A case report of complex auricular neuralgia treated with the great auricular nerve and facet blocks

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Introduction

The great auricular nerve (GAN) is a cutaneous branch of the cervical plexus originating from the C2 and C3 spinal nerves.1 It innervates the skin over the external ear, the angle of the mandible and the parotid gland. It communicates with the ansa cervicalis. Great auricular neuralgia is rarely diagnosed in clinical practice and can be refractory. We present a new approach using ultrasound-guided nerve blocks.

Case: We present a case of a 41-year-old female with paroxysmal ear pain accompanied by dysautonomia, tingling in the tongue, dysphagia, dysarthria and abdominal symptoms. No significant findings were found on cervical and brain imaging. The patient responded partially to a great auricular nerve block. A combined approach using this block with facet block of C2 and C3 induced a more pronounced and prolonged benefit.

Conclusion: Great auricular neuralgia is not often encountered in practice and can be accompanied by symptoms originating from the ansa cervicalis network. A combined approach of nerve blocks can be considered in refractory cases.

Keywords: ansa cervicalis, neuropathic pain, facet block, red ear syndrome, great auricular nerve

Case presentation

The patient is a 41-year-old female who presented with a 1-year history of paroxysmal left ear pain and tingling sensation in her tongue. The pain in her ear radiated to the ipsilateral side of her face as well as posteriorly in the distribution of the lesser occipital nerve. She reported associated jaw clenching and allodynia to teeth brushing and mastication. The pain worsened at the end of the day to a severe intensity, at which time there was red discoloration of the ear. The patient alternated between a constant dull aching pain at 5 out of 10 and periods of paroxysmal pain reaching 8 out of 10 on a numerical rating scale and lasting between 1 and 5 consecutive days.
During exacerbations, the patient also complained of dysphagia and dysarthria. Other symptoms included post-prandial abdominal pain and diarrhea.

There was a history of sports-related mild traumatic head injury as a young adult and a whiplash involving subluxation of the C4–C5 cervical vertebrae. Past medical history also included mononucleosis, two previous superficial lower extremity venous thromboses postpartum and heterozygous factor V Leiden thrombophilia.

Examination of her cranial nerves revealed diffuse hypoesthesia to touch, pinprick and temperature in the left trigeminal distribution. No intraoral trigger points were identified. There was decreased sensation to touch and pinprick in the area of the greater and lesser occipital nerves. The gag reflex on the left side was absent. Lateral neck flexion to the right was limited with diffuse tenderness on palpation of the left paracervical muscles. Neck compression testing for radiculopathy and facet syndrome using Spurling’s maneuver was able to provoke the patient’s red ear discoloration after a 15-second compression.

The patient’s workup included a 3T cerebral angiographic magnetic resonance imaging with contrast, Constructive Interference in Steady State sequence imaging of the circle of Willis and magnetic resonance imaging of the cervical spine. A contact between the superior cerebellar artery and the cisternal portion of the left trigeminal nerve was seen, but without evidence of compression. A chest X-ray was normal. Rheumatological workup showed mildly elevated antinuclear antibodies, but was otherwise inconclusive for collagenosis. Gastrointestinal workup, including food allergy testing, gastroscopy and colonoscopy with biopsies were negative.

Drug trials included acetaminophen, naproxen, tricyclic antidepressant, gabapentinoids, carbamazepine, topical analgesics, bupropion, muscle relaxant, tapentadol, morphine, hydromorphone and methadone. Pharmacotherapy was unsuccessful in this patient. In addition, she developed a multiple drug intolerance syndrome, including a probable Stevens–Johnson with carbamazepine.

She was evaluated by an allergist. Due to the serious nature of side effects, all pain medications had been stopped at least 2 weeks prior to the ultrasound-guided infiltrations described below.

A diet at the patient’s own initiative proved slightly successful in reducing her postprandial abdominal pains. The patient was initially diagnosed with trigeminal neuralgia. The diagnosis was later revised to great auricular neuralgia, given the distribution of the pain. Additional symptoms were attributed to a dysfunction at the level of the ansa cervicalis.

We initially performed a GAN block using ultrasound guidance (Figure 1). We used a GE Logiq e® Ultrasound machine equipped with a 12 MHz high-frequency transducer. The GAN was identified at its emergence from the posterior border of the sternocleidomastoid muscle, at the level of the cricoid cartilage located by palpation. Once the hypoechoic round shaped nerve was visualized, a 25G, 50 mm needle was introduced in plane with the ultrasound probe, and we injected a mix of 1 mL 2% xylocaine, 1 mL 0.5% bupivacaine and 1 mL of triamcinolone (40 mg). This technique proved successful in relieving the patient’s paroxysmal earaches for over 1 month with reduction of pain intensity of 60%. The patient reported improvement within 48 hours following nerve block of the GAN. No adverse events were reported.

Three months later, the patient’s pain was back at baseline. We combined the GAN block with ultrasound-guided facet blocks of the C2 and C3 spinal nerves as well as lesser occipital nerve block. We injected 2 mL of the same analgesic mix at each infiltration point. The combination of these blocks reduced not only the intensity of her paroxysmal ear pain, but also the duration of exacerbations from 1 to 5 days to <24 hours and the background pain by >50%. The relief was more prolonged, lasting for over 2 months.

Figure 1 Ultrasound-guided transverse view of the GAN.
Abbreviations: GAN, great auricular nerve; PSCM, posterior border of sternocleidomastoid muscle.
Discussion

The GAN arises from the ventral rami of C2–C3 roots, along with two other sensory branches, the lesser occipital nerve and the transverse cervical nerve. The fourth cervical cutaneous nerve and the supraclavicular nerve arise from C3–C4.

The GAN has anastomoses with the auricular branch of the vagus nerve, the posterior auricular branch of the facial nerve, the lesser occipital nerve and the auriculotemporal nerve of the trigeminal nerve. The ansa cervicalis, also known as goose’s neck or ansa hypoglossi, is a loop of nerves formed by the ventral rami of C1 to C4 which join the hypoglossal nerve via a branch of C1 (Figure 2). It is also involved in the innervation of infrahyoid muscles, which are important for speech modulation and swallowing.

The case presented above illustrates this complex network of connections and suggests a dysfunction at a more proximal level than the peripheral branches. However, the mechanical disturbance and trigger at the neck, along with the presence of digestive symptoms prompt us to believe that the primary dysfunction is proximal to the peripheral cutaneous branches of the cervical plexus, most likely within the ansa cervicalis.

However, a central dysregulation of the brainstem trigeminal-autonomic circuits cannot be ruled out.

Contrary to cases of red ear syndrome described in the literature, this patient did not have another primary headache disorder, such as migraine. History and physical examination revealed pain of neuropathic origin as per the Douleur Neuropathique 4 questionnaire, but the patient suffered serious adverse reactions to usual neuropathic pharmacotherapy.

Our nerve blocks targeted the GAN peripherally in addition to its spinal nerve root origins at C2 and C3. The combination of these blocks provided more relief to the patient in terms of intensity, duration and frequency of paroxysmal earaches as well as her constant dull background pain, in comparison to blocking the GAN peripherally alone. The improvement was noticeable up to 3 months after the infiltrations. More long-term outcome is not yet known. Advantages to this technique versus fluoroscopic guidance included radiation-free imaging in a young female patient and ability to identify blood vessels in needle trajectory of a highly vascularized region, all in real time. Adverse events were not reported, but rare and serious complications could include injury to the vertebral artery, spinal radicular arteries, nerve roots and spinal cord direct or indirect trauma with leakage of anesthetic into the spinal canal causing motor and sensory blockade. Minor potential events could include local hematoma, bruising, soreness, transient pain exacerbation and vasovagal reactions. A technique aiming for the ansa cervicalis itself could be more successful, as we suspected this was the primary structure of dysfunction that could explain our patient’s constellation of symptoms. We are not aware of any such case reports and this kind of technique is not yet described. Furthermore, we were unable to

Figure 2 Cervical plexus.

to identify a cause or trigger to this dysfunction. However, a better understanding of the anatomical source guided our choices for better targeted nerve blocks that were able to provide our patient with symptomatic relief.

**Conclusion**
Great auricular neuralgia is an uncommon cause of facial pain that can be accompanied by other symptoms related to the cervical plexus. Cases refractory to usual pain medication could be treated by a combination of ultrasound-guided nerve blocks. Future potential therapeutic options would involve medial branch blocks of C2–C3, followed by radiofrequency ablation with or without peripheral neuromodulation of the GAN.

**Acknowledgments**
The authors wish to thank Dr Werner Becker, Dr Carrie Robertson and Dr Ivan Garza for their review of this report.

Written informed consent was obtained from the patient for publication of this case report and the accompanying images.

**Disclosure**
The authors report no conflicts of interest in this work.

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