

Case Number:	CM14-0212065		
Date Assigned:	01/02/2015	Date of Injury:	08/06/2010
Decision Date:	02/25/2015	UR Denial Date:	11/21/2014
Priority:	Standard	Application Received:	12/17/2014

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, Michigan, 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:

The patient is a 37-year-old man who sustained a work-related injury on August 6, 2010. Subsequently, the patient developed chronic ankle pain. According to a progress report dated October 30, 2014, the patient reported right ankle pain mostly with movement and activities. The patient denied having any spasms lately but admitted to burning sensation in the right ankle. He also admitted to swelling in all the right toes with prolonged standing and walking. On exam, the patient was not in acute distress. He was asymptomatic. Range of motion of the right ankle was satisfactory. The patient was diagnosed with ankle pain related to the contusion of the Achilles tendon and injury to the calcaneus posteriorly in 2010. The provider requested authorization for Naproxen and Protonix.

IMR ISSUES, DECISIONS AND RATIONALES

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

1 Prescription of Naproxen 500mg #60: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines NSAIDs, Specific Drug List & Adverse Effects; and NSAIDs, GI Sympt.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Non Selective NSAIDS Page(s): 72.

Decision rationale: Naproxen (Naprosyn): delayed release (Naprosyn): as Sodium salt (Anaprox, Anaprox DS, Aleve [otc]) Generic available; extended-release (Naprelan): 375 mg. Different dose strengths and formulations of the drug are not necessarily bioequivalent. Dosing Information: Osteoarthritis or ankylosing spondylitis: Dividing the daily dose into 3 doses versus 2 doses for immediate-release and delayed-release formulations generally does not affect response. Morning and evening doses do not have to be equal in size. The dose may be increased to 1500 mg/day of naproxyn for limited periods when a higher level of analgesic/anti-inflammatory activity is required (for up to 6 months). Naprosyn or naproxyn: 250-500 mg PO twice daily. Anaprox: 275-550 mg PO twice daily. (total dose may be increased to 1650 mg a day for limited periods). EC-Naprosyn: 375 mg or 500 mg twice daily. The tablet should not be broken, crushed or chewed to maintain integrity of the enteric coating. Naprelan: Two 375 mg tablets (750 mg) PO once daily or two 500 mg tablets (1000 mg) once daily. If required (and a lower dose was tolerated) Naprelan can be increased to 1500 mg once daily for limited periods (when higher analgesia is required). Pain: Naprosyn or naproxyn: 250-500 mg PO twice daily. The maximum dose on day one should not exceed 1250 mg and 1000 mg on subsequent days. Anaprox: 275-550 mg PO twice daily. The maximum dose on day one should not exceed 1375 mg and 1100 mg on subsequent days. Extended-release Naprelan: Not recommended due to delay in absorption. (Naprelan Package Insert) > NSAID should be used for the shortest duration and the lowest dose. There is no documentation from the patient file that the provider titrated Naproxen to the lowest effective dose and used it for the shortest period possible. Naproxen was used without clear documentation of its efficacy. Furthermore, there is no documentation that the provider followed the patient for NSAID adverse reactions that are not limited to GI side effect, but also may affect the renal function. Therefore, the request for Naproxen 500mg #60 is not medically necessary.

1 Prescription of Protonix 20mg #30: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Proton Pump Inhibitors (PPIs). Decision based on Non-MTUS Citation Official Disability Guidelines (ODG), Pain

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines NSAIDs, GI symptoms & cardiovascular risk Page(s): 102.

Decision rationale: According to MTUS guidelines, Protonix is indicated when NSAID are used in patients with intermediate or high risk for gastrointestinal events. The risk for gastrointestinal events are: (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). Recent studies tend to show that H. Pylori does not act synergistically with NSAIDs to develop gastroduodenal lesions. There is no documentation that the patient is at an increased risk of GI bleeding. Therefore the prescription of Protonix 20mg # 30 is not medically necessary.

