Background: Evidence of a paradigm shift towards epicranial neurostimulation treatment techniques aimed at the site of headache pain is beginning to populate the literature. This is most apparent by 2 recently published reports describing alternative approaches to peripheral nerve stimulation techniques for refractory migraine, including hemiplegic migraine.

Objectives: To contribute to the emerging literature on epicranial-based neuroaugmentative approaches which target site-specific areas of distinct, but relatively diffuse, headache pain. Specifically, we describe the feasibility of a novel neurostimulation technique: occipital nerve stimulation, combined with bilateral subcutaneous electrical stimulation over the temporal region, to treat a patient dually diagnosed with "complicated migraine" and occipital neuralgia. Integral to this report, key stimulation programming data are also presented to better distinguish the role of this form of therapy in migraine, or other headache forms, from both the clinical and biomedical perspectives.

Methods: Case presentation with literature review.

Results: At 24-month follow-up, headache onset had been reduced by more than 50%, including cessation of neurologic deficits that accompanied the patient's migraines. No complications or adverse side effects are reported. The programming data reported here supports a proposed mechanism of action concerning stimulation of the auriculotemporal nerve distribution/anterior temporal region for management of refractory pain in migraine.

Limitations: Case presentation provides only initial assessment of treatment safety, not conclusive evidence of treatment effectiveness. Future studies which consider “follow-the-path” epicranial approaches to peripheral nerve stimulation techniques for refractory headache pain are needed to better quantify outcomes and mechanisms of action.

Conclusions: In the single case reported here, the feasibility of a novel neurostimulation technique (occipital nerve stimulation/bilateral subcutaneous temporal region stimulation) to treat headache is presented. At the 24-month follow-up, no complications (such as loss of coverage due to lead displacement or lead fracture or erosion) or adverse side effects were reported. Finally, inclusion of fundamental programming data in reports on neuroaugmentative approaches to headache care will complement initiatives in research from the clinical and biomedical communities involved in this field.

Key words: Epicranial neurostimulation technique, peripheral nerve stimulation, stimulation parameters, complicated migraine, occipital neuralgia.
Oc
cipital neuralgia is a paroxysmal jabbing pain in the distribution of the greater occipital or lesser occipital nerves or of the third occipital nerve, and typically occurs from the effects of irritation secondary to entrapment of a nerve along its course (1). When the associated headache is medically intractable, favorable clinical evidence exists for the use of occipital nerve stimulation (ONS) (2-7).

In contrast, “complicated migraine” is a non-standardized term associated with migraine with aura (1). In the literature the term is often used interchangeably with migraine variants, and synonymously with complex migraine and migraine accompagnée (8). Rothner (8) defined complicated migraine as “syndromes associated with episodic, transient, and reversible neurologic dysfunction such as hemiplegic migraine, basilar artery (basilar-type) migraine, ophthalmoplegic migraine, and retinal migraine.” Treatment can be difficult due to the transient nature of the neurological impairments which confound pharmaceutical regimens. At the minimum, prophylactic use of acetylsalicylic acid is prudent (9,10). Conversely, triptans and the other vasoconstricting agents are contra-indicated due to risks of worsening the patient’s condition, if an episode is, in fact, vasoconstrictive rather than neurologic in origin (11).

In a recently published case series, Reed et al (12) discuss ONS combined with supraorbital nerve stimulation (SONS) to manage primary headache—and one case involved the treatment of a patient with hemiplegic migraine. In addition, other headache forms have been treated by combined neurostimulation techniques. For example, occipital and craniofacial/juxta-orbital pain managed by ONS combined with SONS (13,14) or combined with intraorbital nerve stimulation (IONS) (13).

By means of an alternative neurostimulation technique for chronic migraine, Simopoulos et al (15) reported successful use of bilateral stimulation of the auriculotemporal nerve to treat refractory pain in the bilateral temporal distribution. The authors propose that stimulation influences nociceptive fibers traversing the sutures of the calvaria (citing unreliable paresthesia coverage in the trigeminal distributions associated with ONS) (15,16).

Here we discuss the feasibility of a novel neurostimulation technique in a challenging case of headache based on 24-month follow-up: ONS combined with bilateral subcutaneous stimulation over the temporal region (Fig. 1) to treat a patient dually diagnosed with complicated migraine and occipital neuralgia. It is noted that the time-frame for this report closely parallels the treatment timelines reported by Reed et al (12) and Simopoulos et al (15). Hence, this article contributes to the emerging literature on epicranial-based neuroaugmentative approaches which target site-specific areas of distinct—but relatively diffuse—headache pain. Integral to our report, key neurostimulation programming data are also presented—to help improve our understanding of the role of this form of therapy in migraine, or other headache forms, from both the clinical and biomedical perspectives.

Case Report

Our patient is a 44-year-old Caucasian woman under the care of an experienced neurology headache specialist for 8 years prior to referral to our practice. During the patient’s consultation in 2007, she complained of a 28-year history of headaches which emanated from the occipital base without injury or inciting event, with an incidence rate of 4-5/week. These headaches were diagnosed as occipital neuralgia based on the International Classification of Headache Disorders-2 (1). In addition, her past medical history included absence seizures (with familial history); however, the last episode was 8 years ago, and subsequently her regimen of divalproex sodium was more recently discontinued. In 2002, a Le Fort I osteotomy was performed to correct malocclusion as opposed to treat temporomandibular joint dysfunction.

Notably, in 2004, the patient began experiencing headaches accompanied by marked phonophobia, temporary loss of vision (both eyes), slurred speech, ptosis (right eye), and hemiplegia (left leg). Not only were these headaches incapacitating, but the auras also imposed temporary but prolonged physical and communicative limitations. As a result of such symptoms, the patient’s overall quality of life was negatively affected. The timing of such migraine episodes was variable with no known precipitating factors; incidence was reported as one per month with a maximum duration of headache of 72 hours. Neuroimaging and electroencephalographic studies were unremarkable, and no risk factors for atherosclerotic stroke were present. Based on symptomology and the neurological workup, complicated migraine was added to the patient’s headache profile by the referring neurologist.

Because of the characteristics of the patient’s migraines, the ergot derivatives and triptans or other vasoconstricting agents were contra-indicated. Dark room
retreat and cold pack or moist heat applications were ineffective at aborting onset or alleviating severity. Pain management consisted of analgesics and opioids for mild to moderate onset of headaches. For severe onset, the patient either sought treatment at her local emergency department or her neurologist administered rescue medications, which consisted of acute pain and antiemetic drugs and steroids. Such treatments produced minimal abatement of her headaches. Furthermore, the duration of therapeutic greater occipital nerve blocks (and botulinum toxin injections administered at multiple sites in a halo/crown fashion to follow-the-path of her pain) was short-lived.

**Neurostimulation Trial**

At the time of our initial evaluation, the patient described her headaches with dual origins—a sharp jabbing pain at the occipital base, and a separate and distinct bilateral sensation of tightening/stabbing that encompassed the temporal and retro-orbital regions. She further qualified her headache pain as more intense on the right, but more frequent on the left. We focused on suppressing the site of headache pain (temporal and occipital regions). In January 2008, a 7-day trial of peripheral nerve stimulation was implemented with 3 wire leads (Medtronic Inc., Minneapolis, MN).
Bilaterally, one Quad lead (1x4) was subcutaneously placed superficial to the anterior border of the temporalis muscle (and over the sphenotemporal suture) to achieve stimulation over the auriculotemporal nerve distribution; and a single Standard lead (1x8) was placed from the right to apply ONS bilaterally (Fig. 2A). At follow-up the patient reported she had been headache free throughout the trialing period.

**Implant Procedure**

Subsequently, in June 2008, in accord with the patient’s goals and informed consent, the described lead arrangement was implanted with one modification. We postulated that clinical benefit might be gained by expanding the available coverage over the temporal region—based on the terminal branching patterns of the trigeminal nerve (as well as the communicating branches between the auriculotemporal nerve and the lesser/greater occipital nerves) (17,18). Therefore, a Standard lead (1x8) was substituted for the Quad lead (1x4), and was placed bitemporally and subcutaneously in a horizontal fashion over the inferior border of the temporalis muscle (Fig. 2B). This intervention did not require approval from the institutional review board, and the patient tolerated the implant surgery well. Due to the nature of the patient’s migraines, it was felt unethical to leave the system turned off following implant.

It is important to also point out that 2 RestoreUltra neuropulse generators (Medtronic Inc., Minneapolis, MN,) were implanted. One generator (right gluteal pocket) accepted the ONS lead and the right temporal region lead, a second generator (left gluteal pocket) accepted the lead from the left temporal region. It was reasoned this battery configuration would 1) benefit the patient by providing dedicated power sources, and 2) permit tunneling of each lead respectively along an ipsilateral course. The latter might, in conjunction with the strain-relief loops, also help reduce the risks of displacement or lead fracture secondary to biomechanical-related strain and friction.

**Follow-up**

At the 24-month follow-up, no complications (such as loss of coverage due to lead displacement or lead fracture or erosion) or adverse side effects were reported. More than a 50% reduction in headache onset was reported, and notably, the patient has not experienced the neurologic deficits that defined her migraines. Finally, only one emergency department treatment was sought and the use of rescue medications has de-

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Fig. 2. Panel A: Lead placement during the trial period. Panel B: During the implant procedure, following bitemporal placement for subcutaneous electrical simulation superficial to the temporalis muscle, an introducer needle was positioned posterior to the C1-C2 level to accept a lead for occipital nerve stimulation.
creased. Such outcomes have contributed to an overall improvement to the patient’s quality of life.

Stimulation use is continuous over a 24 hour period. Both pulse rate and amplitude are controlled by the patient. Interestingly, for each of the 3 leads, the patient prefers slower “drumming” pulse rates (10 Hz), and attributes this frequency selection to providing a more satisfactory prophylactic stimulus. Figure 3 provides detailed information concerning programming and stimulation parameters.

**Discussion**

Given the complex presentation of headache in our patient, the most notable outcome reported was cessation of the transient neurologic deficits which characterized the patient’s migraines. Although a placebo effect cannot be completely excluded for the results achieved here, given the continuation of response over the follow-up period, there is probably minimal placebo effect.

![Diagram](image.png)

Fig. 3. A schematic representing the most utilized program setting: anode (+) and cathode (-) configuration; stimulation parameters; and electric fields. All programs (B1, B2, and B3) run simultaneously. Note: B3 is programmed to activate the distal end of the right temporal lead (identified here by the oval ring).
**The Neurostimulation Technique**

The use and positioning of the 1x8 lead at implant relative to the 1x4 lead used during the trialing period, ideally, permits stimulation over the auriculotemporal nerve as well as both the sphenotemporal and temporo-occipital sutures (based on the employed cathode/anode configuration). The significance of this is seen in a supposition made by Simopoulos et al (15) on the possible mechanism of pain relief from stimulation over the auriculotemporal nerve distribution/anterior temporal region. Specifically, the authors make use of a theory that stimulation (and modulation) of nociceptive fibers traversing the bony sutures of the calvaria might be achieved based on the animal model (Fig. 4) (15). In our opinion, such a hypothesis appears reasonable given the limited evidence coupled with the lead placement and cathode/anode selection reported here (which might best facilitate regulation of the electric field adjacent to the suture lines).

It is noted that all patients in the Trentman et al study (16), with respect to ONS-only paresthesia coverage (which included information on neuropulse settings), were implanted with a constant-voltage generator/system, analogous to our case, with follow-up over a nearly identical period since implant. This enabled us to easily compare usage/stimulation parameters between techniques (i.e., ONS-only in contrast to the technique introduced here). Moreover, other authors have described common ONS parameters and use patterns with respect to treatment for various headache forms: Weiner and Reed (2), Popeney and Aló (19), as well as Trentman et al (20) in a pioneering report on the Bion microstimulator. Unfortunately, information about programming settings and stimulation parameters was not found in the reports on patients who received bilateral auriculotemporal nerve stimulation (15) or the combined techniques (ONS/SONS [12-14] or ONS/Ions [13]) for management of headache pain, in-

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**Fig. 4.** (Panel A) Extracranial origin of intracranial pain – action potentials generated at extracranial collaterals of meningeal pain fibers (1) spread antidromically to collaterals that terminate inside the cranium (2). Result: local release of proinflammatory neuropeptides and activation of neighboring meningeal nociceptors (2). (Panel B) Intracranial origin of extracranial pain – action potentials generated at intracranial meningeal pain fibers (1) spread antidromically to collaterals that terminate outside the cranium (2). Result: local release of proinflammatory neuropeptides in the scalp and activation of neighboring somatic nociceptors (3). *Original site of activation. Red dots represent local release of inflammatory neuropeptides (e.g., calcitonin gene-related protein [CGRP], substance P. (Copyright permission granted by the American Headache Society [19].)
Combined Temporal & Occipital Epicranial Neurostimulation

cluding migraine. In addition, in our opinion, such data is not routinely made available in the neuroaugmentative migraine literature. Therefore, presented as a quick reference for the reader, Table 1 summarizes relevant literature according to data on 1) headache form treated; 2) neurostimulation technique – stimulation system; and 3) stimulation parameters.

The inclusion of key data on neuropulse settings (programming options, cathode/anode designation, stimulation parameters) is important not only for further peripheral nerve stimulation research, but is of specific interest for clinical management and neuroaugmentative device research concerning headache (19). To this point, analysis of the neuropulse settings, with careful attention to patient preferences on the “feel” of stimulation (as reported here) and usage patterns is beneficial to advancing neurostimulation therapies, and is valuable for engineering next generation power sources (generator/battery) or modes of stimulation delivery (21).

**CONCLUSION**

In summary, in the single case reported here, the feasibility of a novel neurostimulation technique (ONS/bilateral subcutaneous temporal region stimulation) to treat headache was presented. Clinical outcomes were based on 24-month follow-up. No complications or adverse events were noted. Headache onset has been decreased by more than 50%, and notably, the neurologic symptoms which accompanied the patient’s migraines ceased. Case presentation provides only initial assessment of treatment safety, not conclusive evidence of treatment effectiveness. However, our report supports the initial experiences presented by both Reed et al (12) and Simopoulos et al (15). Clearly, future studies which consider “follow-the-path” approaches to peripheral nerve stimulation techniques for refractory pain in migraine are necessary to better quantify outcomes and mechanisms of action. Finally, while parameter settings might be largely determined by individual patient preferences (i.e., the slow, “drumming” pulse rates our patient enjoys bitemporally and occipitally), from both clinical and biomedical perspectives such information is important to track as research initiatives on neurostimulation techniques are advanced, especially when new techniques are utilized.

Table 1. Stimulation parameters.

<table>
<thead>
<tr>
<th>Headache Form</th>
<th>Neurostimulation Technique – System</th>
<th>Pulse rate (Hz)</th>
<th>Amplitude (V)</th>
<th>Pulse-width (μS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deshpande et al (present report)</td>
<td>Complicated Migraine with Occipital Neuralgia - ONS with bilateral temporal region – constant-voltage, single channel</td>
<td>10</td>
<td>1.43</td>
<td>390</td>
</tr>
<tr>
<td>Simopoulos et al (15)</td>
<td>Migraine - Bilateral auriculotemporal nerve distribution – constant-current, single channel</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Not reported</td>
</tr>
<tr>
<td>Reed et al (12)</td>
<td>Migraine (one case with Hemiplegic Migraine) - ONS/SONS – constant-current, single channel</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Not reported</td>
</tr>
<tr>
<td>Slavin et al (13)</td>
<td>Craniofacial pain - ONS/SONS and ONS/IONS – constant voltage, single channel; constant-current, single channel; constant-current, multiple channel</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Not reported</td>
</tr>
<tr>
<td>Trentman et al (20)</td>
<td>Cluster, Migraine, Hemicrania Continua - ONS – microstimulator – current controlled</td>
<td>45-60</td>
<td>Not reported</td>
<td>200-350</td>
</tr>
<tr>
<td>Trentman et al (16)</td>
<td>Cluster, Migraine, Hemicrania Continua - ONS – constant voltage, single channel</td>
<td>26-40</td>
<td>1.07</td>
<td>400</td>
</tr>
<tr>
<td>Popeney et al (19)</td>
<td>Transformed Migraine - ONS – constant voltage, single channel</td>
<td>55</td>
<td>3.2</td>
<td>400</td>
</tr>
<tr>
<td>Weiner and Reed (2)</td>
<td>Occipital Neuralgia - ONS – constant voltage, single channel</td>
<td>60-130</td>
<td>0.5-2</td>
<td>60-180</td>
</tr>
</tbody>
</table>
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