New York

Mid and Low Back Injury

Medical Treatment Guidelines


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A GENERAL GUIDELINE PRINCIPLES

The principles summarized in this section are key to the intended application of the New York State Medical Treatment Guidelines (MTG).

Medical Care

A.1 MEDICAL CARE

Medical care and treatment required as a result of a work-related injury should be focused on restoring functional ability required to meet the patient’s daily and work activities and return to work, while striving to restore the patient’s health to its pre-injury status in so far as is feasible.

A.2 RENDERING OF MEDICAL SERVICES

Any medical provider rendering services to a workers’ compensation patient must utilize the Treatment Guidelines as provided for with respect to all work-related injuries and/or illnesses.

A.3 POSITIVE PATIENT RESPONSE

Positive results are defined primarily as functional gains which can be objectively measured. Objective functional gains include, but are not limited to, positional tolerances, range of motion, strength, endurance, activities of daily living (ADL), cognition, psychological behavior, and efficiency/velocity measures which can be quantified. Subjective reports of pain and function should be considered and given relative weight when the pain has anatomic and physiologic correlation.

A.4 RE-EVALUATE TREATMENT

If a given treatment or modality is not producing positive results, the provider should either modify or discontinue the treatment regime. The provider should evaluate the efficacy of the treatment or modality 2 to 3 weeks after the initial visit and 3 to 4 weeks thereafter. Recognizing that treatment failure is at times attributable to an incorrect diagnosis should prompt the clinician to reconsider the diagnosis in the event of an unexpected poor response to an otherwise rational intervention.
Education

A.5 EDUCATION

Education of the patient and family, as well as the employer, insurer, policy makers and the community should be a primary emphasis in the treatment of work-related injury or illness. Practitioners should develop and implement effective educational strategies and skills. An education-based paradigm should always start with communication providing reassuring information to the patient. No treatment plan is complete without addressing issues of individual and/or group patient education as a means of facilitating self-management of symptoms and prevention of future injury.

Time Frames

A.6 DIAGNOSTIC TIME FRAMES

Diagnostic time frames for conducting diagnostic testing commence on the date of injury. Clinical judgment may substantiate the need to accelerate or decelerate the time frames discussed in this document.

A.7 TREATMENT TIME FRAMES

Treatment time frames for specific interventions commence once treatments have been initiated, not on the date of injury. Obviously, duration may be impacted by disease process and severity, patient compliance, as well as availability of services. Clinical judgment may substantiate the need to accelerate or decelerate the time frames discussed in this document.

A.8 DELAYED RECOVERY

For those patients who fail to make expected progress 6-12 weeks after an injury, reexamination in order to confirm the accuracy of the diagnosis and re-evaluation of the treatment program should be performed. Assessment for potential barriers to recovery (yellow flags/psychological issues) should be ongoing throughout the care of the patient. However, at 6-12 weeks, alternate treatment programs, including formal psychological or psychosocial evaluation, should be considered. Referrals to mental health providers (i.e.: psychology/psychiatry) for the evaluation and management of delayed recovery do not indicate or require the establishment of a psychiatric or psychological condition. The evaluation and management of delayed recovery does not require the establishment of a psychiatric or psychological claim.
Treatment Approaches

A.9  ACTIVE INTERVENTIONS

Active interventions emphasizing patient responsibility, such as therapeutic exercise and/or functional treatment, are generally emphasized over passive modalities, especially as treatment progresses. Generally, passive and palliative interventions are viewed as a means to facilitate progress in an active rehabilitation program with concomitant attainment of objective functional gains.

A.10  ACTIVE THERAPEUTIC EXERCISE PROGRAM

Active therapeutic exercise program goals should incorporate patient strength, endurance, flexibility, range of motion, sensory integration, coordination, and education as clinically indicated. This includes functional application in vocational or community settings.

A.11  DIAGNOSTIC IMAGING AND TESTING PROCEDURES

Clinical information obtained by history taking and physical examination should be the basis for selection and interpretation of imaging procedure results. All diagnostic procedures have variable specificity and sensitivity for various diagnoses.

When a diagnostic procedure, in conjunction with clinical information, provides sufficient information to establish an accurate diagnosis, a second diagnostic procedure will be redundant if it is performed only for diagnostic purposes. At the same time, a subsequent diagnostic procedure (that may be a repeat of the same procedure, when the rehabilitation physician, radiologist or surgeon documents the study was of inadequate quality to make a diagnosis) can be a complementary diagnostic procedure if the first or preceding procedures, in conjunction with clinical information, cannot provide an accurate diagnosis, and is permissible under the MTG.

It is recognized that repeat imaging studies and other tests may be warranted by the clinical course and to follow the progress of treatment in some cases. It may be of value to repeat diagnostic procedures (e.g., imaging studies) during the course of care to reassess or stage the pathology when there is progression of symptoms or findings, prior to surgical interventions and therapeutic injections when warranted, and post-operatively to follow the healing process. Regarding CT examinations, it must be recognized that repeat procedures result in an increase in cumulative radiation dose and associated risks.
A.12 SURGICAL INTERVENTIONS

Contemplation of surgery should be within the context of expected functional outcome. The concept of "cure" with respect to surgical treatment by itself is generally a misnomer. All operative interventions must be based upon positive correlation of clinical findings, clinical course and imaging and other diagnostic tests. A comprehensive assimilation of these factors must lead to a specific diagnosis with positive identification of pathologic condition(s). For surgery to be performed to treat pain, there must be clear correlation between the pain symptoms and objective evidence of its cause. In all cases, shared decision making with the patient is advised. The patient should be given the opportunity to understand the pros and cons of surgery, potential for rehabilitation as an alternative where applicable, evidence-based outcomes, and specific surgical experience.

A.13 PRE-AUTHORIZATION

All diagnostic imaging, testing procedures, non-surgical and surgical therapeutic procedures within the criteria of the Medical Treatment Guidelines and based on a correct application of the Medical Treatment Guidelines are considered authorized, with the exception of the following procedures: Lumbar Fusion, Artificial Disc Replacements, Vertebroplasty, Kyphoplasty, Electrical Bone Growth Stimulators, Spinal Cord Stimulators, Intrathecal Drug Delivery (Pain Pumps), Osteochondral Autograft, Autologous Chondrocyte Implantation, Meniscal Allograft Transplantation and Knee Arthroplasty (Total or Partial Knee Joint Replacement). These are not included on the list of pre-authorized procedures. Providers who want to perform one of these procedures must request pre-authorization from the carrier before performing the procedure.

Second or subsequent procedures (the repeat performance of a surgical procedure due to failure of, or incomplete success from the same surgical procedure performed earlier, if the Medical Treatment Guidelines do not specifically address multiple procedures) also require pre-authorization.

A.14 PERSONALITY/PSYCHOLOGICAL/PSYCHOSOCIAL EVALUATIONS

In select patients, diagnostic testing procedures may be useful when there is a discrepancy between diagnosis, signs, symptoms, clinical concerns or functional recovery. Psychological testing should provide differentiation between pre-existing depression versus injury-caused depression, as well as post-traumatic stress disorder, and other psychosocial issues that may include work or non-work-related issues when such conditions are identified in the patient.

For those patients who fail to make expected progress 6-12 weeks after an injury and whose subjective symptoms do not correlate with objective signs
and tests, reexamination in order to confirm the accuracy of the diagnosis should be made. Formal psychological or psychosocial evaluation may be considered.

A professional fluent in the primary language of the patient is strongly preferred. When such a provider is not available, services of a professional language interpreter must be provided.

Frequency: One time visit for evaluation. If psychometric testing is indicated by findings in the initial evaluation, time for such testing should not exceed an additional two hours of professional time.

A.15 PERSONALITY/PSYCHOLOGICAL/PSYCHOSOCIAL INTERVENTION

Following psychosocial evaluation, when intervention is recommended, such intervention should be implemented as soon as possible. This can be used alone or in conjunction with other treatment modalities.

- Time to produce effect: 2 to 8 weeks.
- Optimum duration: 6 weeks to 3 months.
- Maximum duration: 3 to 6 months. Counseling is not intended to delay but to enhance functional recovery. For select patients, longer supervision may be required, and if further counseling is indicated, documentation of the nature of the psychological factors, as well as projecting a realistic functional prognosis, should be provided by the authorized treating practitioner every 4 to 6 weeks during treatment.

Return to Work

A.16 FUNCTIONAL CAPACITY EVALUATION (FCE)

Functional capacity evaluation is a comprehensive or more restricted evaluation of the various aspects of function as they relate to the patient’s ability to return to work. Areas such as endurance, lifting (dynamic and static), postural tolerance, specific range-of-motion, coordination and strength, worker habits, employability, as well as psychosocial, cognitive, and sensory perceptual aspects of competitive employment may be evaluated. Components of this evaluation may include: (a) musculoskeletal screen; (b) cardiovascular profile/aerobic capacity; (c) coordination; (d) lift/carrying analysis; (e) job-specific activity tolerance; (f) maximum voluntary effort; (g) pain assessment/psychological screening; (h) non-material and material handling activities; (i) cognitive; (j) visual; and (k) sensory perceptual factors.

In most cases, the question of whether a patient can return to work can be answered without an FCE.
When an FCE is being used to determine return to a specific job site, the treating physician is responsible for understanding and considering the job duties. FCEs cannot be used in isolation to determine work restrictions. The authorized treating physician must interpret the FCE in light of the individual patient’s presentation and medical and personal perceptions. FCEs should not be used as the sole criteria to diagnose malingering.

An FCE may be considered at time of MMI, following reasonable prior attempts to return to full duty throughout course of treatment, when the treating physician is unable to make a clear determination on work status on case closure.

A.17 RETURN TO WORK

For purposes of these guidelines, return to work is defined as any work or duty that the patient is able to perform safely. It may not be the patient’s regular work. Ascertaining a return to work status is part of medical care, and should be included in the treatment and rehabilitation plan. It is normally addressed at every outpatient visit. A description of the patient’s status and task limitations is part of any treatment plan and should provide the basis for restriction of work activities when warranted. Early return to work should be a prime goal in treating occupational injuries. The emphasis within these guidelines is to move patients along a continuum of care and return to work, since the prognosis of returning an injured worker to work drops progressively the longer the worker has been out of work.

A.18 JOB SITE EVALUATION

The treating physician may communicate with the employer or the employer’s designee, either in person or by telephone, to obtain information regarding the demands of the patient’s pre-injury job, including a description of the exertional demands of the job, the need for repetitive activities, load lifting, static or awkward postures, or any other factors that would pose a risk of re-injury or impedance of convalescence. When returning to work at the patient’s previous job task/setting is not feasible, given the clinically determined restrictions on the patient’s activities, inquiry should also be made about modified duty work settings, and a similar set of questions should be posed by the physician about work activities/demands in modified duty jobs.

Ideally, the physician would gain the most information from an on-site inspection of the job settings and activities; but it is recognized that this may not be feasible in most cases. If job videos/CDs/DVDs are available from the employer, these can contribute valuable information.

Frequency: 1 or 2 calls
• 1st call: Patient is in a functional state where the patient can perform some work.

• 2nd call: Patient has advanced to state where the patient is capable of enhanced functional demands in a work environment.

The physician shall document the conversation.

Other

A.19 GUIDELINE RECOMMENDATIONS AND MEDICAL EVIDENCE

The Workers’ Compensation Board and its Medical Advisory Committee have not independently evaluated or vetted the scientific medical literature used in support of the guidelines, but have relied on the methodology used by the developers of various guidelines utilized and referenced in these Guidelines.

A.20 EXPERIMENTAL/INVESTIGATIONAL TREATMENT

Medical treatment that is experimental/investigational and not approved for any purpose, application or indication by the FDA is not permitted under these Guidelines.

A.21 INJURED WORKERS AS PATIENTS

In these Guidelines, injured workers are referred to as patients recognizing that in certain circumstances there is no doctor-patient relationship.

A.22 SCOPE OF PRACTICE

These Guidelines do not address scope of practice or change the scope of practice.
B  INTRODUCTION

B.1  HISTORY TAKING AND PHYSICAL EXAMINATION

History taking and physical examination establish the foundation/basis for and dictate subsequent stages of diagnostic and therapeutic procedures. When findings of clinical evaluations and those of other diagnostic procedures are not consistent with each other, the objective clinical findings have greater weight. The medical records should reasonably document the following:

B.1.a  History of Present Illness

A detailed history, taken in temporal proximity to the time of injury, should primarily guide evaluation and treatment. The history should include:

B.1.a.i  Mechanism of Injury: This includes details of symptom onset and progression. The mechanism of injury should include a detailed description of the incident and the position of the body before, during, and at the end of the incident. Inclusion of work body postures, frequency during the workday and lifting/push/pull requirements should be included in the absence of a known specific incident.

B.1.a.ii  Location of pain, nature of symptoms, and alleviating/exacerbating factors (e.g. sitting tolerance). The history should include both the primary and secondary complaints (e.g., primary back pain, secondary hip, groin pain).

B.1.a.iii  The use of an accepted pain assessment tool, (e.g. the Visual Analog Scale [VAS]) is highly recommended, especially during the first two weeks following injury, to assure that all work-related symptoms, including pain, are being addressed.

B.1.a.iv  Presence and distribution of lower extremity numbness, paresthesias, or weakness, especially if precipitated or worsened by coughing or sneezing.

B.1.a.v  Alteration in bowel, bladder or sexual function.

B.1.a.vi  Prior occupational and non-occupational injuries to the same area including specific prior treatment, history of specific prior motor vehicle accidents, chronic or recurrent symptoms, and any functional limitations.
B.1.a.vii  History of emotional and/or psychological reactions to the current injury/illness.

B.1.a.viii  Ability to perform job duties and activities of daily living.

B.1.b  **Past History**

B.1.b.i  Comprehensive past medical history.

B.1.b.ii  Review of systems includes symptoms of rheumatologic, neurologic, endocrine, neoplastic, infectious, and other systemic diseases.

B.1.b.iii  Smoking history.

B.1.b.iv  Vocational and recreational pursuits.

B.1.b.v  History of depression, anxiety, or other psychiatric illness.

B.1.c  **Physical Examination**

Guided by the medical history, should include accepted tests and exam techniques applicable to the area being examined, including:

B.1.c.i  Vital signs;

B.1.c.ii  General inspection, including posture, stance and gait;

B.1.c.iii  Visual inspection;

B.1.c.iv  Palpation;

B.1.c.v  Lumbar range of motion, quality of motion, and presence of muscle spasm. Motion evaluation of specific joints may be indicated.

B.1.c.vi  Examination of thoracic spine and pelvis;

B.1.c.vii  Nerve tension testing. When the Lassegue Test (Straight Leg Raise test) is performed, a result is generally not considered to be positive at an elevation less than 25 or greater than 60 degrees (and degrees should always be reported).

B.1.c.viii  Sensory and motor examination of the lower extremities with specific nerve root focus.

B.1.c.ix  Deep tendon reflexes.
B.1.c.x If applicable, abdominal examination, vascular examination, circumferential lower extremity measurements, or evaluation of hip or other lower extremity abnormalities.

B.1.d **Spinal Cord Evaluation**

In cases where the mechanism of injury, history, or clinical presentation suggests a possible severe injury, additional evaluation is indicated. A full neurological examination for possible spinal cord injury may include:

B.1.d.i Sharp and light touch, deep pressure, temperature, and proprioceptive sensory function;

B.1.d.ii Strength testing;

B.1.d.iii Anal sphincter tone and/or perianal sensation;

B.1.d.iv Presence of pathological reflexes.

B.1.d.v Spinal cord lesions should be classified according to the American Spine Injury Association (ASIA) impairment scale.

<table>
<thead>
<tr>
<th>ASIA IMPAIRMENT SCALE</th>
</tr>
</thead>
<tbody>
<tr>
<td>A=Complete: No motor or sensory function is preserved in the sacral segments S4-S5</td>
</tr>
<tr>
<td>B=Incomplete: Sensory but not motor function is preserved below the neurological level and includes the sacral segments S4-S5</td>
</tr>
<tr>
<td>C=Incomplete: Motor function is preserved below the neurological level, and more than half of key muscles below the neurological level have a muscle grade less than 3</td>
</tr>
<tr>
<td>D=Incomplete: Motor function is preserved below the neurological level, and at least half of key muscles below the neurological level have a grade of 3 or more</td>
</tr>
<tr>
<td>E= Normal: Motor and sensory function are normal</td>
</tr>
</tbody>
</table>
A worksheet which details dermatomes and muscle testing required is available from ASIA.

**B.1.e Relationship to Work**

This includes a statement of the probability that the illness or injury is work-related. If further information is necessary to determine work-relatedness, the physician should clearly state what additional diagnostic tests or job information is required.

**B.1.f Red Flags**

Certain findings, “red flags,” raise suspicion of potentially serious and urgent medical conditions. Assessment (history and physical examination) should include evaluation for red flags. In the mid and low back, these findings or indicators may include: acute fractures, dislocations, infection, tumor, progressive neurologic deficit or cauda equina syndrome, and extraspinal disorders. Further evaluation/consultation or urgent/emergent intervention may be indicated, and the *New York Mid and Low Back Injury Guidelines* incorporate changes in clinical management triggered by the presence of “red flags.”

**B.2 Imaging/Anatomical Tests**

Imaging studies should not be routinely performed without indications.

Physicians should be aware that “abnormal” findings on x-rays, magnetic resonance images, and other diagnostic tests are frequently seen by age 40 even in asymptomatic individuals. Bulging discs continue to increase after that point and by approximately age 60, will be encountered in a majority of patients. This requires that a careful history and physical examination be conducted by a physician in order to correlate historical, clinical, and imaging findings prior to diagnosing and attributing a patient’s complaints to the finding on imaging. The focus of treatment should be improving symptoms and function, and not the correction of abnormalities on imaging studies.

**B.3 Laboratory Testing**

Laboratory tests are rarely indicated at the time of initial evaluation, unless there is a suspicion of systemic illness, infection, neoplasia or underlying rheumatologic disorder, connective tissue disorder, or other findings based on history and/or physical examination. Laboratory tests can provide useful diagnostic information. Tests include, but are not limited to:
B.3.a **Complete Blood Count (CBC)**

Complete blood count (CBC) with differential can detect infection, blood dyscrasias, and medication side effects.

B.3.b **Rheumatologic, Infection or Connective Tissue Disorder**

Erythrocyte sedimentation rate (ESR), rheumatoid factor (RF), anti-nuclear antigen (ANA), human leukocyte antigen (HLA), and C-reactive protein (CRP), among others, can be used to detect evidence of a rheumatologic, infection, or connective tissue disorder.

B.3.c **Metabolic Bone Disease**

Serum calcium, phosphorous, uric acid, alkaline phosphatase, and acid phosphatase can detect metabolic bone disease.

B.3.d **Liver and Kidney Function**

Liver and kidney function may be performed for prolonged anti-inflammatory use or other medications requiring monitoring.
B.4 FOLLOW-UP DIAGNOSTIC IMAGING AND TESTING PROCEDURES

One diagnostic imaging procedure may provide the same or distinctive information as does another procedure. Therefore, prudent choice of a single diagnostic procedure, a complement of procedures, or a sequence of procedures will optimize diagnostic accuracy, and maximize cost effectiveness (by avoiding redundancy), and minimize potential adverse effects to patients.

All imaging procedures have a degree of specificity and sensitivity for various diagnoses. No isolated imaging test can assure a correct diagnosis. Clinical information obtained by history taking and physical examination should form the basis for selecting an imaging procedure and interpreting its results.

Magnetic resonance imaging (MRI), myelography, or computed axial tomography (CT) scanning following myelography may provide useful information for many spinal disorders. When a diagnostic procedure, in conjunction with clinical information, provides sufficient information to establish an accurate diagnosis, the second diagnostic procedure will be redundant if it is performed only for diagnostic purposes. At the same time, a subsequent diagnostic procedure (that may be a repeat of the same procedure, when the rehabilitation physician, radiologist or surgeon documents that the study was of inadequate quality to make a diagnosis) can be a complementary diagnostic procedure if the first or preceding procedures, in conjunction with clinical information, cannot provide an accurate diagnosis. Usually, preference of a procedure over others depends upon availability, a patient’s tolerance, and/or the treating practitioner’s familiarity with the procedure.

It is recognized that repeat imaging studies and other tests may be warranted by the clinical course and to follow the progress of treatment in some cases. It may be of value to repeat diagnostic procedures (e.g. imaging studies) during the course of care to reassess or stage the pathology when there is progression of symptoms or findings, prior to surgical interventions and therapeutic injections when warranted, and post-operatively to follow the healing process. Regarding CT examinations, it must be recognized that repeat procedures result in an increase in cumulative radiation dose and associated risks.
C DIAGNOSTIC STUDIES

C.1 IMAGING STUDIES

C.1.a Roentgenograms (X-Rays)

Recommendations:

C.1.a.i Routine x-rays are not recommended for acute non-specific back pain. In the absence of red flags (indicators of potentially serious disease, such as fever or major trauma), imaging tests are not recommended in the first 4-6 weeks of back pain symptoms.

C.1.a.ii X-rays are recommended for acute back pain with red flags for fracture or serious systemic illness, back pain that is not improving, or non-acute back pain, as an option to rule out other possible conditions.

C.1.a.iii X-rays are an option to rule out other possible conditions. If MRI is used as imaging, plain x-ray may not be needed. MRI is a more sensitive and more specific test, which is much more costly, but which avoids gonadal radiation, which for those still in the age group to potentially reproduce is a significant consideration.

Frequency/Duration: Obtaining x-rays once is generally sufficient. For patients with non-acute back pain, it may be reasonable to obtain a second set months or years subsequently to re-evaluate the patient’s condition, particularly if symptoms change.

C.1.a.iv Flexion and extension views are recommended:

For evaluating symptomatic spondylolisthesis in which there is consideration for surgery or other invasive treatment or occasionally in the setting of trauma.

- Frequency/Duration: Obtaining flexion and extension and lateral flexion and extension views are generally needed no more frequently than every few years, in the absence of a rapidly changing clinical course.
C.1.b  **Magnetic Resonance Imaging (MRI)**

MRI is considered the gold standard in diagnostic imaging for defining anatomy because it has the greatest resolution of any test currently available. While CT remains an important analytical tool especially for evaluating bony or calcified structures of the spine, due to the greater resolution of MRI, particularly with respect to soft tissue of the spine (nerve root compression, myelopathy to evaluate the spinal cord and/or differentiate/rule out masses), there is less need for using CT at the current time. Ferrous material/metallic objects in tissue is a contraindication for the performance of an MRI.

Inadequate resolution on the first scan may require a second MRI using a different technique. A subsequent diagnostic MRI may be a repeat of the same procedure when the rehabilitation physician, radiologist or surgeon documents that the study was of inadequate quality to make a diagnosis. All questions in this regard should be discussed with the MRI center and/or radiologist.

Recommendations:

C.1.b.i MRI is not recommended for acute back pain or acute radicular pain syndromes in the first 6 weeks, in the absence of red flags.

C.1.b.ii MRI is recommended for patients with acute back pain during the first 6 weeks if they have demonstrated progressive neurologic deficit, cauda equina syndrome, significant trauma with no improvement in atypical symptoms, a history of neoplasia (cancer), or atypical presentation (e.g., clinical picture suggests multiple nerve root involvement).

C.1.b.iii MRI is recommended for acute radicular pain syndromes in the first 6 weeks if the symptoms are severe and not trending towards improvement and both the patient and the physician are willing to consider prompt surgical treatment, assuming the MRI confirms ongoing nerve root compression.

- Frequency/Duration: Repeat MRI imaging without significant clinical deterioration in symptoms and/or signs is not recommended.

C.1.b.iv MRI is recommended for patients with non-acute radicular pain syndromes lasting at least 6 weeks, in whom the symptoms are not trending towards improvement, if both
the patient and surgeon are considering prompt surgical treatment, assuming the MRI confirms ongoing nerve root compression.

C.1.b.v In cases where an epidural glucocorticosteroid injection is being considered for temporary relief of acute or subacute radiculopathy, MRI at 3 to 4 weeks (before the epidural steroid injection) may be reasonable (see Section D.6, Injections: Therapeutic).

C.1.b.vi MRI is recommended as an option for the evaluation of select non-acute back pain patients in order to rule out concurrent pathology unrelated to injury. This should rarely be considered before 3 months and failure of several treatment modalities (including NSAIDs, aerobic exercise, other exercise, and considerations for manipulation, and/or acupuncture).

C.1.b.vii Standing or weight-bearing MRI is not indicated for any back or radicular pain syndrome or condition. In the absence of studies demonstrating improved patient outcomes, this technology is currently considered experimental/ investigational.

C.1.c  **Computerized Tomography (CT)**

Due to the far greater resolution of MRIs, particularly with respect to the soft tissue structures of the spine, there is much less need for CT at the current time. However, CT remains a good test to evaluate bony or calcified structures of the spine. CT is most useful to evaluate the spine in patients with contraindications for MRI (most typically an implanted metallic-ferrous device). CT is not invasive (minimally invasive when contrast is needed), has low potential adverse effects, but is costly and entails radiation exposure.

Recommendations:

C.1.c.i Routine CT is not recommended for acute or non-acute non-specific back pain or for radicular pain syndromes.

C.1.c.ii CT (or MRI) is recommended for those with radicular pain syndrome that has failed to improve within 4 to 6 weeks and there is consideration for an epidural glucocorticoid injection or surgical discectomy (see Section D.6, Injections: Therapeutic).
C.1.c.iii  CT is useful in patients with an indication for MRI who cannot undergo MRI examination due to contraindications such as implanted metallic-ferrous device or significant claustrophobia.

Frequency/Duration: Obtaining serial CT exams is not recommended, although if there has been a significant worsening in the patient’s history of examination, repeat imaging may be warranted.

C.1.d  **Myelography (Including CT Myelography and MRI Myelography)**

Myelography is invasive, has complications and is costly. It has almost entirely been replaced by MRI and other imaging procedures.

Recommendations:

C.1.d.i  Myelography (as well as CT myelography and MRI myelography) is not recommended as the first diagnostic study for the diagnosis of lumbar root compromise.

C.1.d.ii  Myelography, including CT myelography, is recommended only in uncommon specific situations (e.g., implanted metal that precludes MRI, equivocal findings of disc herniation on MRI suspected of being falsely positive, spinal stenosis, and/or a post-surgical situation that requires myelography).

C.1.e  **Bone Scans**

Recommendations:

C.1.e.i  Bone scanning is not recommended for routine use in back pain patients.

C.1.e.ii  Bone scanning is a good diagnostic test for specific situations which involve a minority of patients and may be useful in diagnosing suspected metastases, infection (osteomyelitis), inflammatory arthropathies and fractures.

This technology is generally not used for evaluation of most occupational back pain situations.
C.1.f **Fluoroscopy**

Recommendations:

C.1.f.i Fluoroscopy is not recommended for the evaluation of acute or non-acute back pain.

C.1.g **Single Proton Emission Computed Tomography (SPECT)**

Recommendations:

C.1.g.i SPECT is not recommended, and aside from cases of suspected inflammatory arthropathies not diagnosed by more common tests, there is no current evidence that it has a role in the evaluation of patients with back pain and related disorders.

C.1.h **Ultrasound (Diagnostic)**

Recommendations:

C.1.h.i Diagnostic ultrasound is not recommended for patients with back pain.

C.1.i **Videofluoroscopy**

Recommendations:

C.1.i.i Videofluoroscopy is not recommended for the assessment of acute or non-acute back pain.

C.2 **OTHER TESTS/PROCEDURES:**

C.2.a **Electrodiagnostic Studies (EDX)**

EDX include needle EMG, peripheral nerve conduction velocity studies (NCV) and motor and sensory evoked potentials. Needle EMG is usually what substantiates the diagnosis of radiculopathy or spinal stenosis in patients with back pain and/or radiculopathy problems. Needle EMG can help determine if radiculopathy is acute or chronic. NCV are done in addition to needle EMG to rule out other potential causes for the symptoms (co-morbidity or alternate diagnosis involving peripheral nerves) and to confirm radiculopathy. It is recommended and preferred that EDX in the out-patient setting be performed and interpreted by physicians board-certified in Neurology or Physical Medicine and Rehabilitation.
Recommendations:

C.2.a.i EDX are not recommended for patients with acute or non-acute back pain who do not have significant leg pain or numbness.

C.2.a.ii EDX (must include needle EMG and NCV) are recommended where a CT or MRI is equivocal and there are ongoing complaints of pain, weakness, and/or numbness/parasthesias that raise questions about whether there may be a neurological compromise that may be identifiable. This means leg symptoms consistent with radiculopathy, spinal stenosis, peripheral neuropathy, etc.

Nerve conduction studies are done in addition to the needle EMG both to rule out other potential causes for the symptoms (co-morbidity or alternate diagnosis involving peripheral nerves, e.g. compression neuropathies) and to confirm radiculopathy, but the testing must include needle EMG.

C.2.a.iii EDX is recommended where there is failure of suspected radicular pain to resolve or plateau after waiting 4 to 6 weeks (to provide for sufficient time to develop EMG abnormalities as well as time for conservative treatment to resolve the problems), equivocal imaging findings, e.g. on CT or MRI studies, and suspicion by history and physical examination that a neurologic condition other than radiculopathy may be present instead of or in addition to radiculopathy.

C.2.b Surface Electromyography (Surface EMG)

Recommendations:

C.2.b.i There is no established indication for the use of surface EMG in back pain diagnosis and it is not recommended.

Surface EMG may be of use in biofeedback training and gait analysis for neurologic disorders, but it has no established use in any adult back pain scenario.

C.2.c Diagnostic Facet Blocks

See Section D.6.f.
C.2.d  **Lumbar Discography**

Recommendation

Discography, whether performed as a solitary test or when paired with imaging (e.g., MRI), is not recommended for acute or non-acute back pain or radicular pain syndromes. Improvement in surgical outcomes has not been shown to follow the use of discography, and there is evidence that performing discography on normal discs is associated with an enhanced risk of degenerative changes in those discs in later years.

C.2.e  **CT/MRI Discography**

Recommendations: See Lumbar Discography above.

C.2.f  **Myeloscopy**

Recommendations:

Myeloscopy is not recommended for acute or non-acute back pain, spinal stenosis, radicular pain syndromes or post-surgical back pain problems.

C.2.g  **Thermography**

Recommendations:

Thermography is not recommended for the assessment of acute or non-acute back pain, or radicular pain patients.
D THERAPEUTIC PROCEDURES: NON-OPERATIVE

Before initiation of any therapeutic procedure, the authorized treating provider, employer, and insurer must consider these important issues in the care of the patient.

First, patients undergoing therapeutic procedure(s) should be released or returned to modified or restricted duty during their rehabilitation at the earliest appropriate time.

Second, cessation and/or review of treatment modalities should be undertaken when no further significant subjective or objective improvement in the patient’s condition is noted. If patients are not responding within the recommended duration periods, alternative treatment interventions, further diagnostic studies or consultations should be pursued.

Third, providers should provide and document education to the patient. No treatment plan is complete without addressing issues of individual and/or group patient education as a means of facilitating self-management of symptoms.

Lastly, for those patients who fail to make expected progress 6-12 weeks after an injury and whose subjective symptoms do not correlate with objective signs and tests, reexamination in order to confirm the accuracy of the diagnosis should be made. Formal psychological or psychosocial evaluation may be considered.

Home therapy is an important component of therapy and may include active and passive therapeutic procedures as well as other modalities to assist in alleviating pain, swelling, and abnormal muscle tone.

The following are listed in alphabetical order.

D.1 ACUPUNCTURE

Recommendations:

D.1.a.i Routine use of acupuncture is not recommended for acute back pain or radicular pain. Although it is not high cost and its use is not associated with high potential for patient harm, it is not recommended.

D.1.a.ii Acupuncture is recommended for select use in non-acute back pain as an adjunct to more efficacious treatments.

D.1.a.iii Acupuncture may be recommended as treatment of non-acute back pain as a limited course during which time there are clear objective and functional goals that are to be achieved.
Consideration for time-limited use in non-acute back pain patients without underlying serious pathology is as an adjunct to a conditioning program that has both graded aerobic exercise and strengthening exercises. Acupuncture is only recommended to assist in increasing functional activity levels more rapidly and the primary attention should remain on the conditioning program.

This intervention is not recommended for patients not involved in a conditioning program, or who are non-compliant with graded increases in activity levels.

Frequency/Duration:

a) There are different patterns which are used in quality studies. These range from weekly for a month to 20 appointments over 6 months; however the norm is generally no more than 8 to 12 sessions.

b) An initial trial of 5 to 6 appointments would appear reasonable in combination with a conditioning program of aerobic and strengthening exercises.

c) Future appointments should be tied to improvements in objective measures and would justify an additional 6 sessions, for a total of 12 sessions.

Discontinuation: Resolution, intolerance, or non-compliance, including non-compliance with aerobic and strengthening exercises.

D.2 APPLIANCES

Include: shoe insoles, shoe lifts, kinesiotaping and taping, lumbar supports, magnets, mattresses and sleeping surfaces.

D.2.a Shoe Insoles and Shoe Lifts

Recommendations:

D.2.a.i These interventions are recommended for the treatment of acute or non-acute back pain or radicular pain syndrome in the presence of significant leg length discrepancy.

In the absence of significant leg length discrepancy, these are not recommended.
D.2.b  **Kinesiotaping, Taping or Strapping**

Recommendations:

D.2.b.i Other than for acute joint immobilization (for example, acute ankle sprain), kinesiotaping, taping or strapping are not recommended for acute or non-acute pain.

D.2.c  **Lumbar Supports**

Recommendations:

D.2.c.i Lumbar supports may be useful for specific treatment of spondylolisthesis, documented instability, or post-operative treatment.

D.2.c.ii Lumbar supports are not recommended for the prevention or treatment of other back pain conditions.

D.2.d  **Magnets**

Recommendations:

D.2.d.i The use of magnets is not recommended.

D.2.e  **Mattresses, Water Beds, and Sleeping Surfaces**

Recommendations:

D.2.e.i It is recommended that patients select mattresses, pillows, bedding, or other sleeping options that are most comfortable for them.

D.2.e.ii There is no recommendation regarding mattresses other than that providers should be aware that ordering patients to sleep on firm mattresses or on the floor may be incorrect.

D.2.e.iii There is no quality evidence to guide recommendations regarding other optimal sleeping surfaces (e.g., bedding, water beds, and hammocks).
D.3  BED REST

Recommendations:

D.3.a.i Bed rest is not recommended for the management of acute or non-acute back pain, radicular pain syndromes including sciatica or other back pain-related problems including spondylolisthesis, spondylolysis, spinal stenosis, facet-related pain, or pain thought to be related to the sacroiliac joint.

There is no quality evidence that these conditions are successfully treated with bed rest and there are also likely adverse effects. Although it is non-invasive, it is costly, has no documented benefits and is expected to be associated with higher morbidity.

D.3.a.ii Bed rest is recommended in the management of unstable spinal fractures.

Although there are no quality studies evaluating the role of bed rest in the management of unstable spinal fractures or cauda equina syndrome, there is consensus that these require bed rest or other marked activity limitations to prevent adverse events. Although bed rest is costly and has no documented benefits, the hazard of mobilization in this setting is theoretically catastrophic, thus this treatment strategy is recommended.

D.4  BIOFEEDBACK

Recommendations:

D.4.a.i Biofeedback is not recommended in patients with acute back pain. It is suggested that other treatments for which there is quality evidence of efficacy are more appropriate.

D.4.a.ii Biofeedback is recommended for select patients with non-acute back pain, as a component of an interdisciplinary approach. Please consult the New York Non-Acute Pain Medical Treatment Guidelines for further recommendations.
D.5 ELECTRICAL THERAPIES

D.5.a Interferential Therapy

Recommendations:

D.5.a.i Interferential therapy is not recommended for treatment of acute or non-acute back pain, non-acute radicular pain syndromes, or other back-related conditions.

D.5.b Transcutaneous Electrical Neurostimulation (TENS)

Recommendations:

D.5.b.i Transcutaneous Electrical Nerve Stimulation (TENS) treatment should include at least one instructional session for proper application and use. Indications include muscle spasm, atrophy and control of concomitant pain in the office setting. Minimal TENS unit parameters should include a pulse rate, pulse width and amplitude modulation. Consistent, measurable, functional improvement must be documented and a determination made of the likelihood of chronicity prior to the provision of a home unit. TENS treatment should be used in conjunction with active physical therapy.

- Time to Produce Effect: Immediate.
- Frequency: Variable.
- Optimum Duration: 3 sessions.
- Maximum Duration: 3 sessions. Purchase or provide with home unit if effective.

D.5.c Percutaneous Electrical Nerve Stimulation (PENS)

Recommendations:

D.5.c.i PENS is not recommended for acute or subacute back pain or radicular pain syndromes.

D.5.c.ii As PENS is still an investigational treatment, it is not recommended outside of research settings for non-acute non-radicular back pain.
D.5.d **Microcurrent Electrical Stimulation**

Recommendations:

D.5.d.i Microcurrent electrical stimulation is not recommended for acute or non-acute back pain or radicular pain syndrome patients, as other therapies are believed to be more efficacious and less costly.

D.5.e **Electrical Nerve Block**

Electrical Nerve Block is not recommended.

D.5.f **Electrical Stimulation (Physician or Therapist Applied)**

Electrical Stimulation (like other passive modalities) is not recommended as a stand-alone treatment, but may be a component of a comprehensive treatment plan.

Frequency: 2-3 x week for a maximum of up to two months.

D.5.g **Transcutaneous Neurostimulator (TCNS)**

TCNS is not recommended.

D.5.h **H-Wave Stimulation**

Recommendations:

D.5.h.i H-wave stimulation is not recommended for acute or non-acute back pain or radicular pain syndromes.

D.5.i **High-Voltage Galvanic**

Recommendations:

D.5.i.i High-voltage galvanic is not recommended for the treatment of acute or non-acute back pain or radicular pain syndromes or other back-related conditions.

D.5.j **Iontophoresis**

Recommendations:

D.5.j.i Iontophoresis is not recommended for the treatment of acute or non-acute back pain or radicular pain syndromes or other back-related conditions.
D.6 INJECTIONS: THERAPEUTIC

D.6.a Therapeutic Spinal Injections-Introduction

Description:

Therapeutic spinal injections may be used after initial conservative treatments, such as physical and occupational therapy, medication, manual therapy, exercise, acupuncture, have been undertaken.

- Therapeutic injections should be used only after imaging studies and diagnostic injections have established pathology.

- Injections are invasive procedures that can cause catastrophic complications; thus clinical indications and contraindications should be closely adhered to.

- The purpose of spinal injections is to facilitate active therapy by providing short-term relief through reduction of pain and inflammation.

- All patients should continue appropriate exercise with functionally directed rehabilitation.

- Active treatment, which patients should have had prior to injections, will frequently require a repeat of the sessions previously ordered.

- Injections, by themselves, are not likely to provide long-term relief. Rather, active rehabilitation with modified work achieves long-term relief by increasing active range of motion, strength, and stability.

- Injections should not be repeated if the first injection does not provide:
  - Improvement in function
  - Temporary and sustained pain relief as measured by accepted pain scales, i.e., 50% pain reduction on Visual Analog Scale
  - and/or
  - Reduction in the use of prescribed analgesic medication.
• Medical management should be continued or adjusted based upon patient assessment and response.

Special Considerations:

• For all injections (excluding trigger point) multi-planar fluoroscopy during procedures is required to document technique and needle placement.

• All injections (excluding trigger point) must be performed by a physician experienced in the procedure. Trigger point injections may be performed by a physician or a Nurse Practitioner/Physician Assistant experienced in the procedure.

• Permanent images are required to verify needle placement.

• The subspecialty disciplines of the physicians performing injections may be varied, including, but not limited to: anesthesiology, radiology, surgery, or physiatry.

• The practitioner should have completed fellowship training in pain medicine with interventional training or its equivalent. The practitioner must also be knowledgeable in radiation safety.

Complications:

General complications of spinal injections may include:

• transient neurapraxia
• local pain
• nerve injury
• infection
• headache
• vasovagal effects
• Epidural hematoma, permanent neurologic damage, dural perforation and CSF leakage, and/or spinal meningeal abscess may also occur.

• More serious complications are rare but can include spinal cord damage, quadriplegia, permanent ataxia, and death.
• With steroid injections, there may be a dose-dependent suppression of the hypothalamic-pituitary-adrenal axis lasting between one and three months.

Contraindications:

Absolute contraindications to therapeutic injections include:

• bacterial infection, systemic or localized to region of injection
• bleeding diatheses
• hematological conditions
• possible pregnancy

Relative contraindications to diagnostic injections may include:

• allergy to contrast
• poorly controlled Diabetes Mellitus
• hypertension

Drugs affecting coagulation, such as aspirin, NSAIDs, anti-platelets or anticoagulants require restriction from use.

• Decisions regarding the number of restricted days before a procedure should be made in consultation with the prescribing physician and other specialists as indicated.

D.6.b Lumbar Transforaminal/Interlaminar/Caudal Epidural Injections (ESI)

Description:

The purpose of ESI is to reduce pain and inflammation, restoring range of motion and thereby facilitating progress in more active programs.

As with all treatments, it is important to insure that patients have realistic expectations regarding treatment outcomes.

Diabetics who are candidates for ESI should be counseled that a blood glucose increase may be apparent post-intervention, but effects should not last longer than approximately two days.
Needle Placement: Lumbar ESI must be fluoroscopically guided to verify needle placement. Permanent images are required to verify needle placement.

Contrast epidurograms allow one to verify the flow of medication into the epidural space. One epidurogram per series of ESI is recommended as clinically indicated.

Recommendations:

- ESI is useful in patients with symptoms of lumbar radicular pain syndromes.
- ESI is not effective for lumbar axial pain or non-radicular pain syndromes and they are not recommended for this indication.
- Medications for ESI include anesthetics and/or steroids.
- ESI is an option for acute radicular pain syndromes.
- ESI is an option for radicular pain syndromes lasting at least 3 weeks having been treated with NSAIDs and without evidence of trending toward spontaneous resolution.
- ESI is an option for second-line treatment for acute flare-ups of spinal stenosis.
- ESI is an option for symptoms of spinal stenosis of at least 1 to 2 months, with prior treatment that has included NSAIDs and progressive exercise.
- ESI is not recommended for acute or non-acute back pain in the absence of significant radicular symptoms.
- ESI is also not recommended as first or second line treatment in individuals with back pain symptoms that predominate over leg pain.
- ESI is not recommended as treatment for any non-acute axial back pain without a radicular component.

Maximum Frequency:

- Three injections (per spinal region) may be done in one 12-month period, depending on patient response (improved function and pain reduction). No more than 2 levels per treatment session.
It is recommended that each injection be scheduled separately and effects of each injection be evaluated, depending upon patient response (improved function and pain reduction) rather than scheduling a “Series of Three.”

If the first injection does not provide a response with temporary and sustained pain relief (at least 2 weeks) substantiated by accepted pain scales (i.e. 50% pain reduction as measured by tools such as VAS) and improvement in function, repeat injections are not recommended.

A positive result (functional improvement) should include measurable improvement in physical activity goals, and a return to baseline function or to work duties.

Patients should be reassessed after each injection for:

- Improvement in function
- Temporary and sustained pain relief as measured by accepted pain scales, i.e., 50% pain reduction on Visual Analog Scale and/or
- Reduction in the use of prescribed analgesic medication.

Medical management should be continued or adjusted based upon patient assessment and response.

Discontinuation:

- Resolution of symptoms, decrease in symptoms to a tolerable level or absence of response.

**D.6.c Intradiscal Steroids**

Recommendations:

**D.6.c.i** Intradiscal steroid injections are not recommended for the treatment of acute back pain.

There is no quality evidence on the value of intradiscal steroid injections in those with acute back pain. There is also no quality evidence that these injections improve on the natural history of acute back pain.

**D.6.c.ii** This treatment strategy is not recommended for management of non-acute back pain.
D.6.d **Chemonucleolysis (Chymopapain and Collagenase)**

This procedure, while a successful treatment, is not available in the U.S. due to serious adverse effects.

D.6.e **Tender and Trigger Point Injections**

Description:

Myofascial trigger points are localized hyperirritable palpable nodules in extremely sensitive bands of taut skeletal muscle fibers. These nodules are painful on compression and give rise to local pain and pain referred to distant structures.

- Trigger point treatment consists only of dry needling or injection of local anesthetic into myofascial trigger points.

- Trigger point injection is not the equivalent of acupuncture. Please refer to the acupuncture section in each Medical Treatment Guideline.

- There is no evidence that injection of medications improves the results of trigger point injections. Needling alone may account for some of the therapeutic response.

- As with all treatments, it is important to insure that patients have realistic expectations regarding treatment outcomes.

Recommendations:

- Trigger point injections are not recommended for treatment of acute back pain.

- Trigger point injections may be reasonable secondary or tertiary options for non-acute pain that is not resolving with more conservative means (e.g., NSAIDs, progressive aerobic exercises, other exercises) within a 6-week time frame.

- Trigger point injections should be utilized primarily for facilitating functional progress.

- Trigger point injections may be used to relieve myofascial pain and facilitate active therapy and stretching of the affected areas.
• The use of therapeutic injections without participation in an active therapy program or in the context of maintaining employment is not recommended.

• Patients should be reassessed two weeks after each injection for:
  o Improvement in function
  o Temporary and sustained pain relief as measured by accepted pain scales, i.e., 50% pain reduction on Visual Analog Scale
    and/or
  o Reduction in the use of prescribed analgesic medication.

• Medical management should be continued or adjusted based upon patient assessment and response.

• Functional improvement should last for 3 months.

• A positive result (functional improvement) should include measurable improvement in physical activity goals, and a return to baseline function or to work duties.

• The use of ultrasound or other imaging studies for trigger point injections is not recommended.

Frequency:

• Not more than 4 injections per session, not to exceed 4 sessions per 12-month period.

• If there is a partial demonstrated improvement after the first set of injections, a second set may be reasonable.

• It is recommended to allow at least 3 to 4 weeks between these injections.

• If there are no subjective and objective improvements at that point, further injections are not recommended.

• Repeated injections should be linked to subjective and objective improvement.
D.6.f  **Lumbar Diagnostic and Therapeutic Medial Nerve Branch Blocks or Intra-articular Facet (Zygapophyseal) Joint Injections**

**Indications:**

Medial branch block injections or intra-articular facet joint injections are recommended for:

- Patients with pain suspected to be largely facet in origin based on exam findings (i.e., non-radicular axial pain aggravated by extension-facet loading)
  
  and/or

- Documented evidence (i.e., imaging study) of facet disease (facet arthropathy/hypertrophy at the targeted level(s))
  
  and

- Who have completed a documented course of conservative management as defined in the *New York Mid and Low Back Injury Medical Treatment Guidelines*, including but not limited to medication, modalities, and active exercises.

**Description:**

These injections must be fluoroscopically guided.

Lumbar medial nerve branch blocks or intra-articular facet joint injections may consist of a diagnostic and/or a therapeutic component.

- The diagnostic component consists of an anesthetic and the therapeutic component, a corticosteroid.
- The diagnostic component (anesthetic only) may be used individually or may be combined with a steroid into a single diagnostic/therapeutic injection.
- A medial nerve branch block is indicated for the diagnosis of pain that is suspected of arising from the facet joint.
- A history and physical examination should document the rationale for the suspected diagnosis.
A positive response to a therapeutic injection (steroid) is not determinative of the need for radiofrequency ablation.

Recommendations:

D.6.f.i Acute back pain

Intra-articular facet joint (injection into the intra-articular facet joint space) or medial branch block injections (blocking medial nerve innervation) may be indicated for acute back pain when there is continuing axial back pain after an injury that has not responded to conservative management.

In patients with acute back pain, medial branch block injections may aid in identifying pain generators, therapeutically reduce pain and may be useful in facilitating progress in a rehabilitation program.

For acute pain, these injections involve a combination of an anesthetic and a steroid. Steroid is added to provide longer benefit. The goal of the prolonged therapeutic benefit is to decrease pain and increase function, with the ability to participate in an active rehabilitation program (which the patient was unable to do prior to the injection).

Diagnostic medial branch block injections (anesthetic only without steroids) or diagnostic intra-articular facet joint injections (anesthetic only without steroids) are not recommended for acute back pain.

Positive Therapeutic Response (either Medial Branch Block Injection or Intra-articular Facet Joint Injection):

- Patients should be reassessed after each therapeutic injection for a documented 50% improvement in pain as measured by accepted pain scales and evidence of functional improvement for at least 4-6 weeks

- A positive result (functional improvement) should include measurable improvement in physical activity goals including progress towards return to baseline function or work activities.
• Pain should be measured by accepted pain scales, pre-procedure, immediately post-procedure and at identified intervals after the procedure.

• If the first therapeutic injection does not provide sustained pain relief substantiated by accepted pain scales (i.e., 50% documented pain reduction as measured by accepted pain tools) and improvement in function for at least 4-to-6 weeks, repeat steroid injections are not recommended.

Time to produce effect: up to 72 hours.

Recommended frequency: 2 injections for each applicable joint may be done in one 12-month period, not to exceed 3 joint levels (4 medial branch nerves) per session, depending upon patient’s documented response (i.e., improved functional gain and pain reduction). Maximum 2 sessions/year.

D.6.f.ii Non-acute back pain

Diagnostic medial branch block injections are recommended for a select group of patients with non-acute back pain in order to determine whether specific interventions targeting the facet joint (by blocking medial nerve innervation to the facet joint) should be performed.

The decision to proceed with radiofrequency ablation is based on a positive response to diagnostic injections (anesthetic only or anesthetic with steroid) and not on the response to a therapeutic injection (with steroid only.)

When administering a diagnostic injection, consideration should be given to combining the anesthetic agent with steroid to allow for the potential of extended pain relief.

Positive Diagnostic Medial Nerve Branch Block Response:

A positive response to the diagnostic component of a medial nerve branch block consists of an initial temporary improvement, which may be as short as 1-4 hours, and includes a reduction in pain (50% decrease as measured by accepted pain scales), and improvement in function for the duration of the local anesthetic.
The primary goal of a diagnostic medial nerve branch block in the setting of non-acute pain is to determine the need for more definitive treatment (i.e. radiofrequency ablation).

- If a patient has a positive response to a diagnostic medial branch block injection (whether or not steroids are used), a repeat medial branch block injection should be performed to confirm the diagnosis.

- This repeat comparative medial branch block injection should be performed on a different date to confirm the level of involvement.

- If there is a positive response to the repeat diagnostic medial branch block injection, the patient should be evaluated to determine the need for more definitive treatment such as radiofrequency ablation.

- If the first injection does not provide a positive response, the diagnosis should be re-evaluated.

- If the first injection does not provide a positive response, repeat diagnostic injections are not recommended.

Time to produce effect: up to 30 minutes for local anesthetic; corticosteroid up to 72 hours.

Recommended frequency: 2 injections for each applicable joint may be done in one 12-month period, not to exceed 3 joint levels (4 medial branch nerves) per session, depending upon patient’s documented response (i.e., improved functional gain and pain reduction). Maximum 2 sessions/year.

D.6.g **Facet Joint Hyaluronic Acid Injections**

Recommendations:

D.6.g.i Are not recommended.

D.6.h **Sacroiliac Joint Injections**

Recommendations:
D.6.h.i Sacroiliac joint corticosteroid injections are recommended as a treatment option for patients with a specific cause of sacroiliitis, meaning a work-related aggravation of proven rheumatologic inflammatory arthritis involving the sacroiliac joints.

D.6.h.ii Sacroiliac joint injections are recommended for the treatment of sacroiliac joint sprain/dysfunction. Sacroiliac sprain may present with local tenderness corresponding to the anatomical sacroiliac joint. Such presentation is an extra-axial finding, without radiation, and may be the result of inflammation or trauma. The pain may be acute or non-acute.

- Frequency/Duration: If the results after the first injection are not satisfactory, fluoroscopic guidance must be used for the second injection. Subsequent injections are not recommended unless significant improvement is noted after the initial injections.

D.6.i Prolotherapy Injections

Recommendations:

D.6.i.i Prolotherapy is not recommended for acute or non-acute back pain, or for any radicular pain syndrome.

D.6.j Platelet Rich Plasma (PRP)

PRP not recommended.

D.7 MEDICATIONS

Please consult the New York Non-Acute Pain Medical Treatment Guidelines for additional recommendations on the use of medications in non-acute pain.

The following medications are listed in alphabetical order.

D.7.a Acetaminophen

Acetaminophen is an effective analgesic with antipyretic but not anti-inflammatory activity. Acetaminophen is generally well tolerated, causes little or no gastrointestinal irritation, and is not associated with ulcer formation. Acetaminophen has been associated with liver toxicity when the recommended daily dose is exceeded or in patients who chronically use alcohol. Patients may not realize that many over-the-counter preparations may contain acetaminophen. In general, the
total daily dose of acetaminophen should not exceed 3 grams per 24-hour period from all sources, including narcotic-acetaminophen combination preparations. Patients who consume three or more alcoholic drinks per day are at greater risk for liver toxicity, and consideration should be given to the use of other analgesics or limiting the acetaminophen dose to 2 grams per 24-hour period from all sources. Monitoring liver function via blood testing for use beyond 10 days is advisable.

Recommendations:

**D.7.a.i** Acetaminophen is a reasonable alternative to NSAIDs, although evidence suggests it is modestly less efficacious.

**D.7.a.ii** Acetaminophen is recommended for treatment of back pain with or without radicular symptoms, particularly for those with contraindications for NSAIDs.

- **Optimum Duration:** 7 to 10 days.
- **Maximum Duration:** Chronic use as indicated on a case-by-case basis.

**D.7.b Anti-Depressants**

Recommendations:

**D.7.b.i** Tricyclic antidepressants (TCAs) are recommended for the treatment of non-acute back pain that is not fully treated with NSAIDs and an exercise program. This intervention may be helpful where there is nocturnal sleep disruption and mild dysthymia.

- **Frequency/Duration:** Generally prescribed at a very low dose at night and gradually increased (e.g., amitriptyline 25 mg QHS, increase by 25 mg each week) until a sub-maximal or maximal dose is achieved, sufficient effects are achieved, or adverse effects occur. Most practitioners use lower doses, (e.g., amitriptyline 25 to 75 mg a day to avoid the adverse effects and necessity of blood level monitoring), as there is no evidence of increased pain relief at higher doses. Imipramine is less sedating, thus if there is carryover daytime sedation, this may be a better option.
• Discontinuation: Resolution of pain, intolerance, or development of adverse effects.

D.7.b.ii Tricyclic antidepressants (TCAs) are recommended for the treatment of radicular pain.

There is limited evidence that TCAs result in modest reductions in pain ratings in the treatment of radicular pain compared with placebo.

Recommendations regarding usage, frequency, duration and discontinuation are as above for non-acute back pain.

D.7.b.iii The selective serotonin reuptake inhibitors, (e.g., paroxetine, as well as bupropion and trazodone) are not recommended for treatment of non-acute pain. They may be recommended for the treatment of non-acute back pain with concomitant depression.

There is strong evidence that treatment with these SSRI medications is not of benefit; thus their use is not recommended for the management of non-acute back pain without depression.

Absent other indicators of a need for such treatment, this intervention is not recommended for the management of acute back pain.

D.7.c **Anti-Seizure Drugs**

Recommendations:

Topiramate

D.7.c.i Topiramate is recommended for limited use in select non-acute back pain patients, where there has been failure of multiple other modalities including trials of different NSAIDs, aerobic exercise, specific stretching exercise, strengthening exercise, tricyclic antidepressants, distractants, and manipulation.

• Frequency/Dose: This medication is initiated by gradually increasing the dose. It has been initiated with a beginning dose of 50 mg and increasing by 50 mg a week. The most appropriate steady dose is unclear, but appears to be 300 mg. Patients should
be carefully monitored for the development of adverse events.

- Discontinuation: Resolution, development of adverse effects, or failure to adhere to a functional restoration program. Careful monitoring of employed patients is indicated due in part to elevated risks for central nervous system (CNS) sedating effects.

Topiramate is not recommended for neuropathic pain, including peripheral neuropathy.

Carbamazepine

D.7.c.ii Carbamazepine is recommended as a potential adjunct for non-acute radicular or neuropathic pain after attempting other treatments (e.g., other medications, aerobic exercise, other exercise, manipulation).

While there is not quality evidence for treatment of non-acute radicular back pain, this may be tried if other medications have failed. Oxcarbazepine and lamotrigine may be useful agents to try if the results from carbamazepine are insufficient for pain relief.

- Frequency/Duration: Frequency and dosing are based on the medication prescribed.

- Discontinuation: Resolution of back pain, lack of efficacy, or development of side effects that necessitate discontinuation. Careful monitoring of employed patients is indicated due to elevated risks for CNS sedating effects.

Gabapentin and Pregabalin

D.7.c.iii Gabapentin is recommended for peri-operative management of pain to reduce need for opioids, particularly in those with side effects from opioids.

- Discontinuation: Resolution or intolerance. Careful monitoring of employed patients is indicated due in part to elevated risks for CNS-sedating effects.
D.7.c.iv  Gabapentin may be considered for the treatment of severe neurogenic claudication from spinal stenosis or non-acute radicular pain syndromes with limited walking distance.

- Discontinuation: Resolution or intolerance. Careful monitoring of employed patients is indicated due in part to elevated risks for CNS-sedating effects.

D.7.c.v  Gabapentin is not recommended for non-acute non-neuropathic pain or back pain.

D.7.d  **Colchicine (Oral and IV Colchicine)**

Recommendations:

D.7.d.i  Oral and IV colchicine are not recommended for acute or non-acute back pain.

D.7.e  **Compound Medications**

Recommendations:

D.7.e.i  Topical, oral and/or systemic compound medications are not recommended.

D.7.f  **Nonsteroidal Anti-Inflammatory Drugs (NSAIDs)**

NSAIDs are useful for pain and inflammation. In mild cases, they may be the only drugs required for analgesia. There are several classes of NSAIDs and the response of the individual patient to a specific medication is unpredictable. For this reason a range of orally administered NSAIDs may be tried in each case with the most effective preparation being continued. Patients should be closely monitored for adverse reactions.

The US Food and Drug Administration advises that all NSAIDs may cause an increased risk of serious cardiovascular thrombotic events, myocardial infarction, and stroke, which can be fatal. Prescribers should be aware of the most updated information on this topic. Some NSAIDs may have more favorable cardiovascular risk factors than others.

Administration of proton pump inhibitors, histamine 2 blockers, or misoprostol, a prostaglandin analog, along with these NSAIDs may reduce the risk of duodenal and gastric ulceration associated with NSAID use but do not impact possible cardiovascular complications. Due to the cross reactivity between aspirin and NSAIDs, NSAIDs should not be used in aspirin-sensitive patients, and should be used
with caution in all asthma patients. NSAIDs are associated with abnormal renal function, including renal failure, as well as abnormal liver function. Certain NSAIDs may have interactions with various other medications. Individuals may have adverse events not listed above. Intervals for metabolic screening are dependent upon the patient’s age, general health status and should be within parameters listed for each specific medication. Complete blood count (CBC), liver and renal function should be monitored in patients on chronic NSAIDs.

In patients receiving low-dose aspirin for primary or secondary cardiovascular disease prevention, to minimize the potential for the NSAID to counteract the beneficial effects of aspirin, aspirin should be taken two hours before or at least eight hours after the NSAID.

Chronic use of NSAIDs is generally not recommended. Chronic NSAIDs may be used cautiously in selected cases with regular monitoring.

D.7.f.i Non-selective Nonsteroidal Anti-Inflammatory Drugs:

Non-selective NSAIDs are generally recommended as first-line medications.

Serious gastrointestinal toxicity, such as bleeding, perforation, and ulceration can occur at any time, with or without warning symptoms in patients treated with traditional NSAIDs. Patients at particularly high risk for GI bleeding include those with a history of prior GI bleed, diabetes, alcohol use, smoking, corticosteroid or anticoagulant use, patients older than 65 or those who have a longer duration of therapy.

Physicians should inform patients about the signs and/or symptoms of serious gastrointestinal toxicity and what steps to take if they occur.

Anaphylactic reactions may occur in patients taking NSAIDs.

NSAIDs may interfere with platelet function.

Fluid retention and edema, and renal toxicity in those with underlying reduction of renal function have been observed in some patients taking NSAIDs.

D.7.f.ii Selective Cyclo-oxygenase-2 (COX-2) Inhibitors:
COX-2 inhibitors should not be first-line for low risk patients who will be using an NSAID short-term but are indicated in select patients for whom traditional NSAIDs are not tolerated. Patients who receive COX-2 inhibitors should take the lowest effective dose for the shortest time necessary to control symptoms.

The major advantages of selective COX-2 inhibitors over traditional NSAIDs are that they have less GI toxicity and no platelet effects.

Serious upper GI adverse events can occur even in asymptomatic patients who are taking COX-2 inhibitors. Patients at a high risk include those with a history of prior GI bleed, diabetes, alcohol use, smoking, corticosteroid or anticoagulant use, patients older than 65 or those who have a longer duration of therapy.

COX-2 inhibitors can worsen renal function in patients with renal insufficiency; thus, renal function may need monitoring.

Selective COX-2 inhibitors should be used with great caution in patients with ischemic heart disease and/or stroke and avoided in patients with risk factors for coronary heart disease. In these patients it appears to be safest to use acetaminophen, aspirin or non-selective NSAIDs as first-line therapy.

Celecoxib is contraindicated in sulfonamide allergic patients.

D.7.g Opioids – Oral, Transdermal, and Parenteral

Narcotics should be primarily reserved for the treatment of severe back pain. In mild-to-moderate cases of pain, narcotic medication should be used cautiously on a case-by-case basis. Adverse effects include respiratory depression, the development of physical and psychological dependence, and impaired alertness. This medication has physically addictive properties and withdrawal symptoms may follow abrupt discontinuation.

Narcotic medications should be prescribed with strict time, quantity, and duration guidelines, and with definitive cessation parameters. Pain is subjective in nature and should be evaluated using a scale to rate effectiveness of the narcotic prescribed.
• Optimum Duration: 3 to 7 days.

• Maximum Duration: 2 weeks. Use beyond two weeks is acceptable in appropriate cases.

Any use beyond the maximum should be documented and justified based on the diagnosis and/or invasive procedures.

Please consult the New York Non-Acute Pain Medical Treatment Guidelines for detailed recommendations regarding the use of opioids.

Recommendations:

D.7.g.i Routine use of opioids for treatment of any acute or non-acute back pain condition is not recommended. There is quality evidence that other medications and treatments are superior to opioids.

D.7.g.ii Limited use of opioids is sometimes needed for treatment of acute back pain patients with severe pain. Opioids may be recommended as adjuncts to more efficacious treatments (especially NSAIDs, muscle relaxants, progressive aerobic exercise, manipulation, and directional exercise). Parenteral administration outside of obvious acute trauma or surgical emergency conditions is almost never required, and requests for such treatments are clinically viewed as red flags for substance abuse. Caution should be used in prescribing opioids for patients with a history of depression, personality disorder, substance addiction, or abuse including alcohol or tobacco.

• Frequency/Duration: Generally prescribed at night or when patients are not at work. Lower doses are preferable as they tend to have better safety profiles. Taper off in 2 weeks.

• Discontinuation: Resolution of pain, improvement to the point of not requiring these medications, intolerance or adverse effects, non-compliance, surreptitious medication use, or use beyond 2 weeks.

D.7.g.iii Limited use of opioids for post-operative pain management is recommended as adjunctive therapy to more effective treatments.
For post-operative pain management, a brief prescription of opioids as adjuncts to more efficacious treatments (especially NSAIDs, muscle relaxants, progressive aerobic exercise, and directional exercise) is often required, especially for lumbar fusion and other more invasive procedures.

- **Frequency/Duration:** Generally prescribed as needed throughout the day, then later only at night, before weaning off completely.

- **Discontinuation:** Resolution of pain, improvement to the point of not requiring these medications, intolerance or adverse effects, non-compliance, surreptitious medication use, or use beyond 2 to 3 weeks for less extensive procedures. Use for up to 6 weeks may be necessary during recovery from more extensive surgical procedures.

**D.7.h Skeletal Muscle Relaxants**

**Recommendations:**

**D.7.h.i** Muscle relaxants are not recommended for mild to moderate acute back pain due to problems with adverse effects, nor are they recommended for chronic use in non-acute back pain (other than acute exacerbations).

**D.7.h.ii** Muscle relaxants are recommended for selected cases of moderate to severe acute back pain as a second-line treatment.

For most cases, these agents are not recommended, since other medications, progressive walking and other exercises will be sufficient to control the symptoms. Generally, it is recommended that these agents be prescribed nocturnally initially and not during workdays or when patients plan to operate motor vehicles. Caution should be used in prescribing skeletal muscle relaxants for those with a history of depression, personality disorder, substance addiction and/or abuse, including alcohol or tobacco. If a muscle relaxant is felt to be necessary in patients with those problems, cyclobenzaprine should be the first drug tried, since its chemical structure resembles a tricyclic antidepressant, and since addiction and abuse of this drug typically do not occur.
Frequency/Duration: This initial dose should be taken in the evening. It is not recommended that the first dose be taken prior to starting a work shift, or operating a motor vehicle or machinery.

Daytime use is acceptable in circumstances in which the patient has experienced only minimal CNS-sedating effects and little concern about sedation compromising the patient’s function or for the patient’s or others’ safety. There is no evidence of benefit from higher doses of medication (e.g., cyclobenzaprine 10 mg over 5 mg). If significant daytime somnolence results, then the medication may need to be discontinued, particularly if it interferes with performance of the aerobic exercise and other components of the rehabilitation plan. Another option is to decrease a dose of cyclobenzaprine by 50% to as little as 2.5 mg. Discontinuation: Resolution of the pain, non-tolerance, significant sedating effects that carry over into the daytime, or other adverse effects.

D.7.h.iii Muscle relaxants are recommended as second- or third-line agents for acute radicular pain syndromes or acute postsurgical pain thought to be musculoskeletal in nature.

Other agents may be more efficacious for relieving radicular pain. Generally, muscle relaxants should be prescribed nocturnally initially and not during workdays or when patients plan on operating motor vehicles.

Frequency/Duration: The initial dose should be in the evening. Daytime use is acceptable in circumstances in which the patient has experienced only minimal CNS-sedating effects and there is little concern about sedation compromising the patient’s function or for the patient’s or others’ safety. If significant daytime somnolence results, then the medication may need to be discontinued, particularly if it interferes with the performance of aerobic exercise and other components of the rehabilitation plan.

- Optimum Duration: 1 week.
- Maximum Duration: 2 weeks (or longer if used only at night).
- Discontinuation: Resolution of the pain, non-tolerance, significant sedating effects that carry over into the daytime, or other adverse effects.
D.7.i  **Systemic Glucocorticosteroids (aka “Steroids”)**

Recommendations:

D.7.i.i  Glucocorticosteroids are not recommended for acute or non-acute back pain without radicular pain or mild to moderate radiculopathy.

D.7.i.ii  Oral steroids are not recommended for axial pain.

D.7.i.iii  Glucocorticosteroids are recommended for treatment of acute severe radicular pain syndrome for purposes of obtaining a short-term reduction in pain.

- Frequency/Duration: It is unclear whether parenteral administration or oral administration is more efficacious. In the absence of evidence, it is suggested that oral administration is preferable due to lower invasiveness and costs. It is recommended that only one course (5 to 14 days) of oral medication (i.e.: tapering dose of methylprednisolone) be prescribed for a given episode of radicular pain. If additional treatment is needed, epidural steroid injections are preferable, since they better target the medication to the affected tissue.

D.7.i.iv  Intravenous steroids are recommended in the setting of an acute neurological emergency and should be confined only to the hospital setting. The dose and duration of the intravenous steroids should be determined in consultation with spinal cord experts. The risk of permanent neurological damage from acute spinal cord compression generally outweighs the risk of pharmacologic side effects of steroids in an emergency situation.

D.7.j  **Topical Drug Delivery**

Topical drug delivery (e.g., capsaicin, lidocaine, topical NSAIDs and topical salicylates and nonsalicylates) may be an acceptable form of treatment in selected patients. A topical agent should be prescribed with strict instructions for application and maximum number of applications per day to obtain the desired benefit and avoid potential toxicity. For most patients, the effects of long-term use are unknown and thus may be better used episodically. These agents may be used in those patients who prefer topical treatments over oral medications. Localized skin reactions may occur, depending on the medication agent used.
Physicians should consider that topical medication can result in toxic blood levels.

D.7.j.i **Capsaicin** offers a safe and effective alternative to systemic NSAIDs, although its use is limited by local stinging or burning sensation that typically disappears with regular use. Patients should be advised to apply the cream on the affected area with a plastic glove or cotton applicator to avoid inadvertent contact with eyes and mucous membranes.

- Optimal Duration: 1-2 weeks to determine effectiveness
- Discontinuation: Resolution of pain, or development of adverse effects that necessitate discontinuation.
- Long-term use of capsaicin is not recommended.

D.7.j.ii **Topical lidocaine** is only indicated when there is documentation of a diagnosis of neuropathic pain. In this instance, a trial for a period of not greater than four weeks can be considered, with the need for documentation of functional gains as criteria for additional use.

D.7.j.iii **Topical NSAIDs** (e.g., diclofenac gel) may achieve tissue levels that are potentially therapeutic. Overall the low level of systemic absorption can be advantageous, allowing the topical use of these medications when systemic administration is relatively contraindicated (such as patients with hypertension, cardiac failure, peptic ulcer disease or renal insufficiency).

D.7.j.iv **Topical salicylates or nonsalicylates** (e.g., methyl salicylate) overall do not appear to be more effective than topical NSAIDs. May be used for a short-term course especially in patients with chronic conditions in whom systemic medication is relatively contraindicated or as an adjuvant to systemic medication.

D.7.k **Tramadol**

Recommendations:
D.7.k.i Tramadol is useful in relief of pain and has been shown to provide pain relief equivalent to that of commonly prescribed NSAIDs.

Although Tramadol may cause impaired alertness, it is generally well tolerated, does not cause gastrointestinal ulceration, or exacerbate hypertension or congestive heart failure. Tramadol should be used cautiously in patients who have a history of seizures or who are taking medication that may lower the seizure threshold, such as monoamine oxidase (MAO) inhibitors, SSRIs, and tricyclic antidepressants.

- Maximum Duration: 2 weeks. Use beyond 2 weeks is acceptable in appropriate cases.

This medication has physically addictive properties and withdrawal symptoms may follow abrupt discontinuation. It is not recommended for those with prior opioid addiction.

D.7.i **Tumor Necrosis Factor-α Inhibitors**

Recommendations:

D.7.l.i Tumor necrosis factor-α inhibitors are not recommended for treatment of radicular pain syndromes.

D.7.l.ii Tumor necrosis factor-α inhibitors are not recommended for treatment of acute or non-acute back pain.

D.7.m **Vitamins**

Recommendations:

The use of vitamins in the absence of documented deficiencies or other nutritional deficit states for acute, non-acute, or post-operative back pain patients and for patients with radiculopathy is not recommended.

D.8 **SLEEP POSTURE**

Recommendations:

D.8.a.i Alteration of sleep posture may be recommended in acute or non-acute back pain that results in nocturnal awakening, particularly if not amenable to other treatments.
The most appropriate sleep posture is that which is most comfortable for the patient. If a patient habitually chooses a particular sleep posture, it would appear reasonable to recommend altering posture to determine if there is reduction in pain or other symptoms.

- Discontinuation: Non-tolerance.

D.8.a.ii There is no quality evidence that specific commercial products have roles in primary prevention or treatment of acute or non-acute back pain.
D.9 THERAPY: ACTIVE

D.9.a Therapeutic Exercise

Therapeutic Exercise with or without mechanical assistance or resistance, may include isoinertial, isotonic, isometric and isokinetic types of exercises. Indications include the need for cardiovascular fitness, reduced edema, improved muscle strength, improved connective tissue strength and integrity, increased bone density, promotion of circulation to enhance soft tissue healing, improvement of muscle recruitment, improved proprioception and coordination, increased range of motion and are used to promote normal movement patterns. Therapeutic exercise can also include complementary/ alternative exercise movement therapy (with oversight of a physician or appropriate healthcare professional).

- Time to Produce Effect: 2 to 6 treatments.
- Frequency: 3 to 5 times per week.
- Optimum Duration: 4 to 8 weeks.
- Maximum Duration: 8 weeks.

D.9.b Aerobic Exercises

Recommendations:

D.9.b.i Aerobic exercise is recommended for treatment of all patients with acute or non-acute back pain, although most available evidence is from studies treating chronic back pain patients. Consideration should be given, however, to whether an evaluation is required prior to institution of vigorous exercises for those with significant cardiac disease, or significant potential for cardiovascular disease. For most patients, a structured, progressive walking program on level ground or no incline on a treadmill is recommended. There has been some controversy about whether bicycling is helpful or harmful from a biomechanical perspective (lordosis) and the back muscles are less active with bicycling, thus it may be less appropriate. Yet, if bicycling is the preferred exercise for the patient, it is believed to be far superior to performing no aerobic exercise. For those patients who desire other aerobic exercises, there are no specific data, although there are indications that imply that there is a direct correlation between benefit and the amount of aerobic activity that
results in higher MET expenditure. Therefore, the activity that the patient will adhere to is believed to be the one most likely to be effective, given that compliance is a recognized problem.

- Frequency/Duration: For patients with non-acute back pain, walking at least 4 times per week at 60% of predicted maximum heart rate (220-age = maximum heart rate) is recommended. One successful study benchmarked 20 minutes during Week 1, 30 minutes during Week 2, and 45 minutes after that point. For acute back pain patients, a graded walking program is generally desired, often using distance or time as minimum benchmarks. For example, a patient can start with 10 to 15 minutes twice a day for 1 week, and increase in 10 to 15 minute increments per week until at least 30 minutes per day is achieved.

- Discontinuation: Aerobic exercise should be discontinued when there is intolerance (rarely occurs) or development of other disorders. Nearly all patients should be encouraged to maintain aerobic exercises on a long-term basis for both prevention of back pain (see below), and to maintain optimal health.

D.9.b.ii For post-operative patients aerobic exercise is recommended.

In the absence of quality evidence to support this recommendation, it is suggested that recommendation D.9.b.i be used for aerobic exercise for treatment of post-operative back pain patients.

D.9.c Strengthening and Stabilization Exercises

Recommendations:

D.9.c.i For acute or non-acute back pain, or post-operative back pain patients, strengthening exercises are recommended for treatment of back pain. Specific strengthening exercises, such as stabilization exercises, are helpful for the prevention and treatment (including post-operative treatment) of back pain.
As evidence of efficacy of aerobic exercises appears greater, these exercises should be added after either aerobic exercises have already been instituted and additional treatment is needed, or in situations where both are felt to be required. Exercises should be taught and then performed by the patient in a home exercise program. For those patients who do not improve, follow up appointments to verify technique and compliance (by exercise log books) are recommended. Some patients, particularly those lacking motivation to be in a home exercise program may benefit from a supervised exercise program, although there are questions about long-term compliance among patients with non-acute back pain. More intensive programs with more intensive exercises and direct supervision with active coaching have been shown to be effective for non-acute back pain.

- Frequency/Duration: Home program frequency is 1 to 2 times a day for acute back pain, and two to three times a day for non-acute back pain.

- Discontinuation: Indications to discontinue strengthening exercises include development of a strain in the course of treatment or failure to improve.

**D.9.c.ii** Abdominal strengthening exercises particularly as either a sole or central goal of a strengthening program are not recommended for treatment or prevention of back pain.

Strengthening of abdominal muscles (e.g., rectus abdominus and obliques with sit-up exercises) is a frequent goal of back pain rehabilitation or prevention programs. There is no quality evidence that these exercises are effective, there is evidence that suggests they are not effective, and there are other treatment strategies with proven or at least suggested greater efficacy.

**D.9.d** **Aquatic Therapy (Including Swimming)**

Recommendations:

**D.9.d.i** A trial of aquatic therapy is recommended for the treatment of non-acute back pain in a patient who meets criteria for a referral for supervised exercise therapy and has co-morbidities (e.g., extreme obesity, significant degenerative joint disease, etc.) that preclude effective participation in a
weight-bearing physical activity. Osteoarthritis of the knee is not a clear contraindication to a walking program, rather walking may be therapeutically indicated based on high quality evidence.

- Frequency/Duration: A program should generally begin with 3 to 4 visits per week. The patient must have demonstrated evidence of functional improvement within the first 2 weeks to justify additional visits. The program should include up to 4 weeks of aquatic therapy with progression towards a land-based, self-directed physical activity or self-directed aquatic therapy program by 6 weeks.

- Discontinuation: Non-tolerance, failure to progress, or reaching a 4 to 6 week time frame.

For all other non-acute back pain patients, and for all acute back pain, aquatic therapy is not recommended as other therapies are believed to be more efficacious.

**D.9.e MedX Machine**

Recommendations:

D.9.e.i Use of a MedX machine to strengthen the lumbar spine is not recommended for acute or non-acute back pain or for any radicular pain syndrome.

**D.9.f Yoga**

Recommendations:

D.9.f.i There is some evidence to support the effectiveness of yoga therapy in alleviating symptoms and decreasing medication use for patients with uncomplicated back pain.

- Frequency/Duration: 2 to 5 times per week.
- Time to Produce Effect: 2 to 6 treatments.
- Optimum Duration: 4 weeks.
- Maximum Duration: Reassess after 8 weeks
D.10 THERAPY: PASSIVE

D.10.a Manipulation

Manipulative treatment (not therapy) is defined as the therapeutic application of manually guided forces by an operator to improve physiologic function and/or support homeostasis that has been altered by the injury or occupational disease, and has associated clinical significance.

Contraindications to manipulation may include joint instability, fractures, severe osteoporosis, infection, metastatic cancer, active inflammatory arthritides, and signs of progressive neurologic deficits or myelopathy. Relative contraindications include stenosis, spondylosis, and disc herniation.

D.10.a.i Manipulation is recommended for treatment of acute back pain when tied to objective measures of improvement.

- Time to produce effect for all types of manipulative treatment: 1 to 6 treatments.
- Frequency: Up to 3 times per week for the first 4 weeks as indicated by the severity of involvement and the desired effect, then up to 2 treatments per week for the next 4 weeks with re-evaluation for evidence of functional improvement or need for further workup. Continuance of treatment will depend upon functional improvement.
- Optimum Duration: 8 to 12 weeks.
- Maximum Duration: 3 months. Extended durations of care beyond what is considered “maximum” may be necessary in cases of re-injury, interrupted continuity of care, exacerbation of symptoms, and in those patients with comorbidities.

D.10.a.ii A maintenance program of spinal manipulation (by a physician (MD/DO), chiropractor or physical therapist) may be indicated in certain situations, after the determination of MMI, when tied to maintenance of functional status. (See Section D.11, Therapy: Ongoing Maintenance Care.)
D.10.a.iii There is no evidence that prophylactic treatment is effective, either for primary prevention (before the first episode of pain) or for secondary prevention (after recovery from an episode of back pain) and prophylactic treatment is not recommended.

D.10.b Manipulation under Anesthesia (MUA) and Medication-Assisted Spinal Manipulation (MASM)

Recommendations:

D.10.b.i MUA and MASM are not recommended in acute or non-acute back pain patients.

D.10.c Massage (Manual or Mechanical)

Massage (Manual or Mechanical) consists of manipulation of soft tissue with broad-ranging relaxation and circulatory benefits. This may include stimulation of acupuncture points and acupuncture channels (acupressure), application of suction cups and techniques that include pressing, lifting, rubbing, pinching of soft tissues by or with the practitioner's hands. Indications include edema (peripheral or hard and non-pliable edema), muscle spasm, adhesions, the need to improve peripheral circulation and range of motion, or to increase muscle relaxation and flexibility prior to exercise.

As with all passive therapies, massage must be accompanied by exercise and patient education. Objective benefit (functional improvement along with symptom reduction) must be demonstrated in order for further treatment to continue.

D.10.c.i Massage is recommended for select use in non-acute back pain as an adjunct to more efficacious treatments consisting primarily of a graded aerobic and strengthening exercise program.

- Time to Produce Effect: Immediate.
- Frequency: 1 to 2 times per week.
- Optimum Duration: 6 weeks.
- Discontinuation: Resolution, intolerance, lack of benefit, or non-compliance with aerobic and strengthening exercises.
D.10.c.ii  Massage is recommended as a treatment for acute back pain and non-acute radicular syndromes in which back pain is a substantial symptom component.

- Time to Produce Effect: Immediate.
- Frequency: 1 to 2 times per week.
- Optimum Duration: 6 weeks.
- Discontinuation: Resolution, intolerance or lack of benefit.

D.10.c.iii  Massage is recommended for patients with non-acute back pain without underlying serious pathology, such as fracture, tumor, or infection.

- Time to Produce Effect: Immediate.
- Frequency: 1 to 2 times per week.
- Optimum Duration: 6 weeks.
- Discontinuation: Resolution, intolerance or lack of benefit.

D.10.c.iv  Mechanical devices for administering massage are not recommended.

D.10.d  **Mobilization (Joint)**

Mobilization consists of passive movement involving oscillatory motions to the vertebral segment(s). The passive mobility is performed in a graded manner (I, II, III, IV, or V), which depicts the speed and depth of joint motion during the maneuver. It may include skilled manual joint tissue stretching. Indications include the need to improve joint play, segmental alignment, improve intracapsular arthrokinematics, or reduce pain associated with tissue impingement. Mobilization should be accompanied by active therapy. For Level V mobilization, contraindications include joint instability, fractures, severe osteoporosis, infection, metastatic cancer, active inflammatory arthritides, and signs of progressive neurologic deficits, myelopathy, vertebrobasilar insufficiency, or carotid artery disease. Relative contraindications include stenosis, spondylosis, and disc herniation.

- Time to Produce Effect: 6 to 9 treatments.
- Frequency: Up to 3 times per week.
- Optimum Duration: 4 to 6 weeks.
- Maximum Duration: 6 weeks.

**D.10.e Mobilization (Soft Tissue)**

Mobilization of soft tissue is the skilled application of muscle energy, strain/counter strain, manual trigger point release, and other manual therapy techniques designed to improve or normalize movement patterns through the reduction of soft tissue pain and restrictions. These can be interactive with the patient participating or can be with the patient relaxing and letting the practitioner move the body tissues. Indications include muscle spasm around a joint, trigger points, adhesions, and neural compression. Mobilization should be accompanied by active therapy.

- Time to Produce Effect: 4 to 9 treatments.
- Frequency: Up to 3 times per week.
- Optimum Duration: 4 to 6 weeks.
- Maximum Duration: 6 weeks.

**D.10.f Superficial Heat and Cold**

Superficial heat and cold are thermal agents applied in various manners that lower or raise the body temperature for the reduction of pain, inflammation, and/or effusion resulting from injury or induced by exercise. It includes application of heat just above the surface of the skin at acupuncture points.

Recommendations:

**D.10.f.i** Recommended for acute pain, edema, and hemorrhage, need to increase the pain threshold, reduce muscle spasm, and promote stretching/flexibility. Cold and heat packs can be used at home as an extension of therapy in the clinic setting.
• Time to Produce Effect: Immediate.

• Frequency/Duration: Frequency: 2 to 5 times per week.

• Optimum Duration: 3 weeks as primary, or intermittently as an adjunct to other therapeutic procedures up to two months.

D.10.g **Diathermy**

Recommendations:

D.10.g.i Diathermy is not recommended for treatment of any back pain-related conditions.

D.10.h **Infrared Therapy**

Recommendations:

D.10.h.i For those circumstances where this intervention is used for treatment of acute back pain, it is recommended to only be provider-based treatment and only performed in conjunction with an active exercise program, with frequency not to exceed 4 visits.

D.10.i **Ultrasound**

Recommendations:

D.10.i.i In situations where deeper heating is desirable, a limited trial of ultrasound for the treatment of back pain is reasonable, but only if performed as an adjunct with exercise.

  • Frequency/Duration: 3 times per week.

  • Time to produce effect: 6 to 15 treatments.

  • Optimum Duration: 4 to 8 weeks.

  • Maximum Duration: 8 weeks.
D.10.j  **Low Level Laser Therapy**

Recommendations:

D.10.j.i  Low level laser therapy is not recommended for treatment of back pain.

D.10.k  **Neuroflexotherapy**

Recommendations:

D.10.k.i  Neuroflexotherapy is not recommended for the treatment of acute or non-acute back pain, radicular back pain, or other spinal conditions.

D.10.l  **Reflexology**

Recommendations:

D.10.l.i  Reflexology is not recommended for the treatment of acute or non-acute back pain, radicular back pain, or other spinal conditions.

D.10.m  **Traction**

Traction is not recommended for treatment of acute or non-acute back pain or radicular pain syndromes.

D.10.n  **Vertebral Axial Compression (VAX-D) and Other Decompressive Devices**

Recommendations:

D.10.n.i  VAX-D or other spinal decompressive devices is not recommended for acute or non-acute back pain or radicular pain syndromes.
D.11 THERAPY: ONGOING MAINTENANCE CARE

A maintenance program of physical therapy, occupational therapy or spinal manipulation (by a physician (MD/DO), chiropractor or physical or occupational therapist) may be indicated in certain situations, after the determination of MMI, when tied to maintenance of functional status.

- Although the current body of scientific evidence as reviewed does not support the routine use of this intervention, maintenance therapy modalities may be indicated in certain situations in order to maintain functional status, without which an objective deterioration of function has been previously observed and documented in the medical record.

- Specific objective goals should be identified and measured in order to support the need for ongoing maintenance care.

- Progressively longer trials of therapeutic withdrawal are to be attempted to ascertain whether therapeutic goals can be maintained in the absence of clinical interventions.

- Within a year and annually thereafter, a trial without maintenance treatment should be instituted.

- The care of chronic back symptoms should include an ongoing patient self-management plan performed by the patient regularly and a self-directed pain management program initiated as indicated:
  - An ongoing clinically appropriate self-management plan, typically independent, home-based and self-directed, developed jointly by the provider and patient, should be implemented to encourage physical activity and/or work activities despite residual pain, with the goal of preserving function.
  - In addition to the self-management plan, a self-directed pain management plan should be developed which can be initiated by the patient in the event that symptoms worsen and function decreases.

- If deterioration of ability to maintain function is documented, reinstatement of ongoing maintenance may be acceptable.

Frequency: Maximum up to 10 visits/year, after the determination of MMI, according to objectively documented maintenance of functional status. No variance from the maximum frequency is permitted.

Ongoing Maintenance Care is a component of the Functional Maintenance Care recommendations detailed in the New York Non-Acute Pain Medical
Treatment Guidelines. Please consult the New York Non-Acute Pain Medical Treatment Guidelines for additional information.
D.12 RADIOFREQUENCY ABLATION, NEUROTOMY, FACET RHIZOTOMY

D.12.a Radiofrequency Ablation, Neurotomy, Facet Rhizotomy

Description:

A procedure designed to denervate the facet joint by ablating the corresponding sensory medial branches. Continuous percutaneous radiofrequency is the method generally used.

Radiofrequency medial branch neurotomy is recommended as the procedure of choice over alcohol, phenol, other injectable agents, or cryoablation.

Fluoroscopic guidance is required for precise positioning of the probe.

Permanent images should be recorded to verify placement of the device.

Recommendations:

• For patients with proven facet joint pain in whom two diagnostic medial nerve branch blocks have been therapeutically successful, the use of radiofrequency ablation/neurotomy/facet rhizotomy may be indicated.

• This procedure is not recommended for involvement of more than 3 facet joints (4 medial branch nerves).

• All patients should have a successful response to a diagnostic medial nerve branch block and a separate comparative block.

• To be a positive diagnostic block the patient should report a reduction of pain of 50% or greater from baseline for the length of time appropriate for the local anesthetic used correlated with functional improvement.

• The patient should also identify activities of daily living (which may include measurements of range-of-motion) that are impeded by their pain. The physician should observe and document functional improvement in the identified activities in the clinical setting.

Post-Procedure Therapy:

Active therapy

• Implementation of a gentle reconditioning program within the first post-procedure week is recommended, barring complications.

• Instruction and participation in a long-term home-based program of range of motion, cervical, scapular, and thoracic strengthening, postural or
neuromuscular re-education, endurance, and stability exercises should be accomplished over a period of four to ten visits post-procedure.

Requirements for repeat radiofrequency neurotomy (or additional level radiofrequency neurotomies):

- In some cases pain may recur.
- Successful rhizotomy usually provides from six to nine months of relief.
- Before a repeat radiofrequency neurotomy is done, a confirmatory medial branch injection may be performed if the patient’s pain pattern presents differently than in the initial evaluation.

Maximum Frequency:

Twice a year as indicated by improvement in pain and function.

**D.12.b Dorsal Root Ganglia Radiofrequency Lesioning**

Recommendations:

D.12.b.i Radiofrequency lesioning of the dorsal root ganglia is not recommended for chronic sciatica.

Radiofrequency lesioning is invasive, has adverse effects and is costly. It has been shown to not be efficacious in a well-designed, high-quality study.

**D.12.c Intradiscal Electrothermal Therapy (IDET)**

Recommendations:

D.12.c.i IDET is not recommended for treatment of acute or non-acute back pain, or any other back-related disorder.

**D.12.d Percutaneous Intradiscal Radiofrequency Thermocoagulation (PIRFT)**

Recommendations:

D.12.d.i Percutaneous Intradiscal Radiofrequency Thermocoagulation is not recommended for treatment of acute or non-acute back pain, including discogenic back pain.
E  THERAPEUTIC PROCEDURES: OPERATIVE

E.1  Discectomy, Microdiscectomy, Sequestrectomy, Endoscopic Decompression

Recommendations:

E.1.a.i  Lumbar discectomy is recommended as an effective operation to speed recovery in patients who have radiculopathy due to ongoing nerve root compression, who continue to have significant pain and functional limitation after 6 to 12 weeks and who have been provided appropriate conservative therapy during which time they experienced no progressive neurological deficits.

E.1.a.ii  All of the following should be present: 1) radicular pain syndrome with current dermatomal pain and/or numbness, or myotomal muscle weakness - consistent with a herniated disc at the corresponding level; 2) imaging findings by MRI or CT with/without myelography that confirm persisting nerve root compression at the level and on the side predicted by the history and clinical examination; 3) continued significant pain and functional limitation after 6 to 12 weeks and who have been provided appropriate conservative therapy during which time they experienced no progressive neurological deficits.

Patients who are candidates for discectomy should be informed that (other than for cauda equina syndrome and the rare progressive major neurologic deficit), there is evidence that there is no need to rush surgical decisions, since there is no difference in long-term functional recovery whether surgery is performed early or delayed. Open discectomy, microdiscectomy, and endoscopic discectomy are all potentially appropriate ways to perform discectomy. The decision as to which of these procedures to choose should be left to the surgeon and the patient, until quality evidence becomes available to provide evidence-based guidance.

E.1.a.iii  Discectomy is not recommended as treatment of acute or non-acute back pain without radiculopathy.

E.1.a.iv  Percutaneous discectomy (nucleoplasty), laser discectomy, and disc coblation therapy are not recommended as treatment for any back or radicular pain syndrome.
E.2 ADHESIOLYSIS

Recommendations:

E.2.a.i Adhesiolysis is not recommended for acute or non-acute back pain, spinal stenosis, or radicular pain syndromes.

E.3 DECOMPRESSIVE SURGERY (LAMINOTOMY/FACETECTOMY LAMINECTOMY)

Recommendations:

E.3.a.i Decompression surgery is recommended as an effective treatment for patients with symptomatic spinal stenosis (neurogenic claudication) that is intractable to conservative management.

E.4 SPINAL FUSION

Lumbar Fusion is not included on the list of pre-authorized procedures. Providers who want to perform one of these procedures must request pre-authorization from the carrier before performing the procedure.

Recommendations:

E.4.a.i Lumbar fusion is recommended as a treatment for spinal stenosis when concomitant instability has been proven. Lumbar fusion is not recommended for spinal stenosis without instability.

Indications: All of the following should be present: 1) neurogenic claudication (leg pain and/or numbness with standing or walking); 2) imaging findings, by MRI, or CT/myelogram that confirm the nerve roots compressed are consistent with the neurological symptoms; 3) lack of responsiveness or unsatisfactory response(s) to adequate conservative treatment over a minimum 6 to 8 week period that may or may not include an epidural steroid injection.

E.4.a.ii Lumbar fusion is recommended as an effective treatment for isthmic spondylolisthesis.

E.4.a.iii Lumbar fusion is recommended as an effective treatment for degenerative spondylolisthesis.
E.4.a.iv There are no scientific studies, but consensus is that if a patient is having the third lumbar discectomy on the same disc, that spine fusion at the time of discectomy is an option.

E.4.a.v Lumbar fusion is not recommended as a treatment for patients with radiculopathy from herniated nucleus pulposus (disc herniation) or for patients with non-acute back pain after lumbar discectomy.

E.4.a.vi Lumbar fusion is recommended as a treatment for Degenerative Disc Disease/“Discogenic Back Pain”/“Black Disc Disease” without instability in selected patients for whom non-surgical management has failed to relieve symptoms and improve function. If available, an intensive Functional Rehabilitation Program should be tried first.

E.5 ELECTRICAL BONE GROWTH STIMULATORS

Electrical Bone Growth Stimulators are not included on the list of pre-authorized procedures. Providers who want to perform one of these procedures must request pre-authorization from the carrier before performing the procedure.

Recommendations

E.5.a.i Non-invasive Electrical Bone Growth Stimulators may be considered as an adjunct to spinal fusion surgery for those at high risk for pseudoarthrosis, including one or more of the following fusion failure risk factors:

1) One or more previous failed spinal fusion(s)

2) Grade II or worse spondylolisthesis

3) Fusion to be performed at more than one level

4) Presence of other risk factors that may contribute to non-healing:
   - Current smoking
   - Diabetes
   - Renal disease
- Other metabolic diseases where bone healing is likely to be compromised (e.g.: significant osteoporosis)
- Active alcoholism
- Morbid obesity BMI >40

E.5.a.ii Non-invasive Electrical Bone Growth Stimulators may be considered as treatment for individuals with failed spinal fusion. Failed spinal fusion is defined as a spinal fusion that has not healed at a minimum of 6 months after the original surgery, as evidenced by serial x-rays over a course of 3 months during the latter portion of the 6-month period.

E.6 DISC REPLACEMENT

Artificial Disc Replacement is not included on the list of pre-authorized procedures. Providers who want to perform one of these procedures must request pre-authorization from the carrier before performing the procedure.

Recommendations:

E.6.a.i Artificial disc replacement is recommended as treatment for lumbar degenerative disc disease at one level with radiculopathy that is unresponsive to conservative management.

The following criteria must be met:

1) Skeletally mature patient without renal failure, severe diabetes, osteoporosis, severe spondylosis, severe facet pathology, lumbar instability, localized fracture, or localized or systemic infections.

AND

2) Single-level disc degeneration of L3 to S1 confirmed by imaging studies such as CT or MRI, with one of the following diagnoses:

- Herniated disc; or
- Osteophyte formation; or
- Loss of disc height.

AND
3) The patient must present with **symptoms**, which must correspond with the planned level of disc replacement:

- Intractable radiculopathy (nerve root compression) and/or myelopathy (functional disturbance or pathological change in the spinal cord) causing radicular pain in the lower extremity; **or**

- Functional and/or neurological deficit.

**AND**

4) Six weeks of non-operative alternative treatments have failed. These treatments may include physical therapy, medications, braces, chiropractic care, bed rest, spinal injections or exercise programs. Documentation of treatments and failure to improve is required.

The disc must be approved by the U.S. Food and Drug Administration (FDA). All other artificial disc systems are considered experimental and investigational.

All other indications, including multilevel degenerative disc disease, are considered experimental and investigational.

**Artificial disc replacement is NOT recommended under the following conditions, since safety and effectiveness of the replacement discs has not been established for patients with:**

- Previous surgical intervention at the involved level;

- Prior or proposed fusion at an adjacent cervical level;

- More than one lumbar level requiring artificial disc replacement;

- Clinically compromised vertebral bodies at the affected level due to current or past trauma (including but not limited to the radiographic appearance of fracture callus, malunion or nonunion);

- Active systemic infection or infection at the operating site;
• Allergy to titanium, polyurethane, or ethylene oxide residues;

• Osteoporosis defined as a DEXA bone mineral density T score equal to or worse than -2.5.

### E.7 VERTEBROPLASTY AND KYPHOPLASTY

Vertebroplasty and Kyphoplasty are not included on the list of pre-authorized procedures. Providers who want to perform one of these procedures must request pre-authorization from the carrier before performing the procedure.

Recommendations:

E.7.a.i Vertebroplasty and Kyphoplasty may be considered for treatment of select patients with vertebral body compression fractures with associated non-acute or severe pain. Patients who have had fractures despite bisphosphonate therapy are particularly appropriate candidates.

### E.8 SACROILIAC SURGERY

Recommendations:

E.8.a.i SI joint fusion surgery and other SI joint surgical procedures are not recommended.

### E.9 INTRAOPERATIVE MONITORING

Intraoperative Monitoring is a common intraoperative electrodiagnostic technique that may include somatosensory evoked potentials (SSEP), motor evoked potentials (MEP), or pedicle screw monitoring. The monitoring procedure may be used to evaluate spinal cord integrity and screw placement during the operative procedure. The use of intraoperative monitoring can be anticipated to become more common as percutaneous spinal procedures gain greater acceptance.

### E.10 IMPLANTABLE SPINAL CORD STIMULATORS (SCS)

Please consult the New York Non-Acute Pain Medical Treatment Guidelines for information about Spinal Cord Stimulators. Spinal Cord Stimulators are not included on the list of pre-authorized procedures. Providers who want to perform one of these procedures must request pre-authorization from the carrier before performing the procedure.
Sources:

This Medical Treatment Guideline is adopted, with modification, from ACOEM’s Occupational Medicine Treatment Guidelines for Low Back Disorders with supplementation from the State of Colorado’s Low Back Medical Treatment Guidelines.
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