

Case Number:	CM15-0116891		
Date Assigned:	06/25/2015	Date of Injury:	01/28/1993
Decision Date:	07/24/2015	UR Denial Date:	06/15/2015
Priority:	Standard	Application Received:	06/17/2015

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: Arizona, California

Certification(s)/Specialty: Family Practice

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 59 year old female, who sustained an industrial injury on January 28, 1993. The injured worker was diagnosed as having chronic pain syndrome, low back pain, radicular symptoms and history of coccygeal fracture. Treatment to date has included knee surgery, medication and injections. A progress note dated April 28, 2015 provides the injured worker complains of back and leg pain. She reports prior injections helped until nerve pain returned three weeks ago. Old magnetic resonance imaging (MRI) from 2009 was reviewed and notable for disc bulges. Physical exam notes full lumbar range of motion (ROM) with pain on extension. There is a request for medial branch block injections.

IMR ISSUES, DECISIONS AND RATIONALES

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

Right L4-L5 medial branch block injection: Upheld

Claims Administrator guideline: Decision based on MTUS ACOEM Chapter 12 Low Back Complaints.

MAXIMUS guideline: Decision based on MTUS ACOEM Chapter 12 Low Back Complaints Page(s): 300. Decision based on Non-MTUS Citation ODG- Low back chapter and pg 36.

Decision rationale: According to the guidelines, Criteria for the use of diagnostic blocks for facet "mediated" pain: Clinical presentation should be consistent with facet joint pain, signs & symptoms. 1. One set of diagnostic medial branch blocks is required with a response of 70%. The pain response should last at least 2 hours for Lidocaine. 2. Limited to patients with low-back pain that is non-radicular and at no more than two levels bilaterally. 3. There is documentation of failure of conservative treatment (including home exercise, PT and NSAIDs) prior to the procedure for at least 4-6 weeks. 4. No more than 2 facet joint levels are injected in one session (see above for medial branch block levels). 5. Recommended volume of no more than 0.5 cc of injectate is given to each joint. 6. No pain medication from home should be taken for at least 4 hours prior to the diagnostic block and for 4 to 6 hours afterward. 7. Opioids should not be given as a "sedative" during the procedure. 8. The use of IV sedation (including other agents such as midazolam) may be grounds to negate the results of a diagnostic block, and should only be given in cases of extreme anxiety. 9. The patient should document pain relief with an instrument such as a VAS scale, emphasizing the importance of recording the maximum pain relief and maximum duration of pain. The patient should also keep medication use and activity logs to support subjective reports of better pain control. 10. Diagnostic facet blocks should not be performed in patients in whom a surgical procedure is anticipated. (Resnick, 2005) 11. Diagnostic facet blocks should not be performed in patients who have had a previous fusion procedure at the planned injection level. In this case, the claimant's pain scores were not provided to determine pain level that would justify a MBB. In addition, the claimant had prior facet injections and the ACOEM guidelines do not recommend invasive procedures due to their short-term benefit. The request for an MBB of L4-L5 is not medically necessary.

Left L4-L5 and bilateral L5-S1 medial branch block injections, quantity of three: Upheld

Claims Administrator guideline: Decision based on MTUS ACOEM Chapter 12 Low Back Complaints.

MAXIMUS guideline: The Expert Reviewer did not base their decision on the MTUS. Decision based on Non-MTUS Citation ODG- pain chapter and pg 36.

Decision rationale: According to the guidelines, Criteria for the use of diagnostic blocks for facet "mediated" pain: Clinical presentation should be consistent with facet joint pain, signs & symptoms. 1. One set of diagnostic medial branch blocks is required with a response of 70%. The pain response should last at least 2 hours for Lidocaine. 2. Limited to patients with low-back pain that is non-radicular and at no more than two levels bilaterally. 3. There is documentation of failure of conservative treatment (including home exercise, PT and NSAIDs) prior to the procedure for at least 4-6 weeks. 4. No more than 2 facet joint levels are injected in one session (see above for medial branch block levels). 5. Recommended volume of no more than 0.5 cc of injectate is given to each joint. 6. No pain medication from home should be taken for at least 4 hours prior to the diagnostic block and for 4 to 6 hours afterward. 7. Opioids should not be given as a "sedative" during the procedure. 8. The use of IV sedation (including other agents such as midazolam) may be grounds to negate the results of a diagnostic block, and should only be given in cases of extreme anxiety. 9. The patient should document pain relief with an instrument such as a VAS scale, emphasizing the importance of recording the maximum pain relief and maximum duration of pain. The patient should also keep medication use and activity logs to support subjective reports of better pain control. 10. Diagnostic facet blocks should not be performed in patients in whom a surgical procedure is anticipated. (Resnick, 2005) 11. Diagnostic facet blocks should not be performed in patients who have had

a previous fusion procedure at the planned injection level. In this case, the claimant's pain scores was not provided to determine pain level that would justifies a MBB. In addition, the claimant had prior facet injections and the ACOEM guidelines do not recommend invasive procedures due to their short-term benefit. No more than 2 injections should be performed and response to one is unknown nor indicated to determine benefit. The request for an MBB of L4-L5 and L5-S1 x3 is not medically necessary.