

<b>Case Number:</b>	CM15-0097256		
<b>Date Assigned:</b>	05/27/2015	<b>Date of Injury:</b>	08/01/1992
<b>Decision Date:</b>	06/25/2015	<b>UR Denial Date:</b>	05/14/2015
<b>Priority:</b>	Standard	<b>Application Received:</b>	05/20/2015

### 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

### 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 51-year-old female, who sustained an industrial injury on August 1, 1992, incurring, back, neck and shoulder injuries. Cervical Magnetic Resonance Imaging revealed disc protrusion and disc space narrowing. Electromyography studies were abnormal revealing slow motor conduction. She was diagnosed with cervicgia with radiculopathy, and thoracic outlet syndrome. In 2003, a shoulder Magnetic Resonance Imaging showed degenerative changes. In 2013, she was diagnosed with shoulder impingement syndrome and cervical injury. Treatment included physical therapy, anti-inflammatory drugs, pain medications, chiropractic sessions, nerve blocks and work restrictions. Currently, the injured worker complained of persistent cervical pain, stiffness and weakness radiating into both arms. The treatment plan that was requested for authorization included prescriptions for Percocet and Nucynta ER.

### IMR ISSUES, DECISIONS AND RATIONALES

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

**Percocet 5/325mg #180:** Upheld

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

**MAXIMUS guideline:** Decision based on MTUS Chronic Pain Treatment Guidelines Opioids, page(s) 74-96.

**Decision rationale:** Per the MTUS Guidelines cited, opioid use in the setting of chronic, non-malignant, or neuropathic pain is controversial. Patients on opioids should be routinely monitored for signs of impairment and use of opioids in patients with chronic pain should be reserved for those with improved functional outcomes attributable to their use, in the context of an overall approach to pain management that also includes non-opioid analgesics, adjuvant therapies, psychological support, and active treatments (e.g., exercise). Submitted documents show no evidence that the treating physician is prescribing opioids in accordance to change in pain relief, functional goals with demonstrated improvement in daily activities, decreased in medical utilization or change in functional status. There is no evidence presented of random drug testing or utilization of pain contract to adequately monitor for narcotic safety, efficacy, and compliance. The MTUS provides requirements of the treating physician to assess and document for functional improvement with treatment intervention and maintenance of function that would otherwise deteriorate if not supported. From the submitted reports, there is no demonstrated evidence of specific functional benefit derived from the continuing use of opioids with persistent severe pain for this chronic injury without acute flare, new injury, or progressive deterioration. The Percocet 5/325mg #180 is not medically necessary and appropriate.

**Nucynta ER 50mg #60:** Upheld

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

**MAXIMUS guideline:** Decision based on MTUS Chronic Pain Treatment Guidelines Opioids, page(s) 74-96.

**Decision rationale:** Pain symptoms and clinical findings remain unchanged for this chronic injury. Submitted documents show no evidence that the treating physician is prescribing opioids in accordance to change in pain relief, functional goals with demonstrated improvement in daily activities, decreased in medical utilization or returned to work status. There is no evidence presented of random drug testing or utilization of pain contract to adequately monitor for narcotic safety, efficacy, and compliance. The MTUS provides requirements of the treating physician to assess and document for functional improvement with treatment intervention and maintenance of function that would otherwise deteriorate if not supported. From the submitted reports, there is no demonstrated evidence of specific functional benefit derived from the continuing use of opioids with persistent severe pain for this chronic injury. In addition, submitted reports have not adequately demonstrated the specific indication to support for chronic opioid use without acute flare-up, new injuries, or progressive clinical deficits to support for chronic opioids outside recommendations of the guidelines. The Nucynta ER 50mg #60 is not medically necessary and appropriate.