Update on neuropathic pain treatment for trigeminal neuralgia

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ABSTRACT

Trigeminal neuralgia (TN) is one of the most common causes of facial pain seen in dental and neurologic practices. This classic neuropathic pain disorder has been known for centuries. The International Headache Society (IHS) defines TN as a “unilateral disorder characterized by brief electric shock-like pains, abrupt in onset and termination, and limited to the distribution of one or more divisions of the trigeminal nerve.” The age of onset for most idiopathic cases is usually between 40 and 60 years, although onset may occur in the second and third decades, most often in females. A variety of medical and surgical treatments are available for TN. Antiepileptic drugs (AEDs) are considered first line therapy for TN with different efficacy, and carbamazepine is the drug of choice. Surgical interventions are reserved for patients...

Microvascular decompression, gamma knife radiosurgery, and percutaneous rhizotomies are most promising surgical alternatives. This paper reviews the medical and surgical therapeutic options for the treatment of trigeminal neuralgia, based on available evidence and guidelines.

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Trigeminal neuralgia is a syndrome of unilateral, paroxysmal, stabbing facial pain, originating from the trigeminal nerve. Careful history of typical symptoms is crucial for diagnosis. Most cases are caused by vascular compression of the trigeminal root adjacent to the pons leading to focal demyelination and ephaptic axonal transmission. Brain imaging is required to exclude secondary causes. Many medical and surgical treatments are available. Most patients respond well to pharmacotherapy; carbamazepine and oxcarbazepine are first line therapy, while lamotrigine and baclofen are considered second line treatments. Other drugs such as topiramate, levetiracetam, gabapentin, pregabalin, and botulinum toxin-A are alternative treatments. Surgical options are available if medications are no longer effective or tolerated.
who do not respond to adequate medical therapy. Microvascular decompression (MVD), percutaneous trigeminal rhizotomies, and gamma knife radiosurgery (GKRS) are possibly effective in the treatment of TN.11

**Etiology and pathogenesis.** The cause of TN remains unclear.12,13 However, most cases are caused by compression of the trigeminal nerve root within a few millimeters of entry into the pons.9,14 The nerve impingement is often accompanied by a demyelination of sensory fibres within the nerve root or the root entry zone, or less commonly in the brainstem.15 Vascular compression by an aberrant loop of an artery or vein accounts for 80-90% of idiopathic TN.14 Other compressive causes are benign tumors of the posterior fossa such as acoustic neuroma, meningioma, and epidermoid cyst.9,16

**Diagnosis.** The diagnosis of TN is usually based on the characteristic clinical picture. The key feature is a sudden and severe lancinating pain, usually unilateral, precipitated by touching facial zones. This pain occurs in paroxysms, within the trigeminal nerve distribution; typically involving the maxillary nerve (V2) or mandibular nerve (V3) distribution and lasts for a fraction of a second to 2 minutes.9

**Classification.** The IHS divides TN into classic and symptomatic categories based on presumed etiology. Trigeminal neuralgia is termed classic (or idiopathic) when investigation identifies no cause other than a neurovascular compression; and examination shows no clinical evidence of neurological deficit. This accounts for 80-90% of TN cases.14 The classification symptomatic (or secondary) is reserved for patients with TN when major neurological diseases such as MS, skull deformity, or benign compressions in the posterior fossa have been identified.17-19 Neurological examination may show sensory impairment in trigeminal nerve distribution.2

**Therapeutic modalities.** A huge variety of pharmacological and surgical treatments are available for TN. The practice parameters and guidelines published in 2008 from the American Academy of Neurology (AAN), and the European Federation of Neurological Societies (EFNS)11,20 recommend starting treatment with drugs in patients with classic TN (Table 1). Surgical procedures should be reserved for patients who are refractory to medical therapy or when drugs are causing unacceptable adverse effects. There are few studies directly comparing medical and surgical treatments. However, pharmacological therapy for TN has been the subject of several Cochrane systematic reviews,8,21,22 and available evidence shows carbamazepine is the best-studied treatment and drug of choice for initial and long-term management of classical TN.8,9,23 As far as symptomatic TN is concerned, to date, no placebo-controlled trials have evaluated and established effective treatment. However, it is advisable to treat symptomatic TN with the same drugs used in classical TN.

**Pharmacological therapy.** Different medication has been considered for treatment of TN. Based on the level of evidence; carbamazepine and oxcarbazepine should be offered as a first line for pain control.11 There is limited evidence for efficacy of different AEDS to treat TN; other medications such as Baclofen and Botulinum toxin A seem promising treatment for this disorder.11

**First-line therapy.** According to current evidence-based treatment guidelines published in 2008 from the AAN and EFNS,20 carbamazepine is established as effective (level A) and oxcarbazepine is probably effective (level B) for controlling pain in classic TN. These guidelines recommend carbamazepine (200-1200 mg/d) and oxcarbazepine (600-1800 mg/d) as a first-line therapy for classic TN.

**Carbamazepine.** Carbamazepine acts by inhibiting voltage-gated sodium channels, thereby reducing the excitability of neural membranes. Carbamazepine has also been shown to potentiate gamma aminobutyric acid (GABA) receptors made up of alpha1, beta2, and gamma2 subunits. This may be relevant to its efficacy in neuropathic pain.24 In newly diagnosed cases of TN, the usual starting dose is 100 to 200 mg twice daily. The daily dose should be increased by 100 mg every other day until sufficient pain relief is attained or until intolerable side effects prevent further upward titration. The typical total maintenance dose is 300-800 mg/d, given in 2-3 divided doses. The maximum suggested total dose is 1200 mg/d. With appropriate dose adjustments, pain can be controlled in around 75% of patients.5,25 The dose may be tapered once pain is controlled, since remission may occur. Extended release carbamazepine is useful as a night dose in patients with pain attacks during sleep, as drug levels do not fall. This not only keeps patients pain free during sleep, but may reduce side effects, as high serum peaks are not achieved.26,27 Common side effects include sedation, dizziness, nausea, vomiting, diplopia, memory problems, ataxia, elevation of hepatic enzymes, and hyponatremia, which may contraindicate it for elderly patients. Potentially serious but uncommon side effects are carbamazepine-induced leucopenia, aplastic anemia, allergic rash, systemic lupus erythematosus, hepatotoxicity, and Stevens-Johnson syndrome (SJS). It is advisable to order complete blood count, serum sodium, and liver function tests within several weeks after starting therapy to detect any complications quickly.8,15

**Oxcarbazepine.** Oxcarbazepine is a keto-analogue of carbamazepine that is rapidly converted into its
The keto derivative of carbamazepine does not pass through the liver cytochrome system, resulting in an improved side effect profile and fewer drug interactions than with carbamazepine.28-30 Oxcarbazepine is an acceptable alternative to carbamazepine, which may have provided pain relief but has caused unacceptable adverse effects. Better tolerability can also be considered an advantage over carbamazepine.31 A systematic review from the American Academy of Neurology / European Federation of Neurological Societies (AAN/EFNS)11 identified several randomized controlled trials that compare oxcarbazepine with carbamazepine in patients with classic TN. Oxcarbazepine can be started at 150 mg twice daily. The dose can be increased as tolerated in 300 mg increments every third day until pain relief occurs. Maintenance doses range between 300-600 mg twice daily.26 The maximum suggested total dose is 1800 mg/d. The risk of allergic cross reactivity between carbamazepine and oxcarbazepine is around 25%, so oxcarbazepine is best avoided when carbamazepine allergy is evident.9

**Second-line therapy.** Second-line treatment is based on very little evidence. Three drugs are included in this class - lamotrigine, baclofen, and pimozide. Each drug has been studied in single trials,32-34 and per AAN/EFNS guidelines, are possibly effective (level C) for controlling pain in patients with classic TN.20

**Lamotrigine.** This AED has been used in only one small, double-blind, placebo-controlled trial14 investigating 14 patients with TN who were refractory to carbamazepine, where adjunct therapy with lamotrigine (400 mg/d) was beneficial and more effective than placebo. While in an open-label study, 11 of 15 patients with TN found pain relief with 400 mg of lamotrigine.35 The initial dose of lamotrigine is 25 mg twice daily, and can be increased gradually to a maintenance dose of 200-400 mg/d in 2 divided doses.15 The dosage required for adequate pain relief varied widely from 100-400 mg/d.35 Common side effects are sleepiness, dizziness, headache, vertigo, and ataxia. Some (7-10%) of patients report a skin rash during the first 1-2 months of treatment that most often resolves with continued therapy.36 Stevens-Johnson syndrome can occur in one in 10,000 patients taking lamotrigine.15 Such adverse reactions can be prevented by slower titration of the drug.

**Baclofen.** Baclofen, a skeletal muscle relaxant, is a GABA analogue that activates GABAB receptors and thus depresses excitatory neurotransmission.26 It is effective in controlling pain in TN at a dose of 60-80 mg/d. Baclofen can be used alone or in combination with carbamazepine. Studies have supported its benefit in both conditions.37 The initial dose is 10 mg/d for 3 days, which can be increased to 10-20 mg/d every 3 days if needed. The maximum tolerated dose is 60-80 mg/d, administered 3-4 times per day.37 If baclofen is an add-on therapy with carbamazepine, it is advisable to reduce the dose of carbamazepine to 500 mg/d to maintain a putative synergistic effect.38 Typical side effects of baclofen include drowsiness, dizziness, weakness, fatigue, nausea, hypotension, and constipation. Abrupt discontinuation of baclofen can cause withdrawal symptoms (hallucinations and seizures).15 Patients with MS and trigeminal neuralgia obtain additional benefits with baclofen because it is a muscle relaxant.39 To date, baclofen has the strongest evidence for efficacy in the treatment of TN after carbamazepine.15

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**Table 1 - Clinical approach to the management of trigeminal neuralgia (TN).**

| Establish diagnosis of classic TN |
|----------------------------------|
| Paroxysmal lancinating facial pain in the distribution of the trigeminal nerve with normal neurological examination |
| Evaluate for symptomatic TN |
| MRI brain with MRA to explore underlying demyelinating lesion, posterior fossa tumor, or vascular malformation |

**Medical therapy**

| First line: Carbamazepine or oxcarbazepine |
| Second line: Baclofen or lamotrigine |
| Third line: Levetiracetam, gabapentin, pregabalin, topiramate, Botox-A |

**Surgical options**

If there is failure of at least 3 drug trials, or drugs are causing unacceptable side effects, and according to the candidate’s medical status, age, or preference

- Microvascular decompression: invasive, for healthy patients, highest success rate, risk of mortality
- Percutaneous rhizotomy: less invasive, suitable for elderly patients, risk of recurrence
- Gamma knife radiosurgery: non invasive, suitable for most of patients, expensive

MRI - magnetic resonance imaging, MRA - magnetic resonance angiography

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Pimozide. Pimozide, a dopamine receptor antagonist, is used mainly in the management of Tourette syndrome. It was found effective in a randomized, double-blind crossover trial of 48 patients with refractory TN. In this trial, pimozide achieved pain control in all 48 patients. While displaying encouraging results, pimozide, at a dose of 2-12 mg/d in TN treatment, is seldom used clinically because it has multiple potential serious side effects, including arrhythmias, acute extrapyramidal symptoms, and Parkinsonism.

Third-line therapy. The newer AEDs tested within the past few years are gabapentin, pregabalin, topiramate, and levetiracetam. Gabapentin, a GABA receptor agonist, acts primarily on presynaptic calcium channels of neurons to inhibit the release of excitatory neurotransmitters. Gabapentin has been used in randomized control trials (RCTs) of neuropathic pain and was proven effective. Its use and effectiveness were also reported in several TN studies. Gabapentin showed adequate efficacy in only one RCT, where it was used in combination with ropivacaine. This combination was found to be safe and effective. Gabapentin also has demonstrated effectiveness with sustained relief in TN, especially in patients with MS. Treatment can be started at a dose of 300 mg/d, and may be gradually increased by 300 mg every 2-3 days as tolerated. For maximum efficacy, the dose can be increased to 1800 mg/d. Gabapentin has many advantages, including faster titration, no known drug interactions, no known idiosyncratic skin reactions, and a favorable side-effect profile, with mild somnolence, dizziness, headache, confusion, nausea, and ankle edema. Hyperlipidemia is an important side effect to watch for with gabapentin therapy.

Pregabalin. Pregabalin is an analog of GABA, structurally related to gabapentin. It acts by interacting with the alpha-2-delta (α2-δ) subunit of voltage-gated calcium channels. Although a potentially useful drug for neuropathic pain in some patients, evidence is scant in TN. It has been tested and found effective only in a single open-label study. In this well-designed cohort study, pregabalin (150-600 mg/d) was tested in 53 patients over one year, and proved to be effective in reducing TN pain by more than 50-74% of patients. Side effects are similar to other AEDs but less marked; most common are dizziness and sleepiness.

Topiramate. The exact mechanism of action of topiramate is unknown. However, its pain-modulating effect might be related to its property of blockage of the voltage-gated sodium channel and an augmentation of GABA activity by binding to a non benzodiazepine site on the GABAA receptor. Topiramate (100-400 mg/d) was found effective in 75% of patients in one study of 8 patients with classic TN. A recent meta-analysis evaluated the effectiveness and safety of topiramate with carbamazepine in the treatment of classic TN. In this analysis, 6 RCTs involving 354 patients were included. A meta-analysis of these studies showed the overall effectiveness and tolerability of topiramate did not seem to differ from carbamazepine in the treatment of classic TN. In this analysis the results were limited due to the poor methodological quality of these RCTs. The most frequently registered side effects of topiramate were dizziness, somnolence, cognitive impairment, and weight loss.

Levetiracetam. Levetiracetam is a newer AED that has been tried in TN. The exact mechanism by which it acts is unknown, but it is thought to target high-voltage, N-type calcium channels as well as the synaptic vesicle protein 2A (SV2A); by this, it impedes impulse conduction across synapses. Its evidence in TN is scant. Recently, 2 pilot, open-label studies investigated the efficacy and tolerability of levetiracetam in patients with TN. One study with 10 patients reported an improvement of 50-90%. Another study with 23 patients reported a 62% reduction in number of daily attacks in patients receiving levetiracetam as add-on therapy. The effective dose range of levetiracetam in TN is 1000-4000 mg/d. Levetiracetam has advantages, including no need for routine blood tests, less drugs interactions, the absence of auto-induction effect, nasopharyngitis, sleepiness, headaches, and irritability are side effects when starting levetiracetam. Although evidence suggests good efficacy and safety for levetiracetam in medical treatment of TN, a wide-range of randomized and placebo-controlled trials are warranted before making any definitive claim.

Botulinum toxin A. Botulinum toxin A (BTX-A) has been thoroughly studied as a potential tool in the treatment of several pain syndromes, such as migraine, tension headache, ocipital, and postherpetic neuralgias. The BTX-A’s mechanism of analgesic effect is still unclear, but it is postulated it causes local release of anti-nociceptive neuropeptides such as substance P, glutamate, and calcitonin-gene related peptide, inhibiting central and possibly peripheral sensitization. Its therapeutic efficacy and safety has been reviewed recently. In this systemic review, they identified one double-blind RCT study and 5 prospective open-labelled studies. This systemic review found subcutaneous or mucosal injection of BTX-A effective for adult TN patients, and the results of the 2 RCTs showed significant benefit over placebo. Response was achieved in approximately 70-100%
of patients, and mean pain intensity and frequency were reduced by approximately 60-100% at 4 weeks after injection in most studies. In one study, 47% of patients didn’t need further treatment, 33% patients required nonsteroidal anti-inflammatory drugs to alleviate pain, while 20% responded to the AED after receiving a BTX-A injection. The most commonly used dose of BTX-A was 20-75U, however, patients also showed significant reduction in intensity of pain at lower doses (6-9U). Based on this available limited evidence, BTX-A seems promising treatment of TN. However, well-designed RCT studies are required to determine the therapeutic efficacy, optimal dose, time, and indications for repeat injections of BTX-A.

Other drugs. A number of other drugs have been tried in treating TN, showing limited benefit such as phenytoin and intravenous phenytoin, fosphenytoin, clonazepam, valproic acid, misoprostol, tocainide, topical capsaicin cream, intranasal lidocaine, tizanidine, sumatriptan, and amitriptyline.

Surgical treatment. Surgical interventions are reserved for the candidate with incapacitating symptoms of TN despite a trial of at least 3 drugs in sufficient dosage or where medications caused unacceptable adverse effects. Relapse of symptoms might also be another strong factor in opting for surgical treatment. It is estimated that up to 50% of patients with TN will require some form of surgery sooner or later. According to practice parameters from the AAN and EFNS, microvascular decompression (MVD), percutaneous rhizotomy on the Gasserian ganglion, and gamma knife radiosurgery (GKRS) may possibly be effective in the treatment of TN, while evidence for peripheral neurectomy is inconclusive.

Microvascular decompression. Among all surgical procedures, MVD is the most invasive surgery, but offers the highest success rate of pain-free status in patients with classic TN. Generally it is reserved for the patient with TN who is otherwise, healthy and has no other major medical problems, or for patients who may have failed less invasive procedures such as radiosurgery and rhizotomy. It is carried out under general anesthesia (GA), involves retrosigmoid craniotomy and microsurgical exploration of the posterior fossa. A systemic review by AAN/EFNS identified 5 studies on MVD for TN. The studies concluded 90% of patients experience initial pain relief; more than 80% are still pain free after one year, with 75% pain free after 3 years, and 73% after 5 years. In this review, the most common complication was aseptic meningitis (11%), followed by long-term hearing loss (10%), and sensory loss (7%). Major adverse events were noted in 4% of patients in the form of cerebrospinal fluid leaks, infarction, or hematoma. Although complication rates were low, this surgical option has a small risk of death (around 0.20-0.4%).

Percutaneous rhizotomy on the Gasserian ganglion. There are 3 common percutaneous rhizotomies: radiofrequency thermal rhizotomy (that creates a lesion through application of heat), chemical rhizotomy (by injecting 0.1 to 0.4 mL of glycerol into the trigeminal cistern), and mechanical rhizotomy or micro compression (by inflating a balloon into Meckel’s cave to compress the Gasserian ganglion). These techniques are performed by passing a cannula through the foramen ovale, followed by controlled lesion of the trigeminal ganglion or root. The goal of these rhizotomies is selective destruction of pain fibers (A-delta and C-fibers) while preserving touch fibers (A-alpha, and beta fibers) in the trigeminal nerve.

The systemic review by the AAN/EFNS identified 4 uncontrolled case studies of these percutaneous procedures, and concluded initial pain relief is achieved in 90% of patients, but pain free rates decline gradually in successive years, and after 5 years only 50% of patients are pain free. The most common complication reported are trigeminal distribution sensory loss (50%), followed by dysesthesias (6%), anesthesia dolorosa (4%), characterized by persistent, painful anesthesia, or hypesthesia in the denervated region, corneal numbness with risk of keratitis (4%), aseptic meningitis (0.2%), and extremely low mortality. Percutaneous procedures are less invasive and have relative advantages and disadvantages. They are used to a greater extent in patients with high operative risks such as in elderly or ill patients or as a partially diagnostic procedure in atypical disease. All these procedures are associated with significant risk of recurrence.

Gamma knife radiosurgery. The GKRS has recently been established as an effective treatment for TN. It is a noninvasive outpatient procedure in which a high dose (70-90 Gy) of a highly focused beam of radiation is delivered at the trigeminal root entry zone, which over time causes axonal degeneration and necrosis and thus interrupts pain signals. The success of GKRS demands a very accurate stereotactic system. The AAN/EFNS systemic review identified 3 case series of GKRS that used independent outcome assessment, revealing 69% of patients are pain free at one year after radiosurgery, and 52% are still pain free at 3 years. Pain relief can be delayed (mean one month) with this procedure. Side effects are facial numbness (9-37%), which improves over time, and bothersome sensory loss or paresthesia (6-13%) that may develop.
with a delay of up to 6 months. Quality of life improves by 88%.77 The non-invasive nature of GKRS provides many advantages, and makes it available for patients who cannot opt for MVD because of multiple medical comorbidities, or are on anticoagulation therapy, or are unwilling to take anticonvulsants.67 The treatment is expensive, which limits its widespread use.

In conclusion, TN is one of the most painful, but readily treatable conditions, whose diagnosis is usually based purely on clinical grounds by characteristic symptoms and signs. Many medical and surgical treatments are available; however, only a few of them have proven their efficacy to modern evidence-based medicine standards. The AEDs form the mainstay of treatment, among them carbamazepine is the gold standard. Newer drugs such as gabapentin, pregabalin, topiramate, levetiracetam, and botulinum toxin-A are promising and effective with fewer side effects, better efficacy, and tolerability. However, there is still no existing evidence to use them as first-line therapy. Medically refractory cases and patients who cannot tolerate the side effects of medication may benefit from surgical management. Microvascular decompression remains the best approach in patients when MRI shows a loop of the aberrant vessel. Other options are percutaneous ablative procedure and GKRS. The selection of the surgical procedure needs to be individualized, depending upon patient’s preference, age, underlying pathology, and comorbidities. There is a need for wide-ranged randomized and placebo-controlled trials with long-term-follow up to establish the stepwise, standardized medical regimen after failure of first-line medical therapy, to compare surgical and medical therapies, and to determine the optimal timing for surgical intervention.

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