A Bone-Breaking Case of Syncope Due to Glossopharyngeal Neuralgia

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Abstract

Glossopharyngeal neuralgia is an uncommon cause of intense facial pain, and is rarely associated with syncope. We present a patient with malignant syncope precipitated by glossopharyngeal neuralgia (GPN), who sustained lower extremity fractures. A 64-year-old Caucasian female with hypertension, diabetes mellitus type 2 and dyslipidemia, presented to the emergency department (ED) after a witnessed syncopal event. Over the preceding 9 months, she experienced multiple syncopal episodes triggered by neck pain, and was diagnosed with GPN by an ENT specialist. She reported a severe, stabbing pain in the right side of her neck prior to the event. Her vital signs and physical examination were unremarkable, except for bilateral leg tenderness. Laboratory data were within normal limits. Electrocardiogram revealed normal sinus rhythm, a rate of 72 beats per minute (BPM), and otherwise normal. Lower extremity X-rays revealed bilateral non-displaced fibular fractures. In the ED, she sustained another syncopal event, and bradycardia with transient heart rates as low as 22 BPM were documented. A temporary transvenous pacemaker was placed. She required pacing repeatedly for repeated bradycardic episodes preceded by right-sided neck pain. She was started on carbamazepine, without complete resolution of her symptoms on escalating doses. Due to recurrent bradycardia with frequent neck pains, requiring temporary pacing, an MRI-compatible permanent pacemaker was placed, to allow for further MRI recommended by neurology service. At 8-week follow-up, she had fully recovered from her neuralgic pain and fractures, without further syncope. This case emphasizes the potentially serious morbidity of GPN-induced syncope and importance of a prompt clinical diagnosis and treatment.

Keywords: Glossopharyngeal neuralgia; Syncope; Bradycardia; Pacemaker; Bone fracture

Introduction

Glossopharyngeal neuralgia (GPN) is an uncommon cause of intense facial pain, and is rarely associated with syncope. GPN-associated syncope has been attributed to a combination of vasodepressor and cardioinhibitory mechanisms. Such syncopal episodes tend to be debilitating and may result in injury, necessitating aggressive neuropathic pain management and, in refractory cases, insertion of a permanent pacemaker. We present a patient who sustained lower extremity fractures due to malignant syncope precipitated by GPN, necessitating the insertion of a permanent pacemaker. We review the pertinent literature with regard to the incidence, mechanism, triggers, reported complications and potential therapeutic options of this rare entity.

Case Report

A 64-year-old Caucasian female with hypertension, diabetes mellitus type 2 and dyslipidemia, presented to the emergency department (ED) after a witnessed syncopal event. Over the preceding 9 months, she experienced multiple syncopal episodes triggered by neck pain, and was diagnosed with GPN by an ENT specialist. She reported a severe, stabbing pain in the right side of her neck prior to the event. Her vital signs and physical examination were unremarkable, except for bilateral leg tenderness. Laboratory data were within normal limits. Electrocardiogram revealed normal sinus rhythm, a rate of 72 beats per minute (BPM), and otherwise normal. Lower extremity X-rays revealed bilateral non-displaced fibular fractures (Fig. 1). In the ED, she sustained another syncopal event; bradycardia with heart rates as low as 22 BPM were documented, and ECG performed promptly revealed sinus bradycardia (Fig. 2). A temporary transvenous pacemaker was placed. She required pacing repeatedly for repeated bradycardic episodes preceded by right-sided neck pain. The episodes could be precipitated by opening her mouth, talking and application of pressure on the right side of neck. Due to the severity of the symptoms, the patient often communicated by writing on a note pad instead of talk-
ing, and also declined examination of oral cavity by physicians. She was started on carbamazepine, without complete resolution of her symptoms. Due to recurrent bradycardia with frequent neck pains, requiring temporary pacing, an MRI-compatible permanent pacemaker was placed, to allow for further MRI recommended by neurology service. At the time of her discharge from the hospital, she had a modest improvement in the severity of her neuralgic pain on an escalated dose of carbamazepine with reduction in frequency of episodes. At 8-week follow-up, she had fully recovered from her neuralgic pain and fractures, without further syncope.

Discussion

GPN causes paroxysmal attacks of brief stabbing pain, localized to the external ear canal, the base of the tongue, the tonsil or the area beneath the angle of the jaw, with triggers such as speaking, swallowing, eating, breathing, cold air and slight touch of the mouth and pharyngeal region [1]. Although it is mostly idiopathic without major pathology, and often caused by compression of glossopharyngeal nerve as it exits or enters the brainstem, secondary causes have been described including trauma, neoplasm, infection or vascular malformations [2]. GPN is rare in the general population, with an incidence rate of approximately 0.7 per 100,000 [3].

Syncope has been reported in four out of 216 GPN cases (approximately 2%) of a Mayo case series [4]. The syncope appears to be hemodynamically mediated, with a drop in blood pressure and heart rate coinciding with the onset of pain, as demonstrated by Kim et al in a 53-year-old man admitted with recurrent syncope associated with pain from GPN [5]. One plausible mechanism explaining the association between bradycardia, hypotension and syncope, and GPN is the close connection between the vagus nerve and glossopharyngeal nerve with respect to visceral sensations, creating a vagoglossopharyngeal reflex arch [6]. Barbash et al [7] examined the mechanism of syncope in a patient with GPN. They reported that while atropine prevented asystole (without relieving pain), blood pressure continued to decrease due to inhibition of the sympathetic tone, with decrease in the serum norepinephrine levels during the attacks. Weinstein et al [8] also reported a case of GPN due to metastatic head and neck tumor with hypotension but without

Figure 1. Bilateral foot X-rays reveal fibular fractures (white arrows).
bradycardia. Atropine resulted in sinus tachycardia but did not abolish the hypotension, suggesting selective stimulation of baroreceptor vasodepressor activity.

However, pain may not always be the trigger for the syncope seen in GPN. Korkes et al [9] reported a 71-year-old male with sporadic syncopepisodes, who complained only of ill-characterized spasms in the throat. A loop monitor revealed 4.4 s sinus arrest with presyncope while eating; his symptoms resolved after placement of a permanent pacemaker. Reddy et al [10] reported another case of syncope necessitating a permanent pacemaker insertion in a patient with GPN, but without pain in the distribution of the glossopharyngeal nerve.

The consequences of GPN-induced syncope vary. Our patient sustained bilateral fibular fractures due to an episode of GPN-induced malignant syncope. She also had stopped communicating verbally due to induction of pain while talking, and these symptoms resolved after placement of a permanent pacemaker. Bodily injury can also be a complication of GPN-induced syncope, as in our patient. Elias et al [13] reported an 80-year-old white male with GPN who sustained head trauma and right maxillary sinus fracture following a syncopal episode triggered by lancinating pain in the pharynx. Seizures have also been reported in patients with GPN-induced syncope, likely due to the transient hemodynamic compromise and decrease in cerebrovascular perfusion [14-16].

Therapy for GPN-induced syncope varies among reports. In some reports [17, 18] treatment with carbamazepine abolished the pain and associated cardiovascular manifestations. However, in other cases, treatment with carbamazepine alone [19, 20], carbamazepine and gabapentin combination [21] or clonazepam [22] was not sufficient to alleviate the pain and syncope episodes in GPN, necessitating the insertion of a permanent pacemaker to prevent syncope. Other successful treatments of refractory cases included section of the ninth (glossopharyngeal) nerve alone [23, 24], or combined with rhizotomy and microvascular decompression of the lower cranial nerves [25].

Our patient’s syncope resulted in severe bodily injury causing lower extremity fractures. MRI of the head and neck was recommended by neurology service. However, since the presence of temporary myocardial pacing leads is considered a safety contraindication for MRI [26], it was deemed unsafe for her to undergo MRI testing due to her dependence on temporary pacing. Therefore, the permanent pacemaker brand inserted was REVO (Medtronic), which is compatible with 1.5-Tesla MRI scanners [27]. She later was discharged in good condition, and was scheduled to undergo an outpatient MRI for further evaluation.

Conclusion

In conclusion, GPN is a rare, but serious cause of cardiac
syncope, which is associated with a well-defined presentation. It is paroxysmal, but episodes can be frequent enough to result in bodily injury. Treatment may address pharmacologic or surgical pain relief, versus addition of a pacemaker in refractory cases to prevent episodic asystole. Insertion of a novel MRI-safe pacemaker allows for safe future use of MRI for further workup when indicated.

References


