

Case Number:	CM15-0014425		
Date Assigned:	02/02/2015	Date of Injury:	08/09/2002
Decision Date:	03/24/2015	UR Denial Date:	01/13/2015
Priority:	Standard	Application Received:	01/26/2015

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, 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:

This 63 year old female sustained an industrial injury on 8/9/02, with subsequent ongoing low back pain. Treatment included multilevel lumbar laminectomy and fusion. Following the procedure the injured worker complained of ongoing low back pain and persistent foot drop. In a PR-2 dated 10/21/14, the injured worker complained of ongoing low back pain 8-9/10 on the visual analog scale with medications and 3-4/10 with medications. The injured worker admitted to having problems with control of the medications and had given control to her husband who now dispensed medications for her. Physical exam was unchanged with decreased range of motion to the lumbar spine, numbness to the right leg and foot and diminished lower extremity reflexes. The injured worker was awaiting authorization for additional lumbar fusion. On 1/13/15, Utilization Review modified a request for Norco 10/325mg #120 with 1 refill to Norco 10/325mg #60 with 1 refill, Ambien 10mg #30 to Ambien 10mg #15 and Valium 10mg #60 x 2 to Valium 10mg #30, citing CA MTUS Chronic Pain Medical Treatment Guidelines. As a result of the UR denial, an IMR was filed with the Division of Workers Comp.

IMR ISSUES, DECISIONS AND RATIONALES

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

Norco 10/325mg #120 with 1 refill: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Opioids.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Criteria for use of opioids Page(s): 76-79.

Decision rationale: According to MTUS guidelines, Norco (Hydrocodone/Acetaminophen) is a synthetic opioid indicated for the pain management but not recommended as a first line oral analgesic. In addition and according to MTUS guidelines, ongoing use of opioids should follow specific rules: (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. 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. According to the patient file, there is no objective documentation of pain and functional improvement to justify continuous use of Norco. Norco was used for longtime without documentation of functional improvement or evidence of return to work or improvement of activity of daily living. Therefore, the prescription of Norco 10/325mg #120 with 1 refill is not medically necessary.

Ambien 10mg #30: Upheld

Claims Administrator guideline: The Claims Administrator did not base their decision on the MTUS. Decision based on Non-MTUS Citation Official Disability Guidelines (ODG); Ambien

MAXIMUS guideline: The Expert Reviewer did not base their decision on the MTUS. Decision based on Non-MTUS Citation ODG Non-Benzodiazepine sedative-hypnotics (Benzodiazepine-receptor agonists (<http://worklossdatainstitute.verioiponly.com/odgtwc/pain.htm>

Decision rationale: According to ODG guidelines, Non-Benzodiazepine sedative-hypnotics (Benzodiazepine-receptor agonists): First-line medications for insomnia. This class of medications includes zolpidem (Ambien and Ambien CR), zaleplon (Sonata), and eszopicolone (Lunesta). Benzodiazepine-receptor agonists work by selectively binding to type-1 benzodiazepine receptors in the CNS. All of the benzodiazepine-receptor agonists are schedule IV controlled substances, which means they have potential for abuse and dependency. Ambien is not recommended for long-term use to treat sleep problems. Furthermore, there is no documentation of the use of non pharmacologic treatment for the patient's sleep issue. There is no documentation and characterization of recent sleep issues with the patient. Therefore, the prescription of Ambien 10mg #30 is not medically necessary.

Valium 10mg #60 x 2: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Benzodiazepine. Decision based on Non-MTUS Citation Official Disability Guideline (ODG); Benzodiazepine

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Benzodiazepines Page(s): 24.

Decision rationale: According to MTUS guidelines, benzodiazepines are not recommended for long term use for pain management because of unproven long term efficacy and because of the risk of dependence. Most guidelines limit their use to 4 weeks. There is no recent documentation that the patient have insomnia. Therefore, the prescription of Valium (Diazepam) 10mg #60 x2 is not medically necessary.