Soma and Zantac). The diagnoses include thoracic/lumbar radiculitis, lumbar disc displacement, and status post bilateral lumbar hemilaminectomy. The request for authorization consisted of Tylenol with codeine (#3), Zantac, and Soma.

IMR ISSUES, DECISIONS AND RATIONALES

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

Tylenol with codeine #3 qty:90: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Criteria for Use of Opioids.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Codeine Page(s): 35. Decision based on Non-MTUS Citation Official Disability Guidelines (ODG) Pain, (Tylenol with Codeine).

Decision rationale: MTUS and ODG state regarding codeine, "Recommended as an option for mild to moderate pain, as indicated below. Codeine is a schedule C-II controlled substance. It is similar to morphine. 60 mg of codeine is similar in potency to 600 mg of acetaminophen. It is widely used as a cough suppressant. It is used as a single agent or in combination with acetaminophen (Tylenol with Codeine) and other products for treatment of mild to moderate pain." ODG further states regarding opioid usage, "Not recommended as a first-line treatment for chronic non-malignant pain, and not recommended in patients at high risk for misuse, diversion, or substance abuse. Opioids may be recommended as a 2nd or 3rd line treatment option for chronic non-malignant pain, with caution, especially at doses over 100 mg morphine equivalent dosage/day (MED)." The medical records do not indicate what first-line treatment was tried and failed. Additionally, medical records do not detail how the patient's pain and functional level with Tylenol with Codeine has improved. As such, the request for Tylenol with Codeine # qty 90 is not medically necessary.

Zantac 150mg qty:60: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines NSAIDs, GI symptoms & cardiovascular risk.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Pain (Chronic), NSAIDs, GI symptoms & cardiovascular risk. Decision based on Non-MTUS Citation Uptodate.com, NSAIDs (including aspirin): Primary prevention of gastroduodenal toxicity.

Decision rationale: Ranitidine is an H2 antagonist used for the treatment of stomach ulcers and gastroesophageal reflux. MTUS states, "Determine if the patient is at risk for gastrointestinal events: (1) age > 65 years; (2) history of peptic ulcer, GI bleeding or perforation; (3) concurrent use of ASA, corticosteroids, and/or an anticoagulant; or (4) high dose/multiple NSAID (e.g., NSAID + low-dose ASA)." And "Patients at intermediate risk for gastrointestinal events and no cardiovascular disease: (1) A non-selective NSAID with either a PPI (Proton Pump Inhibitor, for example, 20 mg omeprazole daily) or misoprostol (200 mg four times daily) or (2) a Cox-2 selective agent. Long-term PPI use (> 1 year) has been shown to increase the risk of hip fracture (adjusted odds ratio 1.44)." Uptodate states regarding H2 antagonist for GI prophylaxis, "Standard doses of H2 receptor antagonists were not effective for the prevention of NSAID-induced gastric ulcers in most reports, although they may prevent duodenal ulcers [33]. Studies that detected a benefit on gastric ulcer prevention were short-term (12 to 24 weeks) and focused on endoscopic rather than clinical endpoints." The patient does not meet the age recommendations for increased GI risk. The medical documents provided do not establish the patient has experienced GI discomfort and does not indicate history of peptic ulcer, GI bleeding or perforation. Medical records do not indicate that the patient is on ASA, corticosteroids, and/or an anticoagulant; or high dose/multiple NSAID. Additionally, uptodate suggests that H2

antagonist at this dose is not useful for to prevent ulcers. As such, the request for Ranitidine 150mg Qty 60 is not medically necessary.

Soma 350mg qty:60: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Carisoprodol (Soma), Muscle relaxants (for pain).

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Carisoprodol (Soma) and Muscle relaxants (for pain) Page(s): 29, 63-66. Decision based on Non-MTUS Citation Official Disability Guidelines (ODG) Chronic Pain, Soma (Carisoprodol).

Decision rationale: MTUS states regarding Crisoprodol, "Not recommended. This medication is not indicated for long-term use. Carisoprodol is a commonly prescribed, centrally acting skeletal muscle relaxant whose primary active metabolite is meprobamate (a schedule-IV controlled substance). Carisoprodol is now scheduled in several states but not on a federal level. It has been suggested that the main effect is due to generalized sedation and treatment of anxiety. Abuse has been noted for sedative and relaxant effects. In regular abusers the main concern is the accumulation of meprobamate. Carisoprodol abuse has also been noted in order to augment or alter effects of other drugs." ODG States that Soma is "not recommended. This medication is FDA-approved for symptomatic relief of discomfort associated with acute pain in musculoskeletal conditions as an adjunct to rest and physical therapy (AHFS, 2008). This medication is not indicated for long-term use." The patient has been on the medication since 2013. Guidelines do not recommend long term usage of SOMA. Treating physician does not detail circumstances that would warrant extended usage. As such, the request for SOMA 350 MG #60 is not medically necessary.