Latest Evidence-Based Application for Radiofrequency Neurotomy (LEARN): Best Practice Guidelines from the American Society of Pain and Neuroscience (ASPN)

Abstract: Radiofrequency neurotomy (RFN), also known as radiofrequency ablation (RFA), is a common interventional procedure used to treat pain from an innervated structure. RFN has historically been used to treat chronic facet-joint mediated pain. The use of RFN has more recently expanded beyond facet-joint mediated pain to peripherally innervated targets. In addition, there has also been the emergence of different radiofrequency modalities, including pulsed and cooled RFN. The use of RFN has been particularly important where conservative and/or surgical measures have failed to provide pain relief. With the emergence of this therapeutic option and its novel applications, the American Society of Pain and Neuroscience (ASPN) identified the need for formal evidence-based guidance. The authors formed a multidisciplinary work group tasked to examine the latest evidence-based medicine for the various applications of RFN, including cervical, thoracic, lumbar spine; posterior sacroiliac joint pain; hip and knee joints; and occipital neuralgia. Best practice guidelines, evidence and consensus grading were provided for each anatomical target.

Keywords: radiofrequency neurotomy, radiofrequency ablation, pulsed ablation, cooled ablation, sacroiliac joint neurotomy, peripheral joint neurotomy, occipital neuralgia, chronic pain

Introduction

Radiofrequency neurotomy (RFN), also known as radiofrequency ablation (RFA), is a standard procedure performed to treat pain from an innervated structure. Increased utilization of RFN has been driven by an aging population, a desire to avoid more invasive procedures, and the desire for an alternative to less efficacious and higher risk options, such as chronic long-term opioid therapy. Historically, RFN has been used to treat chronic facet-joint mediated pain diagnosed by facet or medial branch blocks. With the expanded use of this therapy for novel applications involving peripheral joints and nerves, there was an identified need for evidence-based best practice guidance. In addition, there has also been the emergence of different radiofrequency modalities, including pulsed and cooled RFN. This review aims to present the most up-to-date use of radiofrequency neurotomy based on anatomical targets.
History of Radiofrequency Neurotomy

In the early 1930s, Kirschner demonstrated the first known utilization of RFN with thermocoagulation of the Gasserian ganglion for trigeminal neuralgia. Kirschner's initial work demonstrated that continuous radiofrequency (CRF) current created a focal thermal lesion in a neural pathway with the goal to interrupt nociception.1 Two decades later, in the 1950s, work by Aronow and Cosman would lead to the advent of the first commercially available radiofrequency (RF) systems. Shealy and Bogduk would later refine percutaneous medial branch RF neurotomy techniques,2 a procedure that essentially replaced surgical neurotomy. Initially, limitations in technology only allowed for the treatment of cervical and lumbar facet disease. However, CRF has now been studied in the treatment of numerous pain pathologies. Despite the technological advances, the risk of motor deficit remains a concern.3–6

Pulsed radiofrequency (PRF) technology, first introduced in Austria in 1995, was developed to reduce the risk of motor deficit as it does not create a destructive thermal lesion. Ayrapetian proposed that PRF efficacy may be secondary to magnetic field exposure as opposed to thermal coagulation.7 In 1998, the first article on the effects of PRF on the dorsal root ganglion was published.8–10

A recent novel modality for ablation of neural pathways is cooled radiofrequency (CRFN) thermal neurotomy. Despite the name, this technique allows for a larger thermal lesion to be formed than traditional RFN (Figure 1). This method has been increasingly utilized for the interruption of nociceptive pathways after its initial use in cardiac electrophysiology and tumor ablation. Since 2010, there has been emerging evidence supporting the use of CRFN for chronic knee, hip and back pain.11–13

Methods

Based on the identified gap in guidance on the use of RFN, the American Society of Pain and Neuroscience (ASPN) formed a multidisciplinary work group of pain medicine specialists to create a best practice guideline. Selection of panel members was based on expertise, publications, research, clinical experience, practice setting and diversity. All work group members actively utilize RFN in their respective practices. While there have been previous publications on the use of RFN, work group members identified the need to expand best practice guidelines to address sacroiliac joint pain and peripheral joint disease. The aim of the paper was to present the most up-to-date use of this modality based on anatomical targets.

A formal literature search was performed by the authors with clinical experience in RFN procedures. Database searches of Embase, PubMed and Medline, utilizing the following keywords was performed: cervical radiofrequency neurotomy/ablation, thoracic radiofrequency neurotomy/ablation, lumbar radiofrequency neurotomy/ablation, sacroiliac joint radiofrequency neurotomy/ablation, lateral sacral branch neurotomy/ablation, hip radiofrequency neurotomy/ablation, genicular radiofrequency neurotomy/ablation, knee radiofrequency neurotomy/ablation, and occipital radiofrequency neurotomy/ablation. The terms neurotomy and ablation are used interchangeably in this paper. In addition, the terms facet, medial branch, and medial branch nerves are used interchangeably in this paper.

Inclusion criteria for literature search were randomized controlled trials, prospective and retrospective observational studies. Exclusion criteria included conference proceedings, expert opinions, and unpublished data. The records were filtered for “pain management” specific topics. The remaining abstracts were reviewed by two authors utilizing PICOS technique with comparison to other treatment (active, sham or placebo) or no treatment, and outcome measures to include improvement in pain by any scale. Secondary outcomes of interest: function, analgesic use, subsequent need for surgery, healthcare

Figure 1 Lesioning performed in chicken breast (37°C). Left probe, Cooled RF 18G with 4mm active tip, set temperature of 60°C. Right probe, Standard RF with 20G with 5mm active tip, set temperature 80°C. Arrow annotates the location of the probe tips. Used with permission from Avanos Medical, Inc.
utilization, return to work. The screened manuscripts were then separated by anatomic targets:

1. Cervical Medial Branches
2. Thoracic Medial Branches
3. Lumbar Medial Branches
4. Lateral Sacral Branches (Posterior sacroiliac ligaments and joint)
5. Knee Joints
6. Hip Joints
7. Occipital Nerve Branches

A formal meta-analysis was not conducted due to heterogeneity in study types, diagnostic criteria, variances in interventional techniques utilized and discrepancies in primary and secondary endpoints. Literature review was conducted by each work group member for assessment of both quality of evidence and grade of recommendation. Any discrepancies in grading were brought to the work group for discussion so that a consensus could be reached. In the end, there were no discrepancies to disclose. Each section includes a summary of:

1. The most current evidence-based medicine for the use of RFN
2. Patient selection guidance
3. Procedure techniques and best practice recommendations
4. Treatment and research gaps.
5. Quality of evidence for each anatomic target graded from I–III as detailed in Table 1.

Conflict of Interest Management: Each member of the panel recused itself from any decision in which a conflict existed. One author (KA) had the responsibility of evaluating the paper for bias and resolving any potential issues prior to manuscript publication.

Cervical Medial Branch Radiofrequency Neurotomy
Cervical medial branch RFN, also commonly referred to as cervical facet joint neurotomy or ablation, is used for the treatment of chronic neck pain and cervicogenic headache. While physical examination and imaging may support the diagnosis of cervical facet-mediated pain, these alone have not been validated as standalone diagnostic techniques. Neck tenderness or pain with extension and/or rotational movements, though often associated with cervical facet joint mediated pain, are not specific to solely facet-mediated pain. While commonly used diagnostically in pain medicine, diagnostic cervical medial branch/facet blocks are not specifically covered in this manuscript as they are beyond the scope of this paper.

Current Available Evidence of Cervical Medial Branch Radiofrequency Neurotomy
Early descriptive studies of radiofrequency neurotomy date back half a century; recent studies are summarized in Table 3.

Procedure Techniques
The effectiveness of the cervical RFN is highly dependent on proper technique. There is robust evidence for short- and long-term pain relief with denervation protocols involving multiple lesions. This technique has since been adopted as the standard of care.

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Table 1 Quality of Evidence, Based on the USPSTF Criteria

<table>
<thead>
<tr>
<th>Evidence Level</th>
<th>Definition</th>
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<tbody>
<tr>
<td>I</td>
<td>At least one controlled and randomized clinical trial with proper design</td>
</tr>
<tr>
<td>II-1</td>
<td>Well designed, controlled, nonrandomized clinical trial</td>
</tr>
<tr>
<td>II-2</td>
<td>Cohort or case studies add well designed controls, preferably multicenter</td>
</tr>
<tr>
<td>II-3</td>
<td>Multiple series compared over time, with or without intervention, and surprising results</td>
</tr>
<tr>
<td>III</td>
<td>Experience driven opinions, clinical observations</td>
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</table>
**Table 2 Recommendation Grade Level**

<table>
<thead>
<tr>
<th>Grade</th>
<th>Definition</th>
<th>Suggestions for Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>The USPSTF recommends the service. There is high certainty that the net benefit is substantial.</td>
<td>Offer or provide this service.</td>
</tr>
<tr>
<td>B</td>
<td>The USPSTF recommends the service. There is high certainty that the net benefit is moderate or there is moderate certainty that the net benefit is moderate to substantial.</td>
<td>Offer or provide this service.</td>
</tr>
<tr>
<td>C</td>
<td>The USPSTF recommends selectively offering or providing this service to individual patients based on professional judgment and patient preferences. There is at least moderate certainty that the net benefit is small.</td>
<td>Offer or provide this service for selected patients depending on individual circumstances.</td>
</tr>
<tr>
<td>D</td>
<td>The USPSTF recommends against the service. There is moderate or high certainty that the service has no net benefit or that the harms outweigh the benefits.</td>
<td>Discourage the use of this service.</td>
</tr>
<tr>
<td>I</td>
<td>The USPSTF concludes that the current evidence is insufficient to assess the balance of benefits and harms of the service. Evidence is lacking, of poor quality, or conflicting, and the balance of benefits and harms cannot be determined.</td>
<td>Read the clinical considerations section of USPSTF Recommendation Statement. If the service is offered, patients should understand the uncertainty about the balance of benefits and harms.</td>
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</table>

**Table 3 Studies of Cervical Medial Branch Radiofrequency Neurotomy**

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Study Type</th>
<th>Results</th>
</tr>
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<tbody>
<tr>
<td>Lord et al 1995</td>
<td>Clinical study</td>
<td>19 subjects and 21 joints treated, there were C2-3 zygapophysial joints (N=10) and additional C5, C6, C7 medial branches. Only 4 of 10 patients obtained relief following neurotomy for C2–C3 zygapophysial joint pain. Severe out of 10 patients who underwent lower cervical medial branch neurotomy obtained complete pain relief and were able to resume their activities of daily living and employment.</td>
</tr>
<tr>
<td>Lord et al 1996</td>
<td>RCT</td>
<td>24 patients with chronic neck pain underwent cervical radiofrequency. At 6 months, 7/12 (58%) patients in the active treatment group and 1/12 (8%) in the control group were pain-free. Median time of return of pain was 263 days in the treatment group compared to 8 days in the control group.</td>
</tr>
<tr>
<td>Shin et al 2006</td>
<td>Clinical study</td>
<td>28 consecutive patients underwent RFN. 69% of patients treated reported successful outcomes after 6 months, with 29% reporting complete pain relief.</td>
</tr>
<tr>
<td>Sapir and Gorup 2001</td>
<td>Clinical study</td>
<td>RFN was performed on 46 patients (29 litigants and 17 non-litigants). 25 patients were asymptomatic at one-year follow-up. The average time of return of pain of 50%, for those 21 patients who had pain before 12 months, was 8.3 ± 2.3 months.</td>
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<tr>
<td>McDonald et al 1999</td>
<td>Clinical study</td>
<td>28 patients with cervical facet joint pain diagnosed by comparative anesthetic blocks. RFN was performed, and repeated upon return of pain, with primary outcome measures of pain relief and duration of benefit. 18 of the 28 patients had greater than 3 months of complete pain relief. Median duration of pain relief was 421.5 days. The median duration of pain relief for all participants was 218.5 days. The study confirmed that lengthy, but not permanent relief from RFN is possible with repeat utilization.</td>
</tr>
<tr>
<td>MacVicar et al 2012</td>
<td>Clinical study</td>
<td>Cervical RFN was successful in 74% and 61% of patients in two separate sites. Clinical success was defined as at least 80% pain relief for 6 months. Average duration of relief from initial RFN was 17–20 months, and 15 months following repeat RFN treatments.</td>
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<tr>
<td>Barnsley 2005</td>
<td>Observational study</td>
<td>35 consecutive patients met all diagnostic criteria for cervical facet joint pain. For the RFN procedure, the electrode was introduced twice; along a parasagittal path to reach the medial branch as it crosses the lateral articular pillar, and at a 30 degree angle to the sagittal plane to reach the medial branch over the anterolateral aspect of the cervical pillar. Two or three lesions were made at each location to accommodate for possible variation in the course of the nerve (80 degrees Centigrade for 90 seconds). 26 (74%) patients treated obtained complete relief of pain for a mean duration of 35 weeks (8.7 months). Among the responders, 21 obtained complete relief in excess of 12 weeks.</td>
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</table>
In its most rigorous adoption, cervical medial branch RFN is a two-step procedure: (1) oblique insertion and (2) sagittal insertion, which allows for lesions at two separate sites along the trajectory increasing the area of the thermal lesion. The oblique pass is intended to place lesions over the anterior third of the superior articular process. The sagittal pass is intended to place lesions along the lateral aspect of the articular pillar in which the medial branch nerve is located (Figure 2).

Best Practices Summary
- Use of cervical RFN is primarily for symptoms of axial neck pain in the absence of radicular symptoms.
- Prior to RF of the cervical spine, other possible etiologies should be ruled out. The use of magnetic resonance imaging (MRI) and/or computerized tomography (CT) imaging is highly recommended, but x-rays can also be very useful. Although imaging can characterize degenerative changes, it is not a diagnostic for facet-mediated pain. Additionally, discogenic changes do not necessarily rule out the use of RFN.
- Physical examination and radiographic findings have no direct diagnostic or prognostic value in determining success of radiofrequency procedures.
- The efficacy of RFN is directly related to the rigor of diagnostic blocks performed as well as the use of proper technique for both diagnostic and neurotomy procedures.
- Evidence best supports the use of conventional thermal (60–80 degree centigrade) radiofrequency for 60–90 seconds. While the use of pulsed RF on the cervical medial branches has been reported, further studies are necessary.
- Multiple passes, and utilizing two separate approaches, may allow for neurotomy in a larger portion of the medial branch, resulting in improved pain relief with longer duration.
- If neck pain recurs, evidence supports the use of repeat neurotomy with reproducible efficacy. Although timing is highly variable, the procedure may be repeated as early as every six months.

Treatment Gaps & Future Research
Research and evidence gaps exist for cervical medial branch pulsed radiofrequency (PRF). PRF allows for the application of radiofrequency current at lower temperatures, minimizing the damage to surrounding tissue, nerves and/or vasculature. Further research on PRF is required to determine both the efficacy and the safety of this treatment modality, and how it compares to conventional RF. Although several studies have demonstrated the efficacy of both continuous and pulsed RF within the range of 6 to 12 months, more evidence is clearly needed.

Consensus Statement
Cervical medial branch radiofrequency neurotomy may be used for the treatment of axial neck pain when facet joints have been identified as the etiology of pain via diagnostic blocks. GRADE I A.
Thoracic Medial Branch Radiofrequency Neurotomy

Thoracic medial branch RFN is used for the treatment of chronic thoracic/midback pain. Since physical exam and radiographic techniques are variable and difficult to interpret in these instances, diagnosis of thoracic facet joint pain relies mainly on diagnostic injections. In those cases of positive diagnostic medial branch blocks, the patient may find long-term relief with thoracic RFN.

Current Available Evidence of Thoracic Medial Branch Radiofrequency Neurotomy

The literature surrounding the use of thoracic medial branch RFN is relatively scarce when compared to other levels of the spine. Interestingly, multiple modalities of RFN have been studied in the thoracic spine, albeit with low level of evidence. The published literature is summarized in Table 4.

Procedure Techniques

As with all ablation techniques, the effectiveness of the therapy is highly dependent on proper technique and appropriate lesioning of the thoracic medial branch nerves. Using either an anteroposterior (AP) or ipsilateral oblique fluoroscopic image, the radiofrequency cannula is directed in a caudal-to-cephalad direction and advanced towards the superolateral edge of the targeted thoracic transverse process. This is the site of the thoracic medial branch as it courses along the transverse process (Figure 3). The ablation is performed per standard protocol based on ablation modality.

Best Practices Summary

- The use of thoracic RFN is indicated for axial midback pain in the absence of radicular symptoms.

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Study Type</th>
<th>Results</th>
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<tbody>
<tr>
<td>Joo et al 2013</td>
<td>RCT</td>
<td>In the only thoracic RFA RCT to date, Joo et al studied the effects of alcohol ablation versus thermal RFA to the thoracic medial branches in patients with recurrent thoracic facet joint pain after prior thermal RFA treatment. Forty patients with recurrent thoracic facet joint pain following prior successful thermal RFA were randomly assigned to one of two groups: repeat thermal RFA or alcohol ablation. The recurrence rate of thoracic facet joint pain was assessed with NRS and ODI during the following 24 months. The results showed that alcohol ablation was significantly more likely to avoid pain recurrence than thermal RFA (defined as NRS &lt;7 and ODI &lt;22%; p&lt;0.000). The median effective time in the thermal RFA group was 10.7 months (range 5.4–24 months) and in the alcohol ablation group was 24 months (range 16.8–24 months). No significant complications were observed in either group.</td>
</tr>
<tr>
<td>Chang 2018</td>
<td>Prospective case series</td>
<td>Chang reported on the effects of pulsed RFA to the thoracic medial branches in 20 patients with chronic thoracic facet joint pain. Before enrollment, all patients experienced ≥80% temporary pain relief with a diagnostic thoracic MBB with 0.5 mL of 1% lidocaine. After successful diagnostic blocks, patients underwent pulsed RFA with settings at 5 Hz and a 5-millisecond pulse width for 360 seconds at 45 V. The NRS was recorded at baseline, and at 1-, 2-, and 3-months following treatment. Mean NRS was 6.0 at baseline and 4.0 at three months post-procedure (p&lt;0.05). Eleven of 20 patients (55%) reported pain relief ≥50% at three months post-procedure.</td>
</tr>
<tr>
<td>Rohof and Chen 2018</td>
<td>Retrospective study</td>
<td>A retrospective study of 71 patients whom following a successful diagnostic medial branch block underwent bipolar RFA. Eighty-two percent of patients had ≥50% pain relief at 12 months post-procedure. Baseline mean NRS was 7.75±1.25 which was decreased to 2.82±1.29 at 12 months post-procedure (mean 63.6%; p&lt;0.001). There were no complications reported.</td>
</tr>
<tr>
<td>Gungor and Candan 2020</td>
<td>Retrospective study</td>
<td>Cooled RFA (CRFA) is a newer RFA technique that can create a larger spherical lesion. Subsequently, there is an increased likelihood of including the medial branches in the thoracic spine within the created lesion. Gungor and Candan performed a retrospective study investigating the efficacy of CRFA in the treatment of thoracic facet-related pain. In this study, 40 CRFA procedures were performed on 23 individual patients following successful diagnostic medial branch blocks (≥80% temporary pain relief). The investigators assessed NRS scores pre- and post-treatment. At 6–12 month follow-up, mean pain relief was 37.6%.</td>
</tr>
</tbody>
</table>
Low-level evidence suggests that traditional RFA and bipolar RFA may provide longer lasting pain relief than pulsed RFA and CRFA, but given the study's methodological differences, it is impossible to draw any firm conclusions.

Given the location and anatomic variability of thoracic medial branches, patients should be counseled about the risk of pneumothorax. Physicians should have a sound understanding of thoracic fluoroscopic anatomy prior to performing thoracic RFN.

If thoracic pain recurs, evidence suggests that repeat RFN is effective. In the only RCT involving thoracic spine RFN, alcohol ablation was shown to be more efficacious than traditional RFA.

Treatment Gaps & Future Research
Overall, there is a paucity of quality research and publications addressing thoracic medial branch RFN. Thoracic facet-related back pain would benefit from high-quality research addressing basic questions, including comparison of different RFN modalities to conventional medical management and to each other in a prospective, long-term manner.

Consensus Statement
Thoracic medial branch radiofrequency neurotomy may be used for the treatment of thoracic/midback pain when facet joints have been identified as the etiology of pain via diagnostic blocks. GRADE II-3 C.

Lumbar Medial Branch Radiofrequency Neurotomy
Lumbar radiofrequency neurotomy (LRFN) is the second most common procedure performed in interventional pain practices in the United States. This procedure is also commonly referred to as lumbar medial branch nerve RFN. There has been a 130.6% increase in its utilization from 2007 to 2016. Low back pain (LBP) due to lumbar facet joint disease is often treated with LRFN. This diagnosis is prevalent in approximately 45% of lower back pain patients. While LRFN is one of the best studied interventional pain procedures, it is also one of the most debated procedures with constantly evolving evidence.
Current Available Evidence of Lumbar Medial Branch Radiofrequency Neurotomy

Diagnosing lumbar facet pain is challenging due to its heterogeneous etiology, variable clinical findings, radiographic and clinical course, and the multifactorial nature of low back pain. Patients generally complain of axial low back pain with or without radiation into the groin, buttocks, and/or thighs. Axial pain may worsen with extension, lateral bending, rotation, sitting, standing, walking or coughing. Consistent with other arthritic pain, this pain is often worsened by immobility. However, the published evidence does not find these symptoms to correlate well with the success of LRFN. Physical examination findings also have similarly mixed outcomes in the literature. The most studied and utilized physical examination techniques, including “facet loading” or paravertebral tenderness, have minimal ability to reliably predict the efficacy of LRFN. Hence, the literature illustrates that there are no pathognomonic signs with both clinical history and physical examination to reliably diagnose patients with facet-mediated pain or predict the success of LRFN.

Moreover, both MRI and CT scans have no specific findings that help diagnose lumbar facet mediated pathology as the exclusive source of pain. As such, the present standard in the diagnosis of lumbar facet mediated pain is diagnostic facet blocks, including both the use of intra-articular facet joint injections and medial branch nerve blocks. Medial branch blocks are considered more predictive for lumbar medial branch RFN success than intra-articular facet injections; however, both techniques have shown better predictive value compared to sham. The optimal volume of anesthetic injectate for diagnostic medial branch blocks recommended is <0.5 cc in order to improve specificity and reduce the risk of spreading to adjacent nerves or structures.

LRFN has been studied extensively with consensus guidelines indicating moderate to strong evidence. The efficacy of the lumbar medial branch RFN has been described in observational and RCT. Out of the six RCTs, three contained technical flaws in both patient selection and procedural technique, which yielded less reliable and therefore, non-generalizable findings. Of the literature that is published with acceptable methodology and generalizability, patient benefit from lumbar medial branch RFN is well documented. Table 5 summarizes LRFN studies.

Procedure Techniques

Successful treatment of lumbar facet pain by RFN is contingent on nerve coagulation by heat. The lumbar facet joints are paired, with each joint innervated by the medial branches from the dorsal rami of the same level, as well as the level above. Bogduk et al demonstrated the target for lumbar neurotomy to be the medial branches of the lumbar dorsal rami in their landmark study. In order to target these nerves appropriately, RF needles are placed along the posterior elements of the lumbar vertebral column, at the junction of the superior articular and the transverse processes of the vertebrae. At the L5 level, the dorsal ramus itself is amenable to neurotomy, which runs at the junction of the S1 superior articular process and the sacral ala.

The optimal electrode positioning has now been demonstrated to be parallel to the target nerves after the initial thought process that a perpendicular needle placement may be sufficient (Figure 4). While the studies varied in their design, the consensus was that the parallel placement of traditional electrodes is indeed the optimal approach for LRFN.

Motor and sensory testing is often suggested prior to RFN, in order to confirm appropriate and safe placement of the electrode tip. As such, injury to the ventral ramus or any other unintended structures may be avoided. This is a topic that has been widely studied and debated. There have been case studies with nerve root injury where motor stimulation was not performed. Given the limited data, even society guidelines are mixed on utilizing sensory and motor stimulation prior to lumbar medial branch RFN. Finally, a recent expert consensus statement recommended sensory stimulation with single-level lesions, with low level of certainty and inconclusive evidence with multiple lesions. Motor stimulation is recommended for both safety and effectiveness purposes.

Best Practices Summary

- Use of lumbar radiofrequency neurotomy (LRFN) is primarily for symptoms of low back pain in the absence of radicular symptoms.
- Prior to LRFN, other possible etiologies by way of either MRI or CT imaging should be ruled out, although imaging is not itself a diagnostic for facet-mediated pain. Discogenic changes do not necessarily rule-out the use of LRFN.
- Medial branch blocks are generally considered more predictive for success with LRFN than intra-articular facet injections.
In addition to the success of the initial LRFN, the efficacy of repeat LRFN with pain recurrence has been shown to be highly reproducible.

- Optimal electrode positioning is parallel to the target nerves, although more evidence is necessary.
- Motor and sensory testing is recommended prior to LRFN, in order to confirm appropriate placement of the electrode tip and to avoid lesioning of any other unintended structures including the ventral ramus.
- Evidence best supports the use of conventional thermal (60–80 degree centigrade) radiofrequency for 60–90 seconds. While the use of pulsed RF on the lumbar medial branches has been reported, further studies are necessary.

## Treatment Gaps & Future Research

Similar to the cervical region, pulsed radiofrequency (PRF) for the lumbar spine is not covered extensively in the literature. Further research in various forms of RF is needed to determine the efficacy and safety of these treatment modalities. Furthermore, additional research is clearly needed to confirm if a parallel needle placement is indeed the most optimal option, particularly in light of other modalities, such as cooled RF.

## Consensus Statement

Lumbar medial branch radiofrequency neurotomy may be used for the treatment of axial low back pain when facet

### Table 5 Summary of LRFN Studies

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Study Type</th>
<th>Results</th>
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<tbody>
<tr>
<td>Lee et al 2017</td>
<td>Meta-analysis</td>
<td>Included 7 randomized controlled trials (RCT). Concluded RFN groups had significant improvement in pain scores at one-year post-procedure compared to sham and sacroiliac joint pain compared to conservative, non-interventional approaches, revealed a significantly greater improvement in ODI, pain scores and quality of life in the RFN arm compared to the control arm.</td>
</tr>
<tr>
<td>Shih et al 2020</td>
<td>Meta-analysis</td>
<td>Radiofrequency offer significant improvement in lumbar facet joint or sacroiliac joint pain for up to 12 months compared with baseline level of pain.</td>
</tr>
<tr>
<td>Maas et al 2015</td>
<td>Cochrane review</td>
<td>Review of RFN for chronic low back pain evaluated 23 studies; 12 studies examined suspected facet joint pain. The authors found no high-quality evidence to suggest that RF denervation provides pain relief or functional improvement for patients with CLBP.</td>
</tr>
<tr>
<td>Poetscher et al 2014</td>
<td>Literature review</td>
<td>This review examined nine studies and concluded there was low-to-moderate evidence favoring lumbar medial branch RFN for pain control.</td>
</tr>
<tr>
<td>Boswell et al 2015</td>
<td>Systematic review</td>
<td>The available evidence is Level I for lumbar facet joint nerve blocks with the inclusion of 17 studies with dual diagnostic blocks, with at least 75% pain relief with an average prevalence of 16–41% and false-positive rates of 25%-44%.</td>
</tr>
<tr>
<td>Falco et al 2012</td>
<td>Systematic review</td>
<td>11 randomized trials and 14 observational studies met inclusion criteria for methodological quality assessment. The review concluded that evidence for radiofrequency neurotomy is good and fair-to-good for short- and long-term improvement; whereas the evidence for intraarticular injections and pulsed radiofrequency neurotomy is limited.</td>
</tr>
<tr>
<td>Juch et al 2017, and van Kuijk et al 2018</td>
<td>Randomized control trial</td>
<td>This study found no efficacy for lumbar medial branch RFN. However, this study has been widely criticized as rife with flaws including significant issues with the technical aspects such as electrode placement and orientation, statistical analysis, and non-stringent selection criteria which leads the study to lack any generalizability.</td>
</tr>
<tr>
<td>Conger et al 2019 and Tekin et al 2007</td>
<td>Literature review</td>
<td>In addition to success of the initial lumbar medial branch RFN, the efficacy of repeat RFN, has very high success rates in the literature, with 3 retrospective studies and 1 prospective study showing a success rate &gt;85% for repeated RFN treatment with duration of pain relief between 10–12 months.</td>
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</table>
joints have been identified as the etiology of pain via diagnostic blocks, GRADE I A.

**Lateral Sacral Branch Radiofrequency Neurotomy**

Early studies brought attention to the sacroiliac joint as a pain generator through reproduction of symptoms by way of provocative maneuvers and diagnostic/therapeutic injections into the sacroiliac joint (SIJ). Laslett et al found provocative maneuvers to be helpful but lacked sensitivity. There is no evidence to support any provocative SIJ maneuver as pathognomonic for SIJ pain; hence, the use of diagnostic SIJ blocks utilizing local anesthetics with or without steroids has evolved as the current standard. Although, initially, such injections were done blindly, image-guidance has substantially improved the accuracy of this procedure. However, therapeutic intra-articular injections only provide temporary relief for most patients. Recalcitrant cases led to the advent of other treatment options for SIJ dysfunction. RFN emerged as the most viable choice given its known evidence and effectiveness in treating facet-mediated pain.

Histological analysis of the sacroiliac joint has verified the presence of nerve fibers within the joint capsule and adjoining ligaments. The sacroiliac joint is thought to receive its innervation from the ventral rami of L4 and L5, the dorsal rami of L5, S1, S2, and S3, as well as the superior gluteal nerve. There is an additional ventral innervation of the sacroiliac joint that is inaccessible for denervation.

Sacral lateral branch neurotomy addresses the posterior sacroiliac structures, which includes both the posterior joint and ligaments. Therefore, the response to intra-articular sacroiliac joint injections, per se, does not necessarily correlate to radiofrequency success. However, blocking the innervation lateral to the foramen at different depths has been shown to be more effective. However, there are no placebo-controlled trials for lateral sacral nerve blocks to confirm this thought process. This is a treatment gap in appropriately selecting patients for sacral lateral branch neurotomy.

**Current Available Evidence of Lateral Sacral Branch Radiofrequency Neurotomy**

Two controlled and two pragmatic RFN studies of the lateral sacral branch are described in Table 6 with the remaining lateral sacral branch radiofrequency neurotomy studies being observational.

**Procedure Techniques**

The variability in the position of the posterior lateral sacral branches makes diagnostic blocks a technical challenge. As such, monopolar lesions may not necessarily reliably capture the nerve, which has led to the advent of cooled or bipolar RF lesion strategies. There are various studies comparing different forms of RFN of the SIJ; however, results are inconsistent and the study designs have variable patient selection lending to discrepancies in neurotomy techniques used. For this reason, monopolar lesions continue to be used as a treatment option; however, multiple lesions along the suspected trajectory of the lateral branches are recommended.

Roberts et al conducted one of the most comprehensive cadaveric anatomical studies to date on the anatomic location of the lateral sacral branches. The lateral branches were dissected out, and radiopaque wires were directly sutured to them. Anterior-posterior fluoroscopic imaging through the S1 superior endplate was obtained. Lesions by 17 versions of seven current sacroiliac joint RF techniques were mapped on fluoroscopic images. The techniques are meticulously described within this manuscript and were compared for:

1) Percentage of lateral branches captured
2) Percentage of SIJ specimens in which 100% of lateral branches were captured
3) Percentage of lateral branches that would not be captured at each level.

The results demonstrated that bipolar techniques captured a greater percentage of the lateral branches than traditionally used monopolar techniques. Furthermore, monopolar capture rates ranged from 49.6% to 64.6%, likely due to the small lesion size created by this technique.

Regarding the locations of the actual thermal lesion, it was found that the use of the palisade technique (Figure 5) was the most optimal in capturing the lateral branches. The
Table 6 Sacroiliac Joint Radiofrequency Neurotomy Studies

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Type of RF</th>
<th>Diagnostic Criteria</th>
<th>Follow-Up (Months)</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buijs et al 2004&lt;sup&gt;79&lt;/sup&gt;</td>
<td>Unipolar</td>
<td>&gt;50% relief with single SIJB</td>
<td>3</td>
<td>24 patients (56%) with the study reported at least 50% pain relief. 10 patients (23%) had complete pain relief.</td>
</tr>
<tr>
<td>Burnham et al 2007&lt;sup&gt;79&lt;/sup&gt;</td>
<td>Bipolar</td>
<td>Not specified</td>
<td>1, 3, 6, 12</td>
<td>9 patients were included in this pilot study. Median pain score decreased from 8/10 to 3.5/10 at 1 and 3 months and 4.5/10 at 6 and 12 months. The study showed reductions of back and leg pain frequency and severity, and analgesic intake. 8 of 9 subjects were satisfied overall.</td>
</tr>
<tr>
<td>Cheng et al 2013&lt;sup&gt;80&lt;/sup&gt;</td>
<td>Cooled or unipolar</td>
<td>≥50% relief with each of two SIJBs</td>
<td>6</td>
<td>58 patients underwent cooled RF technique and 30 unipolar technique. At six-month follow-up, 28 of the patients (32%) had &gt;50% pain relief with no significant difference between either cooled or unipolar.</td>
</tr>
<tr>
<td>Cohen et al 2003&lt;sup&gt;81&lt;/sup&gt;</td>
<td>Unipolar</td>
<td>80% relief SIJB, 50% after SLBBs</td>
<td>9</td>
<td>8 of 9 patients (89%) reported &gt;50% relief of pain and two of the nine (22%) reported total pain relief.</td>
</tr>
<tr>
<td>Cohen et al 2007&lt;sup&gt;82&lt;/sup&gt;</td>
<td>Cooled and Unipolar</td>
<td>≥75% relief with an intra-articular SIJ with bupivacaine and steroid.</td>
<td>1, 3, 6</td>
<td>At 1-, 3- and 6-months post-procedure, 11, 9, and 8 out of 14 patients treated with active cooled RFN and 7, 6 and 4 out of 11 patients treated with unipolar RFN had at least 50% pain relief based on the NRS scale. Only 2 out of 14 sham patients achieved treatment success at the one-month follow-up, and none at 3 or 6 months. The authors used SIJ injections, rather than sacral lateral branch blocks as the diagnostic measure, which is considered a limitation to this study.</td>
</tr>
<tr>
<td>Cohen et al 2008&lt;sup&gt;83&lt;/sup&gt;</td>
<td>Cooled or Unipolar</td>
<td>≥50% relief with each of two SIJBs</td>
<td>6</td>
<td>40 of 77 patients (52%) reported &gt;50% relief of pain at 6 month follow-up.</td>
</tr>
<tr>
<td>Dutta et al 2018&lt;sup&gt;84&lt;/sup&gt;</td>
<td>Pulsed</td>
<td>≥80% relief with an intra-articular SIJ injection with a local anesthetic</td>
<td>1, 3, 6</td>
<td>At 1, 3 and 6 months 15, 13, and 13 out of 15 patients had a ≥50% reduction in numerical pain score, a positive Global Perceived Effect (GPE), and a 5-point decrease in ODI score. At 3 and 6 months, the mean NRS score was higher in the IA steroid group, with statistical significance. The results suggest that though IA steroid injections may be effective, the use of PRF may be superior in overall improvement and duration of pain relief.</td>
</tr>
<tr>
<td>Ferrante et al 2001&lt;sup&gt;68&lt;/sup&gt;</td>
<td>Bipolar</td>
<td>Not specified</td>
<td>6</td>
<td>12 of 33 patients (36%) reported ≥50% pain relief for 6 months.</td>
</tr>
<tr>
<td>Ho et al 2013&lt;sup&gt;85&lt;/sup&gt;</td>
<td>Cooled</td>
<td>&gt;50% pain relief with SIJ steroid injection</td>
<td>1, 3, 6, 12, 24</td>
<td>15 of 20 patients showed a significant reduction in pain (reduction of at least 3 points on Numeric Rating Scale). Mean for pain was reduced from a NRS of 7.4 ± 1.4 to 3.1 ± 2.5. Mean Patient Global Impression of Change was “improved” (1.4 ± 1.5), and Global Perceived Effect was reported to be positive in 16 patients at two year follow-up.</td>
</tr>
<tr>
<td>Kapural et al 2008&lt;sup&gt;86&lt;/sup&gt;</td>
<td>Cooled</td>
<td>≥50% relief with each of two SIJBs</td>
<td>3–4</td>
<td>13 of 27 patients (48%) reported at least 50% pain reduction at follow-up and three (11%) reported complete relief.</td>
</tr>
</tbody>
</table>

(Continued)
peri-foraminal technique (Figure 6) was found to be an adequate alternative but requires multiple lesions surrounded in the lateral aspect of the S1, S2, S3 neural foramina.

**Best Practices Summary**

- Diagnosing SIJ pain consists of a physical examination including multiple provocative maneuvers (distraction, side thigh thrust, Gaenslen, compression and sacral thrust). The best predictive power is found with positive provocation utilizing 2 of 4 selected tests. If none of the six provocation tests reproduce pain symptoms, SIJ pathology may be ruled-out. Ober’s and Patrick’s tests are secondary options, though these have not been extensively studied.

- Prior to RF of the lateral sacral nerves, other possible etiologies must be ruled out by way of imaging. Radiographs may show degenerative changes and inflammation within the SIJ but are not diagnostic for SIJ pain.

**Table 6 (Continued)**

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Type of RF</th>
<th>Diagnostic Criteria</th>
<th>Follow-Up (Months)</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Karaman et al 2011</td>
<td>Cooled</td>
<td>&gt;75% relief with each of two SIJBs</td>
<td>6</td>
<td>12 of 15 patients (80%) reported at least 50% reduction in pain scores.</td>
</tr>
<tr>
<td>Patel et al 2012 and 2016</td>
<td>Cooled</td>
<td>≥75% relief with each of two SIJBs</td>
<td>3, 6, 9, 12</td>
<td>At 3, 6, 9, and 12 months post-procedure 16/34, 13/27, 20/25, and 10/25 had at least 50% pain relief based on the NRS scale and either a 10-point increase in SF-36BP or 10-point decrease in ODI. At 3 months 2 of the 17 sham patients achieved treatment success. 16 of the sham patients then crossed over to cooled RFN. At 1, 3, and 6 months post-procedure for the cross-over group 7, 7, and 6 of the 16 subjects achieved treatment success. At 12 months post-procedure, 6 achieved treatment success.</td>
</tr>
<tr>
<td>Schmidt et al 2014</td>
<td>Unipolar and Bipolar (Simplicity)</td>
<td>&gt;50% pain relief with SIJ steroid injection</td>
<td>1, 5, 6, 12</td>
<td>Out of the 77 radiofrequency treatments included, 71.4% (N = 55) reportedly had ≥50% pain relief. At 6 months, 54.5% of the treatments continued to have a successful outcome.</td>
</tr>
<tr>
<td>Speldewinde et al 2011</td>
<td>Unipolar</td>
<td>&gt;80% relief with each of two SIJBs</td>
<td>2</td>
<td>20 patients were followed by phone 6–36 months after RF treatment. Twelve patients of 20 (75%) reported at least 50% relief for 2 months. Seven patients (44%) reported having had complete relief.</td>
</tr>
<tr>
<td>Stelzer et al 2013</td>
<td>Cooled</td>
<td>&gt;50% relief with single SIJB</td>
<td>&gt;4</td>
<td>77 of 126 (61%) of patients reported ≥50% pain relief for &gt;4 months.</td>
</tr>
<tr>
<td>Yin et al 2003</td>
<td>Unipolar</td>
<td>&gt;50% relief with each of two SIJBs</td>
<td>6</td>
<td>9 of 14 patients (64%) had &gt;50% decrease in visual integer pain score and 36% had complete relief.</td>
</tr>
<tr>
<td>Zheng et al 2014</td>
<td>Bipolar</td>
<td>≥80% relief with an intra-articular SIJ injection with 40 mg Triamcinolone in 0.5% bupivacaine</td>
<td>3, 6</td>
<td>At 3 and 6 months post procedure, 39 and 37 (out of 76 patients with ankylosing spondylitis) had at least 20% and improvement of 1 unit (on a 0–10 scale). Pain reduction found to be greater in bipolar RFN arm at 12 and 24 weeks in comparison in the celecoxib arm, (P &lt; 0.0001). Bipolar RFN was also more effective in improving physical function and spinal mobility (P &lt; 0.05). The authors concluded that RFN is an effective manner to treat SIJ point in patients with ankylosing spondylitis.</td>
</tr>
</tbody>
</table>
Contraindications for RFN include sacral fracture, tumor, radiculopathy, infection and coagulopathy. A diagnostic block for the lateral sacral branches is highly recommended prior to performing RF, even in cases when previous sacroiliac joint injection was performed. While limited evidence exists regarding diagnostic cutoff, recommendation is for 50% or greater reduction in pain prior to advancing to RF. Multisite and multi-depth technique is recommended to appropriately select RF candidates.

The use of bipolar or cooled RFN is ideal to ensure that the lateral branches are captured with a larger sized lesion. Using a lateral x-ray view, place six (6) cannulae starting superior to S1 and stopping inferior to S3. Cannulae should be parallel to each other, spaced 10–12 mm from the next and perpendicular to the sacral surface. The use of monopolar RFN has been shown to be effective clinically, but multiple RF cannula passes or the use of periforaminal technique is recommended to create larger sized lesions.

When using bipolar or monopolar RFN, temperature of 80 degrees Centigrade for 60–90 seconds per cycle is recommended. The use of a large strip lesion is able to capture most of the lateral branches with a more medial placement of the RF probe.

Treatment Gaps & Future Research
The main limitation found within the literature is the heterogeneity of the studies involved. Most studies utilized intra-articular joint injections to select patients for sacral lateral branch RFN. As identified by Dreyfuss et al, the use of diagnostic nerve blocks in the posterior sacral nerve supply is more appropriate and prognostic for outcomes of RF. Not only is further research needed for the use of diagnostic sacral nerve blocks but also with neurotomy itself. Studies used conventional RF, pulsed RF, and cooled RF, with variable parameters and procedural times. The present literature suggests the clinical efficacy of unipolar, bipolar, cooled and pulsed RF; however, studies are limited and are of fair quality at best.

Consensus Statement
Lateral sacral branch radiofrequency neurotomy may be used for the treatment of posterior sacral ligament and joint pain following positive response to appropriately placed diagnostic blocks. GRADE II-1 B.

Radiofrequency Neurotomy for Peripheral Joints
Large joint RFN has been used predominantly to treat patients with end-stage osteoarthritis (OA). It is estimated that as many as 32 million Americans are affected by OA.31,32 Large joint RFN has been described for the treatment of hip pain, knee pain, and more recently, the
shoulder pain. As the availability of RFN has evolved, so to have the applications. RFN can also be utilized for treating rheumatologic joint disease, post-arthroplasty pain, metastatic disease, osteonecrosis, and chronic infection.  

The capsular and intra-articular innervation of large joints is complex. The afferent nerves often originate from multiple motor-sensory nerves, and there are variances in its procedural execution. Due to this complexity, it is imperative that the treating physician understands the regional anatomy of the target area in order to avoid complications.

**Current Available Evidence of Knee Joint Radiofrequency Neurotomy**

A multitude of studies, including randomized clinical trials (RCT) and meta-analyses, have examined the effectiveness of radiofrequency neurotomy of the knee (Table 7).

**Procedure Techniques**

Recent meticulous dissection studies, including those by both Tran et al and Fonkoue et al, have revealed evidence for both the use of classical and alternate targets for knee RFN.115–120 Tran et al, in two different cadaver studies demonstrated that the articular branches to the knee terminated in their respective quadrants with minimal overlap, though each quadrant could have multiple contributing nerves in addition to the classic superolateral (SL), superomedial (SM), and inferomedial (IM) genicular nerves.116 In a 2020 study, the authors used ultrasound-guided RF cannulae to target the classic SL, SM, and IM nerves and then dissected around the cannulae creating a three-dimensional digitized model showing that classical landmarking can be effective for genicular RFN, though may not capture all the articular innervating branches of the knee.117 In 2019 and 2020, Fonkoue et al used dye and subsequent dissection in 21 and 14 cadavers, respectively, to identify the most effective landmark positions for fluoroscopy-guided genicular nerve blockade and RFN. The results demonstrated the increased effectiveness of revised targets that are, in general, more posterior in the knee than the classical landmarks.118–120

Traditionally, fluoroscopy has been used to guide needle placement for RFN, predominantly targeting the genicular nerves as their course along the periosteum (Figure 7). However, more recently ultrasound (US) has been shown to be an effective method of image guidance for RF knee procedures. A systematic review and meta-analysis on US

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**Table 7** Summary of Major Studies on RFN of Nerves That Supply the Knee

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Study Type</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bhatia et al 2016</td>
<td>Systematic review</td>
<td>This review identified 13 publications, representing 329 patients, which met criteria for inclusion. Of the thirteen included studies, six represented conventional or cooled RFN treatments. Conclusions from these studies included a high rate of success in alleviating chronic pain of the knee joint from 3 to 12 months, improvement in function, in the absence of serious adverse events.</td>
</tr>
<tr>
<td>Choi et al 2011</td>
<td>Randomized control trial</td>
<td>38 patients were randomized into RFN or sham RF. The RFN group had significantly lower pain scores compared to control at 4 and 12 weeks, and functional improvement. (Note that this article was cited in Bhatia et al 2016.)</td>
</tr>
<tr>
<td>Jamison and Cohen 2018</td>
<td>Literature Review</td>
<td>This review identified 9 relevant randomized or comparative clinical trials, representing 592 patients; two studies previously reported in Bhatia et al, and seven new trials. Eight trials specifically compared knee RFN to sham or other treatments such as steroid injections. One study examined intra-articular pulsed RF. This review concluded that there was significant pain reduction and improvement in function as measured time points from 3 to 12 months. Efficacy of diagnostic blocks was questionable. Limitations to this review included considerable variation in the RF targets, technique, as well as selection criteria.</td>
</tr>
<tr>
<td>Hong et al 2019</td>
<td>Meta-analysis</td>
<td>The review consisted of existing RCTs indexed on other databases outside of English-based indices (eg, PubMed) previously not included, such as Chinese-based indices (eg, CNKI). This meta-analysis included 4 studies from the prior reviews and 8 new RCTs, representing 841 patients. All studies described a control group. Seven studies examined RFN of the genicular nerves. Notably, this review excluded patients that were status-post knee arthroplasty and arthroscopic surgery. Results showed decreased pain scores at 1 week, 1 month, and 3 months.</td>
</tr>
</tbody>
</table>
by Huang et al identified 8 publications with 256 patients in 2020. Three of the studies were RCTs and the remaining studies were prospective studies. Meta-analysis revealed US-guided RF knee procedures to result in a pooled mean difference in pain score of $-4.2$ (NRS scale) and a $23.2$ improvement in WOMAC score compared to baseline before RF. An RCT by Sari et al directly compared US-guided RF to fluoroscopic-guided RF, finding similar outcomes between the two modalities. Subgroup analysis found RFN of the genicular nerves as examined by four studies resulted in even greater pain reduction of a mean difference of $-4.9$ (compared to intra-articular modalities such as pulsed-RF). Adverse events reported were infrequent, mild and transient. Overall, US was found to be an effective, safe, non-ionizing, and technically achievable method for RF procedures of the knee, and that the genicular nerves are the preferred targets.

**Best Practice Summary**
- Radiofrequency ablative technologies for the nociceptive sensory innervation of the knee have been shown to be an effective therapy for chronic knee pain due to conditions, such as osteoarthritis and post-surgical pain.
- Utilization of RFN on the knee obviates a comprehensive knowledge and understanding of the anatomical innervation of the knee, experience with fluoroscopy or ultrasound, and knowledge regarding the principles surrounding RFN.
- Evidence-based parameters for electrode settings are the use of 70–80°C for 90–180 seconds.
- In terms of technique, targeted genicular nerves for conventional and cooled RFN include the SM, SL, and IM nerves. Due to the variable anatomy of these branches, larger lesion sizes increase the likelihood of success.
- Pre-ablative diagnostic blocks with a low volume anesthetic may enable more accurate prognostication of the analgesic response to RFN.
- Avoiding unnecessary injury or inadvertent neurotomy of motor nerves, non-targeted sensory nerves, blood vessels, and other non-targeted anatomic structures is essential. Motor testing prior to ablation should be considered.

**Treatment Gaps & Future Research**
In the past several years, there have been multiple studies, including high-quality RCTs, evaluating the efficacy of genicular RFN. These studies have included pain scales and validated measures of function at up to 12 months of follow-up. Further studies will enable the ideal use of this treatment modality. Further research topics should include the examination of demographic factors (eg BMI, gender, severity of OA), improve patient selection, the utility of prognostic blocks and repeat RFN. In general, the ideal timing of RFN in the setting of a multimodal management algorithm should also be established. Long-term outcomes, beyond a year, will also be important due to the chronic nature of the conditions typically treated with this modality.

**Consensus Statement**
Genicular nerve radiofrequency neurotomy may be used for the treatment of osteoarthritis related and post-surgical knee joint pain. GRADE II-1 B.

**Current Available Evidence of Hip Joint Radiofrequency Neurotomy**
The innervation of the hip was first studied and described in the mid-nineteenth century. German anatomist Rüdinger was a pioneer in describing the innervation of the large joints, including the shoulder, knee, and hip. Rüdinger’s work laid the framework for surgical denervation...
techniques and interventional pain procedures. Further study and understanding of the articular branches of the hip was elucidated by Gardner in the 1940s.\textsuperscript{125} The evolution of RFN techniques has renewed the interest in hip pain treatment with RF techniques that maximize benefit while avoiding complications. Recent cadaveric dissections have shown notable anatomic variability with the location of articular branches from femoral (FN), obturator (ON), and accessory obturator nerves.\textsuperscript{126}

The anteromedial capsule of the hip is supplied by the branches of the obturator nerve, while the anterior capsule is innervated by branches of the femoral nerve. The sciatic nerve provides articular branches to the hip, posteriorly. Current ablative techniques focus on targeting the anterior hip. The accessory obturator nerve (AON) has been studied in cadaveric dissections and is present in a minority of individuals (10–30%).\textsuperscript{127,128} The contribution of the AON (if present) can vary. The neurovascular bundle (femoral artery, vein, and nerve) is vulnerable to potential injury when targeting the obturator branches. Traditionally, this risk was mitigated by approaching the inferior aspect of the ischium from medial or lateral trajectories with the goal of passing underneath the bundle. The challenge with fluoroscopic guidance is that patients can have anatomic variations of the local neurovascular structures, and needle approaches must be considered in order to optimize outcomes while decreasing adverse events.

### Current Available Evidence of Hip Radiofrequency Neurotomy

The goal of hip RFN is to target major articular innervation to the anterior hip joint. RFN of hip articular branches was first described by Okada in 1981.\textsuperscript{128} He subsequently reported successful results in a series of 15 patients with neurotomy of the articular branches anteriorly (femoral and obturator) and posteriorly (sciatic).\textsuperscript{129} These early studies led to more robust investigation and interest in RFN for refractory hip pain. As hip RFN techniques evolved, neurotomy of both the femoral and articular branches became standard practice.

With the evolution of ultrasound technology and increased frequency of use in pain medicine, this modality is increasingly being used to increase accuracy and avoid complications. By no means does ultrasound replace fluoroscopy for hip RFN. Rather, it is often used in conjunction with fluoroscopy to aid in needle placement. Ultrasound may allow the proceduralist to avoid the femoral neurovascular bundle, decreasing the incidence of bleeding and nerve injury. These studies also show that it is imperative that motor testing is performed prior to neurotomy, as repositioning of cannula at the incisura acetabuli may be necessary to avoid the ON as it courses through the obturator foramen. Studies of hip RFN are summarized in Table 8.

### Procedure Technique

Careful consideration and shared risk decision-making must be taken in regard to a patient’s comorbidities and anticoagulation status. A paucity of data exists regarding best practices around anti-coagulation management for hip RFN. Many practitioners do not interrupt anticoagulants for diagnostic blocks and subsequent RFN, while others may stop these medications for the RFN portion or even both procedures. Larger cannula size may increase the risk of bleeding, particularly when targeting the ON branches, which run in close proximity to the femoral neurovascular bundle.

Patient body habitus and mobility are also important considerations. In the morbidly obese (BMI >40), it can be extremely difficult to place needles comfortably and easily. The patient must be able to tolerate being in a supine position for a prolonged period of time and have the ability to extend at the hip. If a patient is unable to tolerate these positions, RFN is likely to be challenging or not possible. At a minimum, fluoroscopy is needed to ensure proper identification of hip bony landmarks and to navigate challenging anatomy. Multiple studies have described a combined fluoroscopy and ultrasound technique that may improve safety and decrease the risk of vascular and/or nerve injury.\textsuperscript{132–135}

There is a paucity of data when it comes to prognostic value of hip articular branch blocks. Despite this, it is recommended that patients undergo a diagnostic block before being considered for neurotomy. This is largely due to the multitude of pain generators that may cause pain in the hip area including tendinopathy, bursitis, and lumbosacral pathology. A minority of studies have used relief following intraarticular hip injections as selection criteria for RFN.\textsuperscript{130,136}

Given the anatomic variability of articular branches, larger needles (21G or larger) are preferred. With conventional RFN, cannulae with a 10 mm active tip or larger is optimal. Cooled RFN is an alternative technique that may be
superior given its increased volume of ablated tissue. Prior to neurotomy, we recommend that motor testing be performed to rule out proximity to the obturator or femoral nerves. Sensory testing can also be helpful in confirming proximity to the articular branches. Lastly, multiple neurotomy lesions of the ON and FN articular nerves should be considered given their unpredictable course.

**Best Practices Summary**

- At a minimum, fluoroscopy is necessary to ensure proper identification of hip bony landmarks and to navigate challenging anatomy. More investigation is required to determine the optimal use of ultrasound in hip denervation. Ultrasound is often used in conjunction with fluoroscopy for optimal needle placement.

- Patient body habitus and mobility are also important considerations. In the morbidly obese (BMI >40), it can be extremely difficult to place needles comfortably and easily. Patients must be able to lie supine with the hip in neutral extension.

- Diagnostic blocks should be performed prior to neurotomy but data implying their prognostic value is limited.

- Neurorotomy of both the femoral and articular branches has become standard practice using lateral to medial approaches with the patient in the supine position (Figure 8).

- Evidence best supports the use of conventional RF at 80 degrees for 90 seconds, although efficacy for cooled-RF has been shown in recent cohort and case series.

### Table 8 Summary of Studies on Hip RFN

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Study Type</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Short et al 2018</td>
<td>Research article</td>
<td>Anatomical study of hip innervation in cadavers delineating the variable innervation of the anterior hip. Thirteen hemi-pelvises were dissected and the contribution of the femoral (FN), obturator (ON), and accessory obturator nerves were examined. The FN and ON contributed to the capsular innervation in all specimens. The most consistent landmark was the inferomedial acetabulum (radiographic “teardrop”).</td>
</tr>
<tr>
<td>Kawaguchi et al 2001</td>
<td>Retrospective cohort study</td>
<td>A follow-up study looking at RFN feasibility focused primarily on the obturator articular branches, and to a lesser extent the femoral articular branches. 14 patients with various diagnoses leading to chronic hip pain received either an intraarticular injection or obturator articular block with a local anesthetic. Those with groin and thigh pain had only obturator articular neurotomy. Those who also had trochanteric (lateral) pain also received a second neurotomy for the femoral articular branches. 12 patients (86%) reported at least 50% relief of pain for 1 to 11 months.</td>
</tr>
<tr>
<td>Rivera et al 2012</td>
<td>Clinical trial</td>
<td>18 patients with hip pain (16 with OA and 2 with chronic post-surgical pain) received diagnostic articular blocks with ropivacaine. To lesion the obturator sensory branches, a cannula was introduced just medial to the femoral artery, below the inguinal ligament (medial approach) or 3 cm lateral to the femoral artery (lateral approach), at a 70° angle with the sagittal plane. The needle was advanced to the junction between the ischium and the pubis corresponding to the incisura acetabuli. For the femoral branches, the cannula was inserted via an anterolateral approach with the final position of the tip below anterior inferior iliac spine along the superolateral hip. RFN was performed at each location at 90°C for 90 seconds. Mean VAS scores were 9.52 and 6.35 before radiofrequency and at 6-month follow-up, representing approximately a 38% reduction.</td>
</tr>
<tr>
<td>Kapural 2018</td>
<td>Retrospective cohort study</td>
<td>52 patients undergoing cooled RFN for chronic hip pain with both fluoroscopy and ultrasound guidance. Investigators were able to quantify the close proximity that exists between the ON and the neurovascular bundle. In 21 of the patients undergoing RFN, the distance was less than 1 cm (range 0.5–1.9 cm, with median 0.8). The pain score improved went from 7.61 ± 1.2 at baseline to 2.25 ± 1.4 after the RFN (P &lt; 0.01).</td>
</tr>
<tr>
<td>Naber 2019</td>
<td>Case series</td>
<td>This case series of 11 patients undergoing cooled RFN of hip looked at the efficacy of cooled RFN with combined ultrasound and fluoroscopic imaging in patients with avascular necrosis (AVN). Five patients claimed more than 50% of pain relief. The average time interval of greater than 50% of pain relief from the cooled RFN was 70–250 days.</td>
</tr>
</tbody>
</table>
Larger lesion size is preferred given the anatomic variability of ON and FN articular branches. Multiple neurotomy lesions at each site will likely increase the procedure's success.

**Treatment Gaps & Future Research**

Further investigation is needed to determine the optimal patient selection and protocol for improving outcomes. A review of the available literature found variable outcomes with regard to pain reduction ranging from 8 days to 3 years. Anatomical variations of the FN, ON, and AON are certainly factors in the success of neurotomy. Additionally, RFN modality (cooled vs traditional), number of lesions, and cannula size are also important.

**Consensus Statement**

Hip joint radiofrequency neurotomy targeting the obturator and femoral nerve branches may be used for the treatment of hip joint pain following diagnostic blocks. GRADE II-1 B.

**Occipital Nerve Radiofrequency Neurotomy**

Occipital RFN is used for the treatment of chronic occipital neuralgia (ON) and posterior headache syndromes, including refractory migraine. ON is considered to be rare, while also challenging to diagnosis and treat. Trauma, irritation, and entrapment of the greater (GON), lesser (LON) and/or third occipital (TON) nerves have been suggested as the etiology of ON, which may present as lancinating pain along the respective nerve distributions. Physical examination findings include occipital tenderness, palpable trigger points at the emergence of the greater occipital nerve or in the distribution of C2, and positive Tinel's sign over the occiput at the location of the greater and lesser occipital nerves. In lieu of diagnostic imaging findings, positive response to landmark guided or image-guided diagnostic block of the occipital nerves is confirmatory for the diagnosis, while the mainstay of treatment is considered to be a landmark or image guided corticosteroid injection. While RFN has been utilized in the treatment of ON, it has not garnered universal acceptance. Various modalities of RFN have been described including CRF, PRF, and WCRF.
Current Available Evidence of Occipital Nerve Radiofrequency Neurotomy

Recent publications are summarized in Table 9.

Procedure Techniques
Published studies describe a myriad of techniques for occipital nerve RFN. Most commonly landmark-based and fluoroscopic techniques are described, with scant literature describing the utilization of ultrasound. The most robust techniques describe multiple needles targeting the GON and LON utilizing sensory stimulation relying on patient feedback for localization. The recommendation to place the needle in parallel line as opposed to perpendicular plane is in keeping with recommendations at other anatomic sites. Multiple cycles of PRF at 42 degrees Celsius and single lesions of CRF have been described.

Best Practices Summary
- Use of occipital nerve RFN is primarily for symptoms of ON causing posterior head pain and has also been described for use in migraine syndromes with occipital tenderness
- Implementation of the International Headache Society diagnostic criteria for ON is standard practice
- Diagnostic blockade of the occipital nerves should precede use of RFN
- Prior to RFN of the occipital nerves, other possible etiologies should be ruled out by way of either magnetic resonance imaging (MRI) and/or computerized tomography (CT) imaging. Imaging is not itself a diagnostic for ON
- Evidence best supports the use of PRFN over other RFN options. Time settings for lesion creation range

Table 9: Studies of Occipital Nerve Radiofrequency Neurotomy

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Study Type</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vanelderen et al 2010</td>
<td>Clinical Study</td>
<td>Prospective study in 29-month period, 19 patients underwent PRF. Mean visual analog scale and median Medication Quantification Scale scores declined by 3.6 (P = 0.002) and 8 (P = 0.006), respectively, at 6 months. 52.6% of patients reported a score of 6 (pain improved substantially) or higher on the Likert scale.</td>
</tr>
<tr>
<td>Huang et al 2012</td>
<td>Clinical Study</td>
<td>Retrospective study of 102 consecutive ON patients treated with PRF of the greater and/or lesser occipital nerve. 51% patients reported ≥50% pain relief and satisfaction with treatment for at least 3 months duration. Nerves treated were GON, LON or both GON and LON. Patients with lower volume diagnostic blocks and multiple lesions of PRF did significantly better.</td>
</tr>
<tr>
<td>Choi et al 2012</td>
<td>Clinical Study</td>
<td>10 patients treated with PRF with mean follow-up of 7.5 months show average 6.1 point reduction in VAS score.</td>
</tr>
<tr>
<td>Vanderhoek et al 2013</td>
<td>Case series</td>
<td>2 patients treated successfully with ultrasound guided blocks and PRF</td>
</tr>
<tr>
<td>Yu et al 2014</td>
<td>Case Report</td>
<td>35 yo female with benefit from WCRF of the bilateral GON, reporting 75% reduction in pain.</td>
</tr>
<tr>
<td>Cohen et al 2015</td>
<td>Multicenter randomized double-blind sham controlled RCT</td>
<td>Study compared effectiveness of occipital nerve block with steroid versus PRF. 81 participants with ON or migraine with occipital nerve tenderness. PRF was carried out with three cycles at 120 seconds each. The PRF arm showed significant improvements over control at 6 weeks with sustained benefits seen at 6 months.</td>
</tr>
<tr>
<td>Finiels et al 2016</td>
<td>Clinical Study</td>
<td>Retrospective 111 patients with positive diagnostic blocks, 78 of whom treated with RFN, remainder treated with botulinum toxin injection or occipital nerve stimulation. 89.4% of patients treated with RFN showed “good” and “very good” results out to 6 months, compared to 80% in the other treatment groups.</td>
</tr>
<tr>
<td>Hoffman et al 2018</td>
<td>Clinical Study</td>
<td>A retrospective study of 46 patients who underwent CRF at 80 degrees C for 180 seconds after sensory stimulation by landmark-based technique. A significant difference reduction in pain scores (6.7 vs 2.7, respectively; P &lt; 0.001), equating to a mean reduction in pain scores 1-month post-procedure of 4.0 ± 3.3. The mean patient-defined percent pain relief was 76.3% ± 25.0%. The mean patient-reported length of relief was 6.5 ± 5.1 months.</td>
</tr>
</tbody>
</table>
from a single lesion at 90 seconds to up to three lesions for 120 seconds. Further studies are necessary to compare procedural techniques.

- Six months of pain relief has been documented in several studies. Longer follow-up periods are needed to put forth recommendations for repeat sessions of RFN in successfully treated patients.

Treatment Gaps
Further research is needed to best understand the optimal protocol for diagnostic block, the use of image guidance for occipital RFN, the optimal procedure techniques, and timetable for repeat procedures in successfully treated patients.

Future Research
Research and evidence gaps exist for occipital nerve pulsed radiofrequency (PRF). PRF allows for the application of radiofrequency current at lower temperatures, thereby minimizing the damage to surrounding tissue, nerve or vasculature. They may be the preferred technique given the superficial location of the occipital nerves. Further research on PRF is needed to determine the efficacy and safety of this treatment modality, and how it compares to CRF and WCRF. Although several studies have demonstrated the efficacy of both continuous and pulsed RF within the range of 6 weeks to 6 months more evidence is clearly needed.

Consensus Statement
Occipital neurotomy may be selectively offered for the treatment of occipital neuralgia pain when greater or lesser nerves have been identified as the etiology of pain via diagnostic blocks. GRADE II-2 C.

Conclusion
The use of radiofrequency ablation to treat pain is an established therapy that continues to evolve. This best practice document gives an evaluation as to the current evidence and recommendations. Going forward, these recommendations will be updated as new data is produced by either high-level studies or from large registries. Future guidelines will be modified as evidence is built, innovations arrive at the technology, and new ideas are presented to continue to improve patient safety and efficacy.

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Dr Jonathan M Hagedorn reports personal fees from Abbott, personal fees from Boston Scientific, personal fees from Nevro, personal fees from Speranza Therapeutics, outside the submitted work. Dr Jordan Tate reports personal fees from Abbott. Dr Dawood Sayed reports personal fees from Abbott, personal fees from Medtronic, personal fees from Nevro, PainTeq, Spr, and Vertos, during the conduct of the study. Dr Timothy R Deer reports personal fees from Abbott; also his institution research received funding from Avanos, outside the submitted work. The authors report no other conflicts of interest in this work.

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47. Lee et al


Lee et al


