

Case Number:	CM14-0035014		
Date Assigned:	06/23/2014	Date of Injury:	02/12/2009
Decision Date:	12/26/2014	UR Denial Date:	02/21/2014
Priority:	Standard	Application Received:	03/20/2014

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. The expert reviewer is Board Certified in Occupational Medicine, and is licensed to practice in California. 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/services. He/she is familiar with governing laws and regulations, including the strength of evidence hierarchy that applies to Independent Medical Review determinations.

CLINICAL CASE SUMMARY

The expert reviewer developed the following clinical case summary based on a review of the case file, including all medical records:

According to the records made available for review, this is a 64-year-old female with a 2/12/09 date of injury. At the time (1/7/14) of the request for authorization for Plus 3 month supplies, conductive garment, and X-Force Stimulator Unit (Built in TENS feature) 90 day trial, there is documentation of subjective (none specified) and objective (none specified) findings, current diagnoses (pain in joint and lumbar sprains & strains), and treatment to date (medication, physical therapy, and a brace). There is no documentation of a statement identifying that the TENS unit will be used as an adjunct to a program of evidence-based functional restoration and a treatment plan including the specific short- and long-term goals of treatment with the TENS.

IMR ISSUES, DECISIONS AND RATIONALES

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

Plus 3 month supplies: Upheld

Claims Administrator guideline: The Claims Administrator did not cite any medical evidence for its decision.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Transcutaneous electrical nerve stimulation (TENS) Page(s): 113-117.

Decision rationale: MTUS Chronic Pain Medical Treatment Guidelines identifies documentation of pain of at least three months duration, evidence that other appropriate pain

modalities have been tried (including medication) and failed, a statement identifying that the TENS unit will be used as an adjunct to a program of evidence-based functional restoration, and a treatment plan including the specific short- and long-term goals of treatment with the TENS, as criteria necessary to support the medical necessity of a month trial of a TENS unit. Within the medical information available for review, there is documentation of diagnoses of pain in joint and lumbar sprains & strains. In addition, there is documentation of pain of at least three months duration and evidence that other appropriate pain modalities have been tried (including medication) and failed. However, there is no documentation of a statement identifying that the TENS unit will be used as an adjunct to a program of evidence-based functional restoration and a treatment plan including the specific short- and long-term goals of treatment with the TENS. Therefore, based on guidelines and a review of the evidence, the request for Plus 3 month supplies is not medically necessary.

Conductive garment: Upheld

Claims Administrator guideline: The Claims Administrator did not cite any medical evidence for its decision.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Transcutaneous electrical nerve stimulation (TENS) Page(s): 113-117.

Decision rationale: MTUS Chronic Pain Medical Treatment Guidelines identifies documentation of pain of at least three months duration, evidence that other appropriate pain modalities have been tried (including medication) and failed, a statement identifying that the TENS unit will be used as an adjunct to a program of evidence-based functional restoration, and a treatment plan including the specific short- and long-term goals of treatment with the TENS, as criteria necessary to support the medical necessity of a month trial of a TENS unit. Within the medical information available for review, there is documentation of diagnoses of pain in joint and lumbar sprains & strains. In addition, there is documentation of pain of at least three months duration and evidence that other appropriate pain modalities have been tried (including medication) and failed. However, there is no documentation of a statement identifying that the TENS unit will be used as an adjunct to a program of evidence-based functional restoration and a treatment plan including the specific short- and long-term goals of treatment with the TENS. Therefore, based on guidelines and a review of the evidence, the request for conductive garment is not medically necessary.

X-Force Stimulator Unit (Built in TENS feature) 90 day trial: Upheld

Claims Administrator guideline: The Claims Administrator did not cite any medical evidence for its decision.

MAXIMUS guideline: Decision based on MTUS Chronic Pain Treatment Guidelines Transcutaneous electrical nerve stimulation (TENS) Page(s): 113-117.

Decision rationale: MTUS Chronic Pain Medical Treatment Guidelines identifies documentation of pain of at least three months duration, evidence that other appropriate pain

modalities have been tried (including medication) and failed, a statement identifying that the TENS unit will be used as an adjunct to a program of evidence-based functional restoration, and a treatment plan including the specific short- and long-term goals of treatment with the TENS, as criteria necessary to support the medical necessity of a month trial of a TENS unit. Within the medical information available for review, there is documentation of diagnoses of pain in joint and lumbar sprains & strains. In addition, there is documentation of pain of at least three months duration and evidence that other appropriate pain modalities have been tried (including medication) and failed. However, there is no documentation of a statement identifying that the TENS unit will be used as an adjunct to a program of evidence-based functional restoration and a treatment plan including the specific short- and long-term goals of treatment with the TENS. In addition, the requested 90 day trial exceeds guidelines. Therefore, based on guidelines and a review of the evidence, the request for X-Force Stimulator Unit (Built in TENS feature) 90 day trial is not medically necessary.