DECISION AND ORDER

This case is decided pursuant to Chapter 410 of the Texas Workers’ Compensation Act and Rules of the Division of Workers’ Compensation adopted thereunder.

ISSUE

A contested case hearing was held on March 24, 2009, to decide the following disputed issue:

Whether a preponderance of the evidence is contrary to the Independent Review Organization's decision denying the requested lumbar epidural steroid injection at the L5-S1 level.

PARTIES PRESENT

Petitioner/Claimant appeared, and was assisted by Ombudsman SH; Respondent/Carrier appeared, and was represented by Attorney RJ.

BACKGROUND INFORMATION

Claimant, a construction worker, sustained a compensable injury to his head, neck, and low back when he fell off a scaffold on _______________. He described his symptoms and treatment, specifically indicating that the L5-S1 epidural steroid injection he received in 2006 provided him with an appreciable degree of relief from the ongoing pain of his injury. Since Claimant and his health care providers considered the initial injection successful, it was proposed that Claimant receive another such injection.

The Independent Review Organization denied the requested procedure. In its denial, it cited the Official Disability Guidelines' requirement that radiculopathy be objectively documented in order to proceed with epidural steroid injections, and the lack of objective medical findings to support the contention that Claimant suffers from radiculopathy.

DISCUSSION

Section 408.021 of the Texas Labor Code provides that an employee who sustains a compensable injury is entitled to all health care reasonably required by the nature of the injury as and when needed. Section 401.011(22-a) defines health care reasonably required as “health care that is clinically appropriate and considered effective for the injured employee’s injury and provided in accordance with best practices consistent with: (A) evidence based medicine; or (B) if that evidence is not available, generally accepted standards of medical practice recognized in the medical community.” “Evidence based medicine” is further defined, by Section 401.011(18-a) as the use of the current best quality scientific and medical evidence formulated from credible scientific studies, including peer-reviewed medical literature and other current scientifically based texts, and treatment and practice guidelines in making decisions about the care of individual patients.
In accordance with the above statutory guidance, the Division of Workers' Compensation has adopted treatment guidelines by Division Rule 137.100. This rule directs health care providers to provide treatment in accordance with the current edition of the Official Disability Guidelines (ODG), and such treatment is presumed to be health care reasonably required as defined in the Texas Labor Code. Thus, the focus of any health care dispute starts with the health care set out in the ODG.

With regard to lumbar epidural steroid injections, the ODG sets forth the following:

Recommended as a possible option for short-term treatment of radicular pain (defined as pain in dermatomal distribution with corroborative findings of radiculopathy) with use in conjunction with active rehab efforts. See specific criteria for use below. Radiculopathy symptoms are generally due to herniated nucleus pulposus or spinal stenosis, although ESIs have not been found to be as beneficial a treatment for the latter condition.

**Short-term symptoms:** The American Academy of Neurology recently concluded that epidural steroid injections may lead to an improvement in radicular pain between 2 and 6 weeks following the injection, but they do not affect impairment of function or the need for surgery and do not provide long-term pain relief beyond 3 months. (Armon, 2007) Epidural steroid injection can offer short-term pain relief and use should be in conjunction with other rehab efforts, including continuing a home exercise program. There is little information on improved function or return to work. There is no high-level evidence to support the use of epidural injections of steroids, local anesthetics, and/or opioids as a treatment for acute low back pain without radiculopathy. (Benzon, 1986) (ISIS, 1999) (DePalma, 2005) (Molloy, 2005) (Wilson-MacDonald, 2005)

**Use for chronic pain:** Chronic duration of symptoms (> 6 months) has also been found to decrease success rates with a threefold decrease found in patients with symptom duration > 24 months. The ideal time of either when to initiate treatment or when treatment is no longer thought to be effective has not been determined. (Hopwood, 1993) (Cyteval, 2006) Indications for repeating ESIs in patients with chronic pain at a level previously injected (> 24 months) include a symptom-free interval or indication of a new clinical presentation at the level.

**Transforaminal approach:** Some groups suggest that there may be a preference for a transforaminal approach as the technique allows for delivery of medication at the target tissue site, and an advantage for transforaminal injections in herniated nucleus pulposus over translaminar or caudal injections has been suggested in the best available studies. (Riew, 2000) (Vad, 2002) (Young, 2007) This approach may be particularly helpful in patients with large disc herniations, foraminal stenosis, and lateral disc herniations. (Colorado, 2001) (ICSI, 2004) (McLain, 2005) (Wilson-MacDonald, 2005)

**Fluoroscopic guidance:** Fluoroscopic guidance with use of contrast is recommended for all approaches as needle misplacement may be a cause of treatment failure. (Manchikanti, 1999) (Colorado, 2001) (ICSI, 2004) (Molloy, 2005) (Young, 2007)

**Factors that decrease success:** Decreased success rates have been found in patients who are unemployed due to pain, who smoke, have had previous back surgery, have pain that is not decreased by medication, and/or evidence of substance abuse, disability or litigation. (Jamison, 1991) (Abram, 1999) Research reporting effectiveness of ESIs in the past has been contradictory, but these discrepancies are felt to have been, in part, secondary to numerous methodological flaws in the early studies, including the lack of imaging and contrast administration. Success rates also may depend on the technical skill of the interventionalist. (Carette, 1997) (Bigos, 1999) (Rozenberg, 1999) (Botwin, 2002) (Manchikanti, 2003) (CMS, 2004) (Delport, 2004) (Khot, 2004) (Buttermann, 2004) (Buttermann2, 2004) (Samanta, 2004) (Cigna, 2004) (Benzon, 2005) (Dashfield, 2005) (Arden, 2005) (Price, 2005) (Resnick, 2005) (Abdi, 2007) (Boswell, 2007) Also see Epidural steroid injections, “series of three” and Epidural steroid injections, diagnostic. ESIs may be helpful with radicular symptoms not responsive to 2 to 6 weeks of conservative therapy. (Kinkade, 2007) Epidural steroid injections are an option for short-term pain relief of persistent radiculopathy, although not for nonspecific low back pain or spinal stenosis. (Chou, 2008) As noted above, injections are recommended if they can facilitate a return to functionality (via activity & exercise). If post-injection physical therapy visits are required for instruction in these active self-performed exercise programs, these visits should be included within the overall recommendations under Physical therapy, or at least not require more than 2 additional visits to reinforce the home exercise program.

**With discectomy:** Epidural steroid administration during lumbar discectomy may reduce early neurologic impairment, pain, and convalescence and enhance recovery without increasing risks of complications. (Rasmussen, 2008)

An updated Cochrane review of injection therapies (ESIs, facets, trigger points) for low back pain concluded that there is no strong evidence for or against the use of any type of injection therapy, but it cannot be ruled out that specific subgroups of patients may respond to a specific type of injection therapy. (Staal-Cochrane, 2009) Recent
studies document a 629% increase in expenditures for ESIs, without demonstrated improvements in patient outcomes or disability rates. (Devo, 2009)

Criteria for the use of Epidural steroid injections:

Note: The purpose of ESI is to reduce pain and inflammation, thereby facilitating progress in more active treatment programs, and avoiding surgery, but this treatment alone offers no significant long-term functional benefit.

1. Radiculopathy must be documented. Objective findings on examination need to be present. For unequivocal evidence of radiculopathy, see AMA Guides, 5th Edition, page 382-383. (Andersson, 2000)
2. Initially unresponsive to conservative treatment (exercises, physical methods, NSAIDs and muscle relaxants).
3. Injections should be performed using fluoroscopy (live x-ray) and injection of contrast for guidance.
4. Diagnostic Phase: At the time of initial use of an ESI (formally referred to as the “diagnostic phase” as initial injections indicate whether success will be obtained with this treatment intervention), a maximum of one to two injections should be performed. A repeat block is not recommended if there is inadequate response to the first block (< 30% is a standard placebo response). A second block is also not indicated if the first block is accurately placed unless: (a) there is a question of the pain generator; (b) there was possibility of inaccurate placement; or (c) there is evidence of multilevel pathology. In these cases a different level or approach might be proposed. There should be an interval of at least one to two weeks between injections.
5. No more than two nerve root levels should be injected using transforaminal blocks.
6. No more than one interlaminar level should be injected at one session.
7. Therapeutic phase: If after the initial block/blocks are given (see “Diagnostic Phase” above) and found to produce pain relief of at least 50-70% pain relief for at least 6-8 weeks, additional blocks may be required. This is generally referred to as the “therapeutic phase.” Indications for repeat blocks include acute exacerbation of pain, or new onset of symptoms. The general consensus recommendation is for no more than 4 blocks per region per year. (CMS, 2004) (Boswell, 2007)
8. Repeat injections should be based on continued objective documented pain relief, decreased need for pain medications, and functional response.
9. Current research does not support a routine use of a “series-of-three” injections in either the diagnostic or therapeutic phase. We recommend no more than 2 ESI injections for the initial phase and rarely more than 2 for therapeutic treatment.
10. It is currently not recommended to perform epidural blocks on the same day of treatment as facet blocks or sacroiliac blocks or lumbar sympathetic blocks or trigger point injections as this may lead to improper diagnosis or unnecessary treatment.
11. Cervical and lumbar epidural steroid injection should not be performed on the same day. (Doing both injections on the same day could result in an excessive dose of steroids, which can be dangerous, and not worth the risk for a treatment that has no long-term benefit.)

The initial criterion for the use of an epidural steroid injection is radiculopathy documented by objective findings. Since Claimant's medical records do not demonstrate such findings, and since Claimant has presented no evidence-based medical opinion to justify a departure from the ODG, and the IRO opinion based upon the ODG, a decision in Carrier's favor is appropriate with respect to the sole issue presented for resolution herein.

Even though all the evidence presented was not discussed, it was considered. The Findings of Fact and Conclusions of Law are based on all of the evidence presented.

**FINDINGS OF FACT**

1. On ______________, Claimant was employed by (Employer).
2. On ______________, Employer subscribed to a policy of workers' compensation insurance issued by the Liberty Insurance Corporation, Carrier.
3. On ______________, Claimant sustained an injury arising out of the course and scope of his employment with Employer.
4. Carrier delivered to Claimant a single document stating the true corporate name of Carrier, and the name and street address of Carrier's registered agent, which document was admitted into evidence as Hearing Officer’s Exhibit Number 2.

5. Claimant's pain management doctor, BS, M.D., recommended that Claimant undergo an epidural steroid injection at the L5-S1 spinal level.

6. The Independent Review Organization (IRO) determined that the requested service was not reasonable and necessary health care for Claimant's compensable injury of ______________.

7. Claimant does not meet the criteria for a lumbar epidural steroid injection as set forth in the ODG.

8. An epidural steroid injection at the L5-S1 spinal level is not health care reasonably required for Claimant's compensable injury of ______________.

CONCLUSIONS OF LAW

1. The Texas Department of Insurance, Division of Workers’ Compensation, has jurisdiction to hear this case.

2. Venue is proper in the (City) Field Office.

3. The preponderance of the evidence is not contrary to the IRO's decision to the effect that an epidural steroid injection at the L5-S1 spinal level is not health care reasonably required to treat Claimant's compensable injury of ______________.

DECISION

Claimant is not entitled to an epidural steroid injection at the L5-S1 spinal level for his compensable injury of ______________.

ORDER

Carrier is not liable for the benefits at issue in this hearing. Claimant remains entitled to medical benefits for the compensable injury in accordance with §408.021.

The true corporate name of the insurance carrier is LIBERTY INSURANCE CORPORATION, and the name and address of its registered agent for service of process is

CT CORPORATION SYSTEMS
350 NORTH SAINT PAUL STREET
DALLAS, TEXAS 75201

Signed this 27th day of March, 2009.

Ellen Vannah
Hearing Officer