

Case Number:	CM15-0020507		
Date Assigned:	02/10/2015	Date of Injury:	11/26/2001
Decision Date:	04/01/2015	UR Denial Date:	01/07/2015
Priority:	Standard	Application Received:	02/03/2015

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 York, Tennessee
 Certification(s)/Specialty: Emergency 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 injured worker is a 51 year old female who sustained an industrial injury on 11/26/2001. Current diagnoses include lumbar post-laminectomy syndrome, degeneration of lumbosacral intervertebral disc, and opioid dependence. Previous treatments included medication management, physical therapy, and surgery. Report dated 01/16/2015 noted that the injured worker presented with complaints that included low back and left shoulder pain. Physical examination was positive for abnormal findings. Utilization review performed on 01/07/2015 non-certified a prescription for electrodes, skin prep pads, lead wires, and batteries x 6 months, based on the clinical information submitted does not support medical necessity. The reviewer referenced the California MTUS in making this decision.

IMR ISSUES, DECISIONS AND RATIONALES

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

Electrodes, skin prep pads, leads wires, and batteries x 6 months: Upheld

Claims Administrator guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Page(s): 114-121.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Pain Interventions and Guidelines Page(s): 114-115.

Decision rationale: TENS units are not recommended as a primary treatment modality, but a one-month home-based TENS trial may be considered as a noninvasive conservative option, if used as an adjunct to a program of evidence-based functional restoration, including reductions in medication use, for neuropathic pain, phantom limb pain, spasticity, and multiple sclerosis. Several published evidence-based assessments of transcutaneous electrical nerve stimulation (TENS) have found that evidence is lacking concerning effectiveness. Functional restoration programs (FRPs) are designed to use a medically directed, interdisciplinary pain management approach geared specifically to patients with chronic disabling occupational musculoskeletal disorders. These programs emphasize the importance of function over the elimination of pain. FRPs incorporate components of exercise progression with disability management and psychosocial intervention. Chronic intractable pain (for the conditions noted above): (1) Documentation of pain of at least three months duration. (2) There is evidence that other appropriate pain modalities have been tried (including medication) and failed. (3) A one-month trial period of the TENS unit should be documented (as an adjunct to ongoing treatment modalities within a functional restoration approach) with documentation of how often the unit was used, as well as outcomes in terms of pain relief and function; rental would be preferred over purchase during this trial. (4) Other ongoing pain treatment should also be documented during the trial period including medication usage. (5) A treatment plan including the specific short and long-term goals of treatment with the TENS unit should be submitted. (6) After a successful 1-month trial, continued TENS treatment may be recommended if the physician documents that the patient is likely to derive significant therapeutic benefit from continuous use of the unit over a long period of time. At this point purchase would be preferred over rental. (7) Use for acute pain (less than three months duration) other than post-operative pain is not recommended. (8) A 2-lead unit is generally recommended; if a 4-lead unit is recommended, there must be documentation of why this is necessary. In this case the patient had received a defective TENS unit and was unable to return it. There is no documentation that the patient had a successful home trial with the TENS unit and that the patient is participating in a functional restoration program. Criteria for TENS unit use have not been met. Therefore electrodes, skin prep pads, leads, wires, and batteries are not medically necessary. The request should not be authorized.