Managing cluster headache with sphenopalatine ganglion stimulation: a review

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Abstract: Cluster headache (CH) is a primary headache and considered as one of the worst pains known to man. The sphenopalatine ganglion (SPG) plays a pivotal role in cranial autonomic symptoms associated with pain. Lesioning procedures involving the SPG and experimental acute SPG stimulation have shown some degree of efficacy with regard to CH. A neuromodulation device, chronically implanted in the pterygopalatine fossa, has been specifically designed for acute on-demand SPG stimulation. In a pilot placebo-controlled study in 28 patients suffering from refractory chronic CH, alleviation of pain was achieved in 67.1% of full stimulation-treated attacks compared to 7% of sham stimulation-treated attacks ($p<0.0001$). Long-term results (24 months; 33 patients) confirmed the efficacy of SPG stimulation as an abortive treatment for CH attacks. Moreover, 35% of the patients observed a >50% reduction in attack frequency, suggesting that repeated use of SPG stimulation might act as a CH-preventive treatment. Globally, 61% of the patients were acute responders, frequency responders, or both, and 39% did not respond to SPG stimulation. The safety of SPG microstimulator implantation procedure was evaluated in a cohort of 99 patients; facial sensory disturbances were observed in 67% of the patients (46% of them being transient), transient allosthenia in 3%, and infection in 5%. SPG stimulation appears as a promising innovative, efficient, and safe therapeutic solution for patients suffering from severe CH. It has shown its efficacy in aborting CH attacks compared to placebo stimulation, suggesting that it is particularly adapted for CH patients who are not sufficiently improved by abortive treatments such as sumatriptan and oxygen. However, further studies comparing SPG stimulation with standard abortive and/or preventive CH treatments will be necessary to define more precisely its place within the management of severe chronic and/or episodic CH.

Keywords: cluster headache, primary headache, sphenopalatine ganglion, stimulation, neuromodulation

Cluster headache

Cluster headache (CH) is a primary headache and is the most common form of trigeminal autonomic cephalalgia that are grouped in the third section of the International Headache Society Classification (ICHD-III).1 CH is considered to be one of the worst pains known to man. Individuals often describe the pain like having a “red hot poker” penetrating one eye. They report that the intensity is so extreme it is unlike anything they have ever experienced, and female patients describe each attack as being worse than childbirth. The intensity of pain is so high that it can induce suicidal thoughts, and so CH has been named as “suicidal headache”. Such severity justifies greater attention to this disease.

CH is characterized by recurrent attacks of very severe unilateral pain, usually located in and around the orbit. During an attack, ipsilateral cranial autonomic
symptoms (lacrimation, conjunctival injection, nasal congestion or rhinorrhea, ptosis, edema of the eyelid or the face, sweating of the forehead or the face, miosis) accompany the pain, and behavior is characterized by a marked sense of agitation and necessity to move. Attacks generally last 15 minutes to 3 hours and occur from once every day to eight times daily, sometimes with a striking circadian rhythmicity, with some patients reporting a predictability of onset during the day or, more frequently, during the night, waking from sleep. The term CH is related to the tendency of attacks to cluster together into bouts that last several weeks or months. In the episodic CH, the bouts occur at certain times of year and are separated by periods of remission, which last at least a month. Frequently, episodic CH has a striking circannual periodicity, with the bouts occurring in the same month of the year. However, about 10%-20% of individuals have the chronic form of CH and have continuous attacks with no respite or remission periods lasting less than 1 month.1

While there is currently no cure for CH, the available treatment has become more effective in the last 20 years, especially the acute treatments.2 Acute treatment is used to stop the pain once it has started, and the key element is speed to reduce the excruciating pain as fast as possible. For this purpose, conventional analgesia is ineffective, and only two treatments are validated, oxygen inhalation and subcutaneous sumatriptan. Preventive and transitional treatment, attempting to reduce frequency and intensity of attacks, is based on verapamil, lithium, or corticosteroids. Due to failures of these treatments in refractory patients, neuromodulation approaches have been introduced in the management of CH during the last 10 years. Invasive methods, namely deep brain stimulation (DBS), occipital nerve stimulation (ONS), and sphenopalatine ganglion (SPG) stimulation are reserved for patients with severe and refractory CH, but emergent noninvasive methods might serve as useful adjuncts to more conventional therapies.

CH physiopathology and SPG involvement

The pathophysiology of CH has not been completely identified yet. The individual CH attack is regarded as an activation of the trigemino-autonomic reflex, and the posterior hypothalamus is supposed to play a pivotal role, probably via a trigemino–hypothalamic pathway.3 This pathophysiological model explains the three major features of CH: unilateral trigeminal distribution of pain, ipsilateral cranial autonomic symptoms, and circadian episodic pattern of attacks.4 Nevertheless, many issues still remain unanswered. Few neuroimaging studies have explored CH patients,5 suggesting structural6 and functional7 changes in a posterior hypothalamic region. Finally, a genetic component is suggested by the increased familial risk (14-fold and twofold increased risk in first- and second-degree relatives, respectively).8

The SPG is believed to play a pivotal role in cranial autonomic symptoms associated with pain and might be involved in headache pain. Recent hypothesis suggests that, during CH attacks, there is an activation of the parasympathetic superior salivatory nucleus and SPG parasympathetic fibers, inducing neuropeptides release (especially calcitonin gene-related peptide) and vasodilatation of the cerebral and dural blood vessels, which activate meningeal nociceptive fibers projecting to the trigeminal ganglion and nuclei. These processes induce referred pain in the periorbital region. Thus, blockage of the SPG, by any means, might theoretically interfere with this pathological process and treat CH attack.

The SPG contains sensory fibers innervating the posterior nasopharynx, sympathetic fibers from the superior cervical ganglion via the vidian nerve, and preganglionic parasympathetic fibers coming from the superior salivatory nucleus via the greater petrosal nerve and connecting, within the SPG, with postganglionic fibers. The SPG is located in the pterygopalatine fossa (PPF), behind the posterior wall of the maxillary sinus (Figure 1), bordered posteriorly by the pterygoid process, superiorly by the sphenoid sinus, and medially by the palatine bone. Laterally, the PPF communicates with the
infratemporal fossa. The maxillary division of the trigeminal nerve (V2) passes the foramen rotundum located superolaterally within the PPF, and is in close contact with the SPG.

**SPG as a therapeutic target for CH**

For more than a century, SPG has been targeted to treat primary headache, either by surgical ganglioneectomy, lidocaine application, percutaneous lesioning, and, more recently, electrical stimulation.

Several lesioning procedures involving the SPG have been proposed for the treatment of chronic CH. The success rates of ganglioneectomy, radiofrequency lesion, or blocks varied from 46% to 85%. However, repeated access to the SPG was often necessary because, in most of the cases, the benefits were transient. Efficacy of SPG radiofrequency lesioning, using a percutaneous infra-zygomatic approach, has been studied in two series, cumulating 25 patients (mean follow-up 12–24 months). The mean frequency of chronic CH (CCH) attacks was decreased by half after the procedure. However, the complication rate was high and included such complications as epistaxis (80%), lesion of the maxillary division of the trigeminal nerve (40%), and transient hypoesthesia of the palatine area (90%).

Stereotactic radiosurgery (SRS) of the SPG had been proposed initially to treat CH in combination with trigeminal nerve SRS and seemed slightly more efficient than trigeminal nerve SRS alone. SRS targeting only the SPG has been reported to be efficient in single cases. Globally destructive procedures on the SPG appeared to be partially efficient on CCH, which has encouraged the development of new nonlesional procedures targeting the SPG.

**Electrical stimulation of the SPG**

Based on the effect of lesioning procedures on the SPG, its stimulation was proposed for the treatment of CH by Ibarra for the first time in 2007, using an implantable device in a single case. The author mentioned a marked reduction of pain with acceptable safety. The only complication reported was hardware failure, after which the patient’s headaches worsened. Interestingly, his condition improved after replacement of the stimulator, suggesting the efficacy of SPG stimulation. In 2010, Ansarinia showed that acute SPG electrical stimulation was able to abolish provoked CH attack within few minutes in six patients suffering from refractory chronic CH. In this study, SPG stimulation was delivered using a transient electrode, implanted percutaneously by an infra-zygomatic approach.

Based on these preliminary results, a chronically implantable neuromodulation device (“Pulsante”), specifically designed for acute SPG stimulation, has been developed by the company Autonomic Technologies (ATI, Redwood City, CA, USA), in order to abort the CH attacks on-demand (Figure 2). The neurostimulator device is implanted in the PPF, along the posterior wall of the maxillary bone, and fixed to the zygomatic process with a screwed plate, with the lead being placed in contact with the SPG (Figure 3). The neurostimulator does not contain a battery but is activated and powered by a remote controller using radiofrequency energy.

To evaluate the safety and efficacy of SPG stimulation, we conducted a Medline search using the terms “sphenopalatine ganglion”, “stimulation”, and “cluster headache”, excluding case reports, technical reports, review articles, and redundant series. Only two articles, originating from the same multicentric group of physicians, were finally considered.

The safety and efficacy of this SPG stimulation device have been assessed in a pilot multicenter study (Pathway CH-1 study), in 28 patients suffering from refractory CCH with at least four attacks per week. In this sham-controlled study, each CH attack was randomly treated with 15-minute full stimulation, subperception stimulation, or sham stimulation.

![Figure 2 Principle of the Pulsante system (Autonomic Technologies, Redwood City, CA, USA) designed for acute on-demand stimulation of the SPG.](image)

**Notes:** (A) External programming hardware. The program software (1) allows the physician to test and program the neurostimulator parameters to customize stimulation for each patient. The remote controller (2) is a rechargeable handheld device used by the patient to activate the neurostimulator, and it is also used by the physician to communicate with the neurostimulator during programming sessions. (B) The miniaturized neurostimulator is implanted in the PPF to stimulate the SPG and is fixed by screws to the maxillary bone. (C) Activation of the stimulation by the patient to abort the CH attack by application of the remote controller against the patient’s cheek.

**Abbreviations:** CH, cluster headache; PPF, pterygopalatine fossa; SPG, sphenopalatine ganglion.
(last two stimulation modes being nonactive). Absence of pain was defined as reduction of pain intensity by 2 points or more according to a Categorical Pain Scale to 0 points, and alleviation of pain to 0 or 1 point(s). Alleviation of pain was achieved in 67.1% of full stimulation-treated attacks compared to 7.4% of sham-treated and 7.3% of subperception-treated attacks \((p<0.0001)\). Absence of pain was achieved in 34.1% and 1.5% of attacks after active stimulation and sham stimulation, respectively \((p<0.0001)\). For CH attacks treated with full stimulation, pain relief was achieved in 55.5% and 60.6%, at 30 and 60 minutes, respectively. Nineteen of 28 (68%) patients experienced a clinically significant improvement, but only 32% achieved a pain relief in more than 50% of the treated attacks. Moreover, 43% of the patients experienced a reduction >50% of attack frequency (average 88%).

Long-term (24 months) results of the open phase of this study have been recently published,\(^20\) in 33 patients. All patients were initially included in the Pathway CH-1 study, although 11 of them were not included in the first analysis (for time reasons). Moreover, 10 patients from the initial study were excluded from this long-term analysis, because they no longer had the stimulator implanted or due to previous protocol noncompliance. Across all 33 patients, a total of 5,956 attacks were treated. Effective treatment (alleviation and/or absence of pain) was achieved in 65% of CH attacks, with a delay of 11.2 minutes on average, including 50% becoming pain free. Fifteen out of 33 patients (45%) were considered as acute responders (at least 50% of attacks were successfully treated). In 79% of the attacks, patients did not report the use of acute medication. Moreover, 11 patients out of 31 (35%) observed a >50% reduction of the weekly attack frequency 24 months after implantation compared with the 4-week period before implantation (average reduction was 83%). These data suggested that repeated use of the SPG stimulation might act as a CH prophylactic treatment, although spontaneous transformation from chronic to episodic forms of the disease cannot be excluded. Globally, 61% of the patients were acute responders, frequency responders, or both, and 39% of the patients did not respond to SPG stimulation.

Headache disability (Headache Impact Test [HIT-6™] scale) was significantly improved (decrease >2.3 points) in 55% of the patients, and 21 patients out of 33 (64%) were able to decrease or stop (9 cases) their prophylactic CH medication.

**Complications and side effects**

In the pilot Pathway-CH1 study,\(^20\) 81% of the patients experienced transient, mild-to-moderate hypoesthesia within the maxillary (V2) nerve territory, resolving within 3 months in most of the cases. More recently, the safety of the surgical implantation procedure of the Pulsante SPG stimulator has been evaluated in a cohort of 99 patients, including 43 patients of the Pathway CH-1 study and 56 patients from the Pathway-R1 registry.\(^21\) Eighty one percent of the patients experienced at least one adverse event, most of them being transient. Sensory disturbances were the most frequent complications, observed in 67% of the patients, 46% of them resolving within a mean delay of 104 days. Transient allodynia was rare (3%). Pain and/or swelling were reported by 47% of the patients, resolving in 80% of the cases with a mean delay of 68 days. Dry eye (3%, resolving in 40% of cases), transient trismus (8%), and limited jaw movements (6%) were also reported.

Infection rate was 5%.

Microstimulator revision procedures were performed in 13 cases due to inappropriate initial
placement of the stimulating electrode within the PPF. Five stimulators were explanted.

Although frequent, most (92%) of the adverse events were transient and evaluated as mild or moderate. The authors concluded that SPG microstimulator insertion procedure has sequelae comparable to other oral cavity procedures including tooth extractions, sinus surgery, and dental implant placement. Moreover, the technique is recent, and the rate of surgical complications will likely decrease with progression of the learning curve, further refinement of the surgical procedure and tools, and the use of neuronavigation systems.22

**Limitations**

Although SPG stimulation appears as a promising, safe, and efficient technique to treat CH attacks, data assessing its safety and efficacy remain limited, and the quality of evidence concerning this therapy, according to the GRADE assessment system,23 is low. All the currently available data come from a single study and a registry, including a limited number of patients (respectively, 43 and 56 patients), and both of these studies were promoted and supported by the company commercializing the implantable device. In the pilot study,20 15 out of the 28 reported patients have had less than 10 CH attacks during the randomized phase, meaning that the conclusions were based mostly on the analysis of the remaining 13 patients. This study compared active stimulation with sham stimulation and demonstrated that SPG stimulation therapeutic effect was not due to a placebo effect. However, no study compared SPG stimulation with the current standard abortive treatment, namely sumatriptan injection and/or oxygen inhalation, in terms of efficacy, delay between treatment administration and pain relief, amplitude of pain relief, respective constraints and safety, and costs. Further studies will have to address these points. Similarly, although a preventive effect has been suggested by the association of repeated SPG stimulation and decrease of attack frequency, no study compared SPG stimulation with CH prophylactic treatment or other surgical alternatives for refractory chronic CH, namely ONS and DBS.

These data will be needed to determine the exact place of SPG stimulation in the management of CH. Moreover, all the previous data have been collected in refractory CH, and no published data are available regarding the interest of SPG stimulation in episodic CH.

**Place of SPG stimulation in the management of CH**

Despite these limitations, SPG stimulation is a promising, innovative, efficient, and safe therapeutic solution for patients in severely disabling painful condition. As an acute treatment to abort attacks, this procedure is particularly adapted for patients who do not respond to current abortive treatment as sumatriptan and oxygen. In a recent study on refractory CCH patient candidates for ONS,24 19% of them did not use subcutaneous sumatriptan or oxygen because these treatments were ineffective, not tolerated, or contraindicated. These patients, experiencing 20 CH attacks per week on average, would benefit dramatically from an alternative abortive treatment such as SPG stimulation. About 30% of refractory CCH attacks do not respond to ONS.24 Considering their respective risks, we consider that ONS and SPG stimulation should be proposed before DBS in medically intractable CCH patients. However, the choice between ONS and SPG stimulation as a first surgical treatment is difficult and should be based on benefits/risks ratio, patient’s anatomy and headache characteristics, local availability of the different techniques, regulatory and reimbursement issues, and national guidelines.

Clinical criteria for invasive surgery, initially DBS, in CCH have been proposed by a group of experts as follows:25 CCH for at least 2 years, at least one attack per day, resistance to pharmacotherapy, headache “locked” to the same side, normal neurological examination, and absence of psychiatric comorbidity. Selection criteria for SPG stimulation may be moderated and/or extended, considering that SPG stimulation is less invasive and risky than DBS and acts as an abortive treatment. An expert consensus proposed in 2014 selection criteria for SPG stimulation to be as follows: 1) CCH according to ICHD classification;26 2) documented for at least 2 years; 3) refractory to medical treatment,27 which is defined as failed adequate trials of preventive medication, alone or in combination (failure being defined as no therapeutic or unsatisfactory effect, intolerable side effects, or contraindications to use; adequate being defined as appropriate dose and appropriate length of time); 4) with substantial number of weekly attacks; 5) severe psychosocial impairment; and 6) attacks being side locked or occurring predominantly (>90%) on the current side for the last 12 months. Patients must be able to understand the device use and ideally be able to document their attacks on a diary. Implantation should be performed in a center with expertise in headache diagnosis and treatment, and having a surgical team with specific training and expertise in SPG stimulator implantation.

No recommendation was given concerning episodic CH, as the clinical data currently published have been obtained in CCH. However, patients with episodic CH (ICH-D-III classification) for at least 2 years, with active periods lasting several months,
and having several daily attacks not or poorly responding to abortive treatments might benefit from SPG stimulation.

Exclusion criteria for SPG stimulation have been identified as follows: previous history of destructive procedure on the SPG ganglion; severe psychiatric disorder (other than reactive depression); particular anatomical conditions as narrow lateral opening of the PPF (evaluated on preoperative computed tomography scan); recent (<4 months) facial surgery in the PPF area or zygomatico-maxillary buttress; active regional infectious process (periodontal disease, open carious lesions, osteomyelitis, abscess, etc); previous (<6 months) facial malignancy or radiotherapy; and craniofacial disease requiring frequent magnetic resonance imaging (Pulsante system seems compatible with 1.5T magnetic resonance imaging but generates image artifacts) or surgery in the next months.

In a position statement on neuromodulation techniques in chronic headaches published in 2013, a group of experts from the European Headache Federation considered that, given that only one placebo-controlled study was available, SPG stimulation had to be seen as experimental until further studies are presented. This opinion was shared by the guidelines for the treatment of CH published by the French Headache Society in 2014, who recommended, in addition, to reserve this therapy to tertiary headache centers. Pulsante system has obtained a CE mark (Conformité Européene [European Conformity]) for the treatment of CH but is not yet approved by the US Food and Drug Administration, and is not yet reimbursed by all the health care providers.

Conclusion
SPG stimulation appears as a promising innovative, efficient, and safe therapeutic solution for patients suffering from severe CH. SPG stimulation has showed its efficacy to abort CH attacks versus placebo stimulation, suggesting that it is particularly adapted for CH patients who are not sufficiently improved by currently available abortive treatments such as sumatriptan and oxygen. Additionally, the pilot study suggested that repeated SPG stimulation might have a preventative action on CH attack frequency. However, further studies comparing SPG stimulation with standard abortive and/or preventative CH treatments will be necessary to define more precisely its place within the management of severe chronic and/or episodic CH.

Disclosure
D Fontaine and M Lanteri-Minet are consultants for Autonomic Technologies and Medtronic. The authors report no other conflicts of interest in this work.

References


