Case Report

Vascular entrapment neuropathy of the tibial nerve within the gastrocnemius muscle

Paul Samuel Page, Stewart Paige, Amgad Hanna

Department of Neurosurgery, University of Wisconsin Hospitals and Clinics, Madison, Wisconsin, United States.

E-mail: Paul Samuel Page - paulsamuelpage@gmail.com; Stewart Paige - tpstewart@wisc.edu; *Amgad Hanna - ah2904@yahoo.com

INTRODUCTION

Tibial neuropathy is a common clinical entity that has been described as a result of compression. The most common location occurs under the soleal arch or in the tarsal tunnel. In addition to these sites, other sites of compression have been identified as well as localized injury and compression of individual branches. While most causes of tibial neuropathy remain idiopathic, other causes include trauma, mechanical irritation, and tumors.

More recently, ultrasonographic literature describes the “punched-nerve-syndrome” as a rare condition where direct vicinity to a vascular structure leads to chronic neural damage due to repetitive pulsatile compression.[5] This pathology has been described in the setting of ulnar neuropathy as well as radial neuropathy; however has not been described affecting the tibial nerve thus far.[1,3] Here, we describe the first case of a “punched-nerve-syndrome” affecting the medial gastrocnemius (MG) branch from the tibial nerve. Given this rare entity, we hope to increase the awareness of this syndrome.

CASE REPORT

A 41-year-old male presented to clinic with a 2.5-year history of painful right MG atrophy and fasciculations following a previous gastrocnemius tear after a fall from a treadmill. He was initially
seen by neurology where his right MG appeared significantly wasted compared to his right lateral gastrocnemius. On examination, fasciculations were present and limited to his right gastrocnemius with no reliable inciting triggers. On motor examination, his strength was MRC Grade 5/5 in all muscle groups, including plantar flexion and inversions, and deep tendon reflexes were 2+ and symmetric. Further workup with electromyography (EMG) revealed a possible focal neuropathy of the right tibial nerve branch supplying the MG, as evidenced by increased insertional activity, 3+ fasciculations, and 1+ spike and wave activity in the MG. All other EMGs on muscle group testing were normal. There was no evidence of radiculopathy, plexopathy, or proximal sciatic neuropathy. Subsequent MRI illustrated his tibial and common peroneal nerve had normal morphology without any structural evidence or injury to explain his EMG findings. Ultimately, the decision was made for a right tibial nerve exploration at the level of the popliteal fossa with decompression and possible grafting of the MG branch.

On exploration, both the medial and lateral gastrocnemius muscles were easily identified along with the sciatic nerve and associated tibial and common fibular nerve branch points. Of interest, there were three distinct muscle head contributions to the MG. It was unclear, however, whether this anomaly was congenital or secondary to his previous injury. Nerve identifications were confirmed with intraoperative checkpoint nerve stimulator under direct visualization. Three MG branches were identified arising from the tibial nerve. Unexpectedly, we found large vasculature including, 1 artery and 2 veins, directly abutting deep to the MG branches [Figure 1]. Vascular surgery was consulted intraoperatively, and ligation of these vessels was performed to completely decompress the nerve. After ligation and full dissection of the MG branches, the muscle fasciculations and ongoing twitching had diminished. In addition, EMG stimulation was completed before closure to ensure no injury to the dissected nerves had occurred.

On follow-up at 3 months, his incision had healed without complication, he regained more strength, and his muscle fasciculations had decreased.

DISCUSSION

Mononeuropathies are nerve dysfunctions commonly secondary to local injury such as impingement, mechanical impact, tearing, and/or other constant pressure sources. Tibial neuropathies, in particular, are typically heavily attributed to both impingement and/or injury such as with tarsal tunnel syndrome and entrapment deep to the flexor retinaculum.[2,4] This same phenomenon could be caused by any other source of impingement such as space-occupying lesions, compartment syndromes, or other morphologic variants.[2,7] For instance, in a 2013 study by Tang et al., they evaluated the effects of acute compartment syndrome and subsequent proximal tibial nerve compression by the soleus tendinous arch.[7] However, vascular compression is not typically high on the differential diagnosis. Unlike the trigeminal and facial nerves where vascular compression is a common and well-studied phenomenon, the same is not true with the tibial nerve.[2,6-8] It seems logical that neuropathy secondary to close vascular proximity could occur at various locations in the body, especially given the tendency of nerves to travel adjacent to arterial and venous structures within the same space.

In the presented case, given the significant improvement in fasciculations following the vasculature ligation, the most likely diagnosis following exploration was due to vascular compression of his right MG branches. Contrary to cranial nerves, his presentation was unusual considering the ample space in the popliteal fossa and planes between gastrocnemius heads and underlying soleus muscle. In addition, the normal anatomic variant includes only one MG head and its nerve supply. Yet, our patient not only had multiple MG heads but
also a total of three small nerve contributions branching off the tibial nerve. Therefore, there may have been more surface area of nerve to cause abutment and compression to. His previously torn gastrocnemius was likely to cause decreased space between adjacent structures in accommodation for increased swelling as well as adhesions. While the etiology of his anatomic variation remains unclear, it remains feasible that the combination of his anatomy in conjunction with his injury provide a rational mechanism of vascular compression of his MG branch.

Overall, while punched nerve syndrome has been described in the setting of vascular compression of peripheral nerves, this syndrome has largely been described in the setting of ultrasonography. This syndrome was first described in 2012 by Loizides et al. in nine cases of isolated mononeuropathy.[5] On real-time high-resolution ultrasound (HRUS), they were able to identify the involved nerve segments with enlarged hypoechoic changes of echotexture at the site of vascular compression. In addition, two case reports exist in the literature these HRUS findings correlating with mononeuropathies of the median and ulnar nerves which improved following surgical decompression.[1,3] In our case, while no HRUS findings were obtained preoperatively, our findings suggest that vascular compression, and thus punched nerve syndrome, likely provided a significant contribution to our patients mononeuropathy.

CONCLUSION
A high level of suspicion is required to identify vascular compression in the setting of a mononeuropathy. With an understanding of the uncommon presentation and location within the popliteal fossa, clinicians should consider the possibility of vascular compression in individuals with a history of localized trauma and lasting associated mononeuropathy. Surgical exploration is worthwhile when the clinical context and EMG findings are compelling, even when the imaging is negative.

Declaration of patient consent
The authors certify that they have obtained all appropriate patient consent.

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Conflicts of interest
There are no conflicts of interest.

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