Surgical Therapy of Temporal Triggered Migraine Headache

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Background: The auriculotemporal and zygomaticotemporal nerves are the 2 primary trigger points in the temporal area of migraine headache. Different surgical approaches are described in literature, either open or endoscopic ones.

Methods: We described and delineated the currently adopted strategies to treat temporal trigger points in migraine headache. Furthermore, we reported our personal experience in the field.

Results: Regardless of the type of approach, outcomes observed were similar and ranged from 89% to 67% elimination / >50% reduction rates. All procedures are minimally invasive and only minor complications are reported, with an incidence ranging from 1% to 5%.

Conclusions: Just like upper limb compressive neuropathies, migraine headache is believed to be caused by chronic compression of peripheral nerves (ie, the terminal branches of trigeminal nerve) caused by surrounding structures (eg, muscles, vessels, and fascial bands) the removal of which eventually results in improvement or elimination of migraine attacks. Particular attention should be paid to the close nerve/artery relationship often described in anatomical studies and clinical reports. (Plast Reconstr Surg Glob Open 2018;6:e1980; doi: 10.1097/GOX.0000000000001980; Published online 17 December 2018.)

INTRODUCTION

Following the studies of Guyuron1–10 and those of other independent groups, migraine surgery has been widely accepted as an effective surgical solution for chronic headaches refractory to medical and conservative treatment. Although the pathophysiology of migraine headaches (MH) remains a matter of debate, it’s a common belief that chronic compression to the terminal branches of trigeminal nerve caused by surrounding structures (eg, muscles, vessels, and fascial bands) is responsible for its origin. Four main trigger zones of MH are amenable to surgical decompression: frontal (site I: supraorbital and supratrochlear nerves), temporal (site II: zygomatic-temporal branch of the trigeminal nerve), endonasal (site III: trigeminal end branches), and occipital (site IV: great occipital nerve).11 The auriculotemporal nerve (ATN) and lesser occipital nerve, site V and VI, respectively, are commonly described as less relevant trigger sites12 (Table 1).

This article aims to describe and carefully delineate the currently adopted strategies to treat temporal trigger points in MH sufferers. The ATN and zygomaticotemporal nerves (ZTN) are the 2 primary trigger points in the temporal area. Different surgical approaches are described in literature, either open or endoscopic ones. Regardless of the surgical approach chosen, the procedure aims at decompressing the putative irritated nerves from the surrounding offending structures. Therefore, a clear anatomical knowledge of this area should be kept in mind.

RELEVANT ANATOMY

Many anatomical studies have been carried out to describe the anatomy of the temple, particularly the course of the ATN and ZTN and the structure that could potentially compress them. The ZTN is one of the terminal branches of the maxillary division (V2) of the trigeminal nerve. It originates from the bifurcation of the zygomatic branch within a canal in the zygomatic bone, from which it exits in the anterior part of the temporal fossa 17 mm posterolateral and 6.5 mm cephalad from the lateral canthus.13 Then, it as-

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It provides innervation to: temporomandibular joint, parotid gland, external acoustic meatus, anterior auricle, zygoma, and superficial temporal region. It originates from mandibular division of the trigeminal nerve within the pterygopalatine fossa, it turns around the inferior border of the zygomatic arch in close proximity to the temporomandibular joint, and then it reaches the facial soft tissue running cranially within the temporoparietal fascia superficial to the superficial temporal artery (STA). The ATN has a single main trunk in 78.5% of the cases while in the remaining 21.5% it’s duplicated at the retromandibular space.\(^\text{11}\)

Whenever the ATN is duplicated, both trunks accede to the superficial layers at approximately 10 mm above the middle of the tragus.\(^\text{18}\) The anterior trunk runs cranially parallel to the STA, while the posterior can be found into the subcutaneous layer 1 mm anterior to the auricle. Communicating branches can be present at this level.

The ATN has been traditionally described to divide into several superficial temporal branches (STb) innervating the skin of the temple. Gülekon et al.\(^\text{19}\) demonstrated that the nerve has one branch in 50% of specimens, while multiple branches (up to 4) can be present in the other 50% of specimens. Baumel et al.\(^\text{20}\) identified 5–9 terminal STBs. Iwanaga et al.\(^\text{17}\) reported from 2 to 7 STB, which branched from the main trunk at a distance of 6.19–25.66 mm vertically and 3.45–11.88 mm horizontally from the middle of the tragus. Furthermore, from 0 to 4 communicating branches were reported, which occasionally formed a loop or “ansa.” Andersen et al.\(^\text{21}\) reported that the ATN might have a single main branch, a diffuse branching pattern, or divide into smaller branches, herein described as STb (Table 2).

Three potential compression points of the ATN along its course have been described: 2 preauricular fascial bands (the most caudal one is found 13.1±5.9 mm anterior and 5.0±7.0 mm superior to the most anterosuperior point of the external auditory meatus, whereas the most cranial one is located 11.9±6.0 mm anterior and 17.2±10.4 mm superior to the same landmark) and the STA.\(^\text{14}\) Indeed, a

### Table 1. Migraine Headache Trigger Sites, Peripheral Nerves Involved, and Putative Anatomical Compressing Structures

<table>
<thead>
<tr>
<th>Migraine Headache Trigger Sites</th>
<th>Involved Trigeminal Nerves</th>
<th>Potential Compressing Points</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site I – frontal</td>
<td>Supraorbital and supratrochlear nerve ((V_1))</td>
<td>Glabellar muscles</td>
</tr>
<tr>
<td></td>
<td></td>
<td>STA</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Temporalis muscle and fascia</td>
</tr>
<tr>
<td>Site II – temporal</td>
<td>Zygomatic-temporal nerve ((V_2))</td>
<td>STA</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sentinal vein</td>
</tr>
<tr>
<td>Site III – endonasal</td>
<td>Trigeminal end branches ((V_3))</td>
<td>Deviated septum</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Turbinates – septum contact</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Concha bullosa</td>
</tr>
<tr>
<td>Site IV – occipital</td>
<td>Greater occipital nerve ((C_2))</td>
<td>Occipitalis, trapezius, and semispinalis capitis</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fascial bands</td>
</tr>
<tr>
<td>Site V – auriculotemporal</td>
<td>Auriculotemporal nerve ((V_2))</td>
<td>Superior temporal artery</td>
</tr>
<tr>
<td>Site VI – lesser occipital</td>
<td>Lesser occipital nerve ((C_2))</td>
<td>Preauricular fascial bands</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trapezius and sternocleidomastoid muscles</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fascial bands</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Occipital artery</td>
</tr>
</tbody>
</table>

### Table 2. Zygomatic-temporal and Auriculotemporal Nerves’ Relevant Anatomy

<table>
<thead>
<tr>
<th>Temporal Trigger Site</th>
<th>Involved Trigeminal Nerve</th>
<th>Anatomy of Relevant Temporal Nerves</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site II – temporal</td>
<td>Zygomatic-temporal nerve ((V_2))</td>
<td>- Origin: bifurcation of the zygomatic branch within a canal in the zygomatic bone</td>
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<tr>
<td></td>
<td></td>
<td>- Exit point: anterior temporal fossa 17 mm posterolateral and 6.5 mm cephalad from the lateral canthus</td>
</tr>
<tr>
<td>Site V – auriculotemporal</td>
<td>Auriculotemporal nerve ((V_2))</td>
<td>- Origin: mandibular division of trigeminal nerve within pterygo-palatine fossa</td>
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<td></td>
<td></td>
<td>- Exit point: turns around the inferior border of the zygomatic arch in close proximity to the temporomandibular joint</td>
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close relationship between these 2 structures is reported with different percentage in literature.

Chim et al.\(^{14}\) found as high as 80% close relationship of the ATN with the STA with 3 patterns of interaction: single STA crossing over of the ATN (62.5%), helical intertwining (18.8%), and single ATN crossing over the STA (18.8%). Conversely, Janis et al.\(^{15}\) found a 34% contiguous relationship between ATN and STA. Furthermore, only 2 patterns of interaction were reported: single, short, discrete intersection (88.2%), and helical intertwining (11.8%; Fig. 1). Anatomic studies describe the course of nerves and their relationships with arteries in a “normal” population, with no recorded history of migraine. Hence, patients with migraine could experience headache because their anatomic relationships are not normal (Table 3).

**TRIGGER POINT DETECTION**

Detecting the precise site of pain onset (trigger point) is of paramount importance for a successful outcome (Table 4). Indeed, partial or nonresponse to surgery may be primarily related to failure in recognizing all the trigger points. Botulinum toxin injections, nerve blocks, portable Doppler devices were all investigated.\(^{12}\) Botulinum toxin administration proved to be useless since lack or incomplete response does not automatically exclude the suspected trigger point, since close nerve-artery relationship might be the main cause of MH.\(^{12}\) Nerve blocks could be helpful; however, patients should be seen in the office during a MH attack. Finally, handheld Doppler devices proved to be useful in detecting migraine trigger points, especially because the nerve-artery crossover or close relationship might be the implicated in the headache pathogenesis.\(^{12}\)

Nevertheless, clinical findings are still the best way to identify trigger points’ location.

Patients suffering from site II temporal migraine usually report diffuse low temporal pain closer to the lateral canthus, whereas those with site V involvement tend to complain of more superior pain localized in the temple or alternatively moderate-to-severe pain on the preauricular area, often spreading to the ipsilateral temple.\(^{22}\)

Patients usually report diffuse headache; however, they can precisely pinpoint with one fingertip the precise location of the tender spot form where the MH attack begins, and that is where the surgical treatment should be carried out. Furthermore, simple compression of the tender spot by the fingertip of the surgeon during physical examination can usually evoke pain confirming the diagnosis. Therefore, it is the authors’ experience that accurate history and careful physical examination can safely detect all patient’s trigger points, while positive handheld Doppler signal can corroborate the diagnosis.

**Table 3. Auriculotemporal Nerve Potential Compression Points Anatomy**

| Auriculotemporal Nerve Potential Compression Points | Superior preauricular facial band | Inferior preauricular facial band | STA  
Chim et al.\(^ {14}\) | Janis et al.\(^ {15}\) |
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<tbody>
<tr>
<td>13.1 ± 5.9 mm anterior and 5.0 ± 7.0 mm superior to the most anterosuperior point of the external auditory meatus.</td>
<td>11.9 ± 6.0 mm anterior and 17.2 ± 10.4 mm superior to the most anterosuperior point of the external auditory meatus.</td>
<td>- Reported 80% close relationship of the ATN with the STA with three patterns of interaction: single STA crossing over of the ATN (62.5%), helical intertwining (18.8%), and single ATN crossing over the STA (18.8%).</td>
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<tr>
<td>- Single intersection site was found at a mean distance of 19.2 ± 10.0 mm anterior and 39.5 ± 16.6 mm superior to the most anterosuperior point of the external auditory meatus.</td>
<td>- Helical intertwining relationship had a mean length of 10.3 ± 0.4 mm; inferior limit was at a mean distance of 20.0 ± 15.6 mm anterior and 53.7 ± 4.7 mm superior to the most anterosuperior point of the external auditory meatus, superior limit was at a mean distance of 24.7 ± 17.9 mm anterior and 62.7 ± 3.8 mm superior to the same anatomical landmarks.</td>
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<tr>
<td>- Helical intertwining relationship had a mean length of 21 mm, starting inferiorly 123 mm lateral to the midline and 37.53 ± 15.29 mm cranial to the nasion-lateral orbit line.</td>
<td>- Reported a 34% contiguous relationship between ATN and STA with only 2 patterns of interaction: single, short, discrete intersection (88.2%), and helical intertwining (11.8%).</td>
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<tr>
<td>- Helical intertwining relationship had a mean length of 21 mm, starting inferiorly 123 mm lateral to the midline and 25 mm cranial to the nasion-lateral orbit line, ending superiorly 117 mm lateral to the midline and 38 mm cranial to the nasion-lateral orbit line.</td>
<td>- Single intersection site was found on average 107.88 ± 17.73 mm lateral to the midline and 37.53 ± 15.29 mm cranial to the nasion-lateral orbit line.</td>
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SURGICAL PROCEDURE

Endoscopic or open surgical approaches are the only viable techniques to alleviate migraine headaches.

Site II—Zygomaticotemporal Nerve Deactivation

Guyuron1,3 has developed the technique for endoscopic decompression of ZTN. The procedure is performed under local anesthesia; two 1.5-cm incisions are made on each side at approximately 7 and 10 cm from the midline in the hair-bearing area of the scalp. Incisions are deepened to the deep temporalis fascia by means of Metzenbaum scissors to expose the deep temporal fascia and then a periosteal elevator is used for medial, lateral, cephalad, and caudal dissection to accommodate the endoscope. Under endoscopic view, the ZTN is exposed, isolated, and either avulsed by removing approximately 2.5 cm of the nerve or decompressed by widening the fascia opening and cauterizing the zygomaticotemporal artery.23

Gfrerer et al.24 and Peled25 have described open approaches for decompression surgery of the ZTN through transpalpebral and temporal hair-bearing area incisions, respectively. Again, careful dissection is performed along the deep temporal fascia to prevent lesions to the temporal branch of the facial nerve. Once exposed, the ZTN is either avulsed or decompressed under direct vision.

Site V—Auriculotemporal Nerve Deactivation

ATN deactivation surgery is usually performed with an open approach only, since the nerve lies within the superficial temporal fascia just beneath the subcutaneous tissue, where the STA can be found too. If site V surgery has to be done in conjunction with site II, 5–7 port incision is regularly designed for endoscopic decompression of ZTN and a further 1.5-cm lateral incision is made locating the tender point with the aid of handheld Doppler. Dissection is made with blunt tipped scissors to expose the ATN and the STA taking care not to injure the temporal branch of the facial nerve (even though it usually lies in a deeper plane). Then once the ATN and STA are isolated, the last one can be either cauterized or simply ligated.26

If the trigger point lies beyond the hairline or in proximity of the temporal branch of facial nerve, ATN and STA exposure, isolation, and ligation of the artery can be performed with a 1.5-cm incision 0.5 cm anterior to the tragus, just above the temporo-mandibular joint. At this level, the ATN is identified first, while the STA is located just deeper to it.

Authors’ Experience

We perform both procedures under local assisted anesthesia.11 Addressing ZTN, we make 3-cm cutaneous incision 8–10 mm behind the temporal hairline and take dissection deep to the deep temporal fascia by the blunt tip scissors. We open the inferior temporal septum exposing the inferior temporal compartment that contains the ZTN, sentinel vessels, and temporal branches of facial nerve (that must be carefully preserved by incorporating it in the roof of the elevated flap). Then we widen the exit

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Table 4. Trigger Point Detection Techniques’ Advantages and Drawbacks

<table>
<thead>
<tr>
<th>Trigger Point Detection Technique</th>
<th>Advantages</th>
<th>Drawbacks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical findings</td>
<td>Still the gold standard detection technique for trigger points’ location, patients can easily recognize the starting point of their MH attacks.</td>
<td>It’s a subjective approach and relies in patients’ ability to report their trigger point locations.</td>
</tr>
<tr>
<td>Botulinum injections</td>
<td>It can be applied even when patients aren’t experiencing MH attacks.</td>
<td>It’s costly, it requires second office evaluation, thus it can’t be easily employed for patients that live far or for off the country ones.</td>
</tr>
<tr>
<td>Local anesthetics injections</td>
<td>It is helpful since it doesn’t require second office-based visit.</td>
<td>Lack or incomplete response does not automatically exclude the suspected trigger point, since close nerve-artery relationship might be the main cause of MH.</td>
</tr>
<tr>
<td>Handheld Doppler</td>
<td>It’s the second most useful detecting technique of migraine trigger points, especially because positive Doppler sign 100% correlate with nerve-artery crossover or close relationship intraoperatively. It’s cheap.</td>
<td>Patients should be seen in the office during a MH attack.</td>
</tr>
</tbody>
</table>

![Fig. 2. Intraoperative picture: Isolation of the left STA.](image-url)
of the ZTN through temporal muscle and fascia, and the sentinel vessels are cauterized only when patients describe a pulsating pain in the temporal region.

For ATN decompression surgery, we usually mark a 1.5-cm incision where patients pinpoint the tender spot. A handheld Doppler is regularly used to locate the STA, since we observed 100% correlation rate between the trigger point identified and a close ATN-STA relationship (being either a simple crossover or a helical intertwining). Once the incision is made, dissection is taken with the help of blunt tipped scissors to expose and isolate both ATN and STA (Fig. 2), which is ligated both proximally and distally at the operative site.

RESULTS

These procedures are usually performed as 1-day surgery. No drainage tubes usually need to be positioned. Patients are commonly permitted to resume ordinary daily activities starting from second postoperative day (POD), light exercise by seventh POD, and regular exercise by 21st POD.11

Outcome

Authors have reported their results following temporal migraine deactivation surgeries.25-28 Regardless of the type of approach, deactivated trigger point, and surgery performed, outcomes observed were similar and are reported in Table 5. Nevertheless, ZTN deactivation surgery as described by Guyuron et al.23 achieved the highest MH elimination / >50% reduction rates. However, results cannot be easily compared since most of the patients included in these studies underwent also other MH sites deactivations, either at the time of, before, or after site II/V surgeries.

Authors’ Experience

From June 2011 till January 2018, we have performed MH decompression surgery over 168 patients with either frontal, occipital, or temporal migraine trigger sites.7-11 Among them, 41 suffered from temporal migraine, 32 patients had monolateral MH, while 9 had bilateral one. We performed a total of 56 temporal migraine deactivation surgeries. Thirty-four surgeries were performed at site V only because ATN-STA close relationship was observed intraoperatively. Whereas site II and V deactivation procedures were performed during the same operative session 6 times, since no ATN-STA close relationship was encountered. Therefore, we observed ATN-STA close relationship in 85.3% of patients; single STA-ATN intersection accounted for 83.7% of the cases, while helical intertwining accounted for 16.3%.
After a mean follow-up of 21 months (range, 3–67 months), patients complaining for temporal MH had 83% positive surgical outcome (50% complete MH elimination, 33% significant improvement). Seventeen percentage of patients did not notice any improvement since secondary trigger points occurred. Among these patients who had negative surgical outcome, 6 had concomitant site II and V deactivation surgeries while one had site V only deactivated (Fig. 5).

Adverse Events

Only rare minor complications are usually reported (eg, edema, hematoma/ecchymosis, numbness, and paresthesia). Numbness can occur (lasting <1 year, 163 days on average) in 5.7% of the patients. Intense itching, hypertrophic scar, incisional cellulitis, transient mild incisional alopecia, and hair thinning are also minor complications that have an incidence ranging from 1% to 5%. The most common complication after site II/V surgery is slight hollowing of the temple (54% incidence rate).

Nerve avulsion may be associated with neuroma formation, although it is not reported in literature of any neuroma following avulsion of ATN and/or ZTN.

Almost 90% of the patients can recognize more than 1 MH trigger site and surgical deactivation may be performed at all sites during the same surgical procedure depending on surgeon and patient preferences. Furthermore, 17.8% of patients can experience secondary trigger point emergence following primary migraine surgery.

MH recurrence may occur from 1 up to 3 months after surgery; thus, the result may be regarded as permanent only after the third postoperative month.

DISCUSSION

Despite strenuous efforts in determining MH etiology, it is still a matter of debate whether it is of central origin.
or peripheral genesis. Nevertheless, the striking results reported by surgeons performing decompression surgeries of trigger points in MH sufferers strongly support peripheral etiology of MH. Indeed various authors independently reported success rate higher than 80% in resolving or at least improving MH by decompressing irritated nerve from surrounding structures. Nonresponders might either have been misdiagnosed in their MH, or have undergone incomplete surgical decompression, or even have trigger point not addressed at all. Just like upper extremity compressive neuropathies, MH may be the resulting epi-phenomenon of chronic compression to cranial peripheral nerves.

Special relevance should be given to the close nerve/artery relationship that may intersect or intertwine each other, perhaps promoting irritation and therefore triggering MH attacks. Several studies have demonstrated a 34–80% rate of ATN-STA relationship, which may explain why some predisposed people have MH seemingly originating from this region. The same pulsatile irritation was addressed as main etiology in site IV MH, where close interaction between great occipital nerve and occipital artery has a 54% rate.

Wolff et al. have provided clear evidences of the vasomotor mechanism producing head pain. While symptoms as scotomas, paresthesia, blurred speech, photo/phono-phobia are due to vasocostriction of terminal branches of internal carotid artery, pulsatile distention of terminal branches of external carotid artery can determine traction and pressure stimuli to surrounding peripheral nerves deriving from the trigeminal nerve. This chronic irritation eventually results in throbbing, pulsating headache. In the later stage, antalgic contraction of surrounding muscles of head and neck can overcome the original vascular pain, resulting in chronic, dull, headache. For historical sake, Tunis and Wolff managed temporal migraine by ligating STA in 1953. Our experience corroborate the “vascular” theory. When dealing with the occipital trigger sites, we usually found a close connection with dilated (or frankly aneurysmatic) occipital arteries, the ligation of whose completely deactivated this trigger point. The same occurs when dealing with the AT site.

CONCLUSIONS

Further studies need to be carried out to precisely delineate MH pathophysiology and to identify all potential compression points at trigger zones. Finally, in our humble opinion, particular attention should be paid in future to the nerve/artery relationship as one of MH causes.

ACKNOWLEDGMENTS

This study did not need any ethical approval since it did not describe experimental studies on either humans or animals. However, the study has been conducted in accordance with the principles outlined in the Declaration of Helsinki. All participants provided informed consent.

REFERENCES