Trigeminal neuralgia secondary to vascular compression and neurocysticercosis: illustrative case

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BACKGROUND Trigeminal neuralgia (TN) is a frequent neurosurgical problem negatively influencing the quality of life of patients. The standard surgical treatment is microvascular decompression for primary cases and decompression of the mass effect, mainly tumors, for secondary cases. Neurocysticercosis (NCC) in the cerebellopontine angle is a rare etiology of TN. The authors report a case in which NCC cysts around the trigeminal nerve coexisted with a vascular loop, which compressed the exit of the trigeminal nerve from the pons.

OBSERVATIONS A 78-year-old woman presented with a 3-year history of persistent severe pain in the left side of her face, refractory to medical treatment. On gadolinium-enhanced magnetic resonance imaging, cystic lesions were observed around the left trigeminal nerve and a vascular loop was also present and in contact with the nerve. A retrosigmoid approach for cyst excision plus microvascular decompression of the trigeminal nerve was successfully performed. There were no complications. The patient was discharged without facial pain.

LESSONS Albeit rare, TN secondary to NCC cysts should be considered in the differential diagnosis in NCC-endemic regions. In this case, the cause of the neuralgia was probably both problems, because when both were treated, the patient improved.

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KEYWORDS trigeminal neuralgia; microvascular compression; neurocysticercosis; retrosigmoid approach; Peru

Trigeminal neuralgia (TN) is the most common type of facial pain, negatively affecting quality of life and work capacity in 34% of patients.1 TN has a prevalence of 4–13 cases per 100,000 inhabitants1–3 and generally affects patients older than 50 years, with a female-to-male ratio of 1.5 to 1. In primary (classic) TN, compression by vascular loops is found at the entrance of the trigeminal nerve to the brainstem.4,5 In secondary TN,6,7 extrinsic compression of the trigeminal nerve triggers the pain. The most common causes of extrinsic TN are tumors (meningiomas, epidermoid cysts, acoustic neuromas, etc.) and, less frequently, aneurysms and arteriovenous malformations,8 and rarer still, neurocysticercosis (NCC). In secondary TN, pathophysiologic changes similar to those in primary TN occur, although the structural lesion depends on the etiology; for example, in multiple sclerosis, it is due to demyelination plaques.9 The standard treatment for TN is microsurgical decompression of the trigeminal nerve. We present a case in which neighboring NCC cysts, arachnoiditis, and vascular compression occurred in a patient with TN, who was treated with excision of the cysts plus microvascular decompression of the trigeminal nerve.

Illustrative Case

History and Examination
A 78-year-old woman presented with intense pain and numbness in the left side of her face that felt like electric discharges and that started especially by chewing food, which prevented her from eating. Her only noticeable medical antecedent was arterial hypertension, which was under regular medical treatment. She was initially treated by the neurology department with anti-inflammatory drugs, analgesics, and carbamazepine (200 mg 3 times/day). As the pain was not controlled, she was referred for neurosurgical evaluation. On examination, there was a left V1–V2 facial hypoesthesia and

ABBREVIATIONS AICA = anterior inferior cerebellar artery; CPA = cerebellopontine angle; MRI = magnetic resonance imaging; MS = multiple sclerosis; NCC = neurocysticercosis; TN = trigeminal neuralgia.

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left sensorineural hearing loss and no other neurological sign. Gadolinium-enhanced magnetic resonance imaging (MRI) revealed cystic lesions in the left cerebellopontine angle (CPA), better seen on the fast imaging employing steady-state acquisition (FIESTA) sequence (Fig. 1). In addition, an elongated trigeminal nerve was observed trapped between two cystic lesions. The fluid-attenuated inversion recovery (FLAIR) sequence revealed pial edema around the pons, reflecting the inflammatory-immunological reaction of the parenchyma in response to the cysts. Toward the emergence of the trigeminal nerve, a branch of the anterior inferior cerebellar artery (AICA) was also seen in contact with the nerve. Immunodiagnosis using enzyme-linked immunoelectrotransfer blot (Western blot; sensitivity of 98% and specificity of 100% in patients with two or more viable brain lesions) and detection of circulating Taenia solium were both strongly positive. After evaluating the case, it was decided to use a retrosigmoid approach for the resection of the NCC cysts plus microvascular decompression of the left trigeminal nerve.

Surgery

The surgical plan was, first, resection of the cysts and, later, exploration and possible decompression of the trigeminal nerve. The first surgery used a microsurgical retrosigmoid approach, with the patient in the right lateral decubitus park-bench position with a 10-cm linear retroauricular incision at 4 cm from the ear posterior implant. Craniotomy was 4 cm in diameter, and the dura mater was opened, referencing the sigmoid sinus. Initially, the cerebrospinal fluid was drained to expand our work space. The cranial nerves VII–VIII complex was initially identified, and then the entire arachnoid of the CPA was observed with the thick, yellowish appearance typical of arachnoiditis secondary to NCC. After opening the arachnoids, the cystic lesions were visualized in the deeper plane (Fig. 2), again typical of NCC. NCC cysts are generally not attached to vascular structures or brain parenchyma, so saline irrigation and gentle traction were sufficient to remove the cysts. The cysts were extracted and the exploration was expanded. A branch of the AICA was found to be in contact with the base of the trigeminal nerve (Fig. 3). The artery was dissected from the arachnoid adhesions and separated from the nerve with a piece of Teflon. After hemostasis was verified, closure was performed in layers with the usual technique.

There were no complications during the procedure. The patient was extubated without added neurological signs and with relief from facial pain and was discharged 3 days later. Pathological examination of the excised membranes confirmed the diagnosis of NCC. Follow-up MRI at 2 months after surgery found no evidence of NCC cysts in the CPA (Fig. 4). She received antiparasitic treatment with albendazole, with concomitant steroids, for 2 months. Currently, the patient is under follow-up with a modified Rankin scale score of 0, without facial pain.

Discussion

Observations

TN is a frequent and well-known neurosurgical problem significantly affecting the quality of life of patients. Its global incidence has remained constant, ranging from 12.6 to 27/100,000 inhabitants per year, which increases in people over 50 years and is slightly more frequent in women and on the right side.8,9 Bilaterality can be seen in 5% of classic cases.10,11 Undoubtedly, vascular compression is the most frequent cause. Of the secondary etiologies, multiple sclerosis (MS) is the most frequent cause of TN, with a prevalence ranging from 1% to 6.3%,12 followed by intracranial tumors. Some cases secondary to NCC cysts or cystercercal arachnoiditis have also been reported. This immunological-inflammatory reaction can also generate irritation of other cranial nerves, such as the reported case of hemifacial spasm secondary to arachnoiditis due to NCC, which improved after release of the facial nerve.13

The diagnosis of TN remains fundamentally clinical.3,6,14 The first-line treatment for TN continues to be anticonvulsant medical therapy, and those cases secondary to tumors or infectious inflammatory problems are treated initially with surgery.5,7 In patients with MS, the prevalence is higher, and in addition to episodic pain, they often report a component of constant pain.10,12,15 Although the pain is mainly unilateral in these patients, bilateral involvement can occur in up to 31% of TN cases secondary to MS.16

TN may also occur in patients with tumoral and/or infectious lesions of the CPA. While meningiomas (classically in the petrous apex) are the most frequently reported lesions associated with TN, small lesions in very specific locations can also cause it, as in the case reported by Ishi et al.17 in 2015, in which a very small meningioma in the suprameatal tubercle, with hyperostosis at the entrance to Meckel’s cave, led to intractable TN in a 72-year-old woman. Other frequent extra-axial tumors that may cause TN are acoustic

FIG. 1. Preoperative axial MRI. A and B: FIESTA sequences showing subarachnoid cystic lesions in the left cerebellopontine angle (yellow arrows) medial and lateral to the trigeminal nerve, which is elongated laterally. C: FLAIR sequence demonstrating hyperintensity (white arrow) in the pial plane of the pons around cysts. D: Gadolinium-enhanced, T1-weighted sequence showing a branch of the AICA at the exit of the trigeminal nerve from (red arrows).
Rarer etiologies for TN have also been reported, such as in the patient who presented with TN as the only presenting symptom secondary to brain metastasis or in association with Charcot-Marie-Tooth disease and bilateral TN, with only 4 reported cases.

There are only a few reports of TN secondary to NCC. Tenuto et al. reported two cases secondary to NCC cysts in Meckel's cave; one of the cases initially evaluated as clinical suspicion and the other as an intraoperative finding. Revuelta et al. presented two cases—interestingly, one of them with contralateral neuralgia. This patient had NCC cysts in the left CPA with right-sided TN, and the pain completely subsided after removing the left cerebellopontine cysts. They suggested that the neuralgia was possibly caused by distortion of the brainstem resulting in compression of the nerve against an arterial loop at its entrance site. Their other case of NCC of the CPA with ipsilateral neuralgia was also alleviated after excision of the cysts. Hamamoto Filho et al. reported another case secondary to ipsilateral NCC cysts in the CPA, in the absence of vascular compression. Aguiar et al. reported another singular case—that is, a patient with bilateral TN with images of cystic lesions in the right CPA, which was finally reported as NCC, who also presented with neuralgia on the left side. The neuralgia improved on both sides after excision of the right CPA cyst. They considered that the neuralgia was caused by two probable mechanisms: a distortion of the brainstem and compression of the nerve against an arterial loop in the entry zone or arachnoiditis caused by NCC in both cisterns of the CPA. It is known that the inflammatory-immunological reaction produced by NCC cysts causes vascular damage in the form of vasculitis and nerve damage in the form of neuritis. The treatment does not conclude with the excision of associated cysts and may involve antiparasitic or anti-inflammatory regimens. Unlike the cases mentioned, in our case we observed both problems—that is, cysts and vascular compression.

Lessons
NCC should be considered as a diagnosis in patients with cystic lesions in the CNS. In our case, in addition to the local inflammatory reaction, vascular compression could be observed, suggesting that both mechanisms contributed to the neuralgia. However, the exact
role of each problem in the pathophysiology of neuralgia remains for analysis and discussion. Surgical resolution of both conditions resolved the patient's facial pain.

References
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