Intradiploic epidermoid cyst with intracranial hypertension syndrome: Report of two cases and literature review

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A B S T R A C T

INTRODUCTION: Intradiploic epidermoid intracranial cysts (IEIC) derive from ectodermal cells and are covered with stratified squamous epithelium. They are extremely rare, and most common locations are in the occipital, frontal and parietal bones. They have a very slow growth and can be asymptomatic until becoming evident by the deformation produced. The treatment is based on the removal of the lesion, and subsequent histopathological confirmation.

PRESENTATION OF CASE: Two cases are reported, with intracranial hypertension syndrome, which is very uncommon because of the slow growth of this type of pathology; however, decompensations occurring in the space-occupying lesions at intracranial level explain this type of clinical presentation.

DISCUSSION: The most common presentation of intracranial intradiploic epidermoid cysts (IEIC) is asymptomatic, which is made evident by the prominence at the level of the soft tissues and then presenting less frequently local pain and cephalgia; rarely the size of the lesion can cause focal neurological signs.

CONCLUSION: These benign lesions, although they are of low incidence, are seen very rarely in intradiploic locations and above all, of significant size, may produce significant mass effect in patients, which was initially tolerated because of its slow growth, however, they may become decompensate and cause intracranial hypertension syndrome.

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1. Introduction

Epidermoid cysts are benign tumors that originate in the spine or intracranially [1]. Love and Kernohan in 1936, referred to epidermoid cysts as congenital epithelial tumors [2]. They originate from granulation alterations producing a defect in the closure of the neural tube during the third to sixth week of gestation, at level surface of the ectoderm [3].

At intracranial level represents 0.2–1.8% of all tumor lesions; they can be located intra- or extradurally [4,5]. Intradural compose 75% of epidermoid cysts and -IEIC-25% are extradural in the diploidal space—IEIC [6,7].

IEICs are very rare; as the name indicates are found among the two tables of the skull bones, having a slow growth [8,9]. The first IEIC was reported by Müller in 1838, Cushing in 1922, also performed a description of an IEIC [12,10] and the first case of giant IEIC in the posterior cranial fossa was reported by Rengachary [11].

Two cases are reported, that in a particular way, go to the emergency room because they display intracranial hypertension syndrome, which is very intriguing because of the slow growth of this type of pathology; however, decompensation occurring in the pathophysiology of space-occupying lesions at intracranial level explain this type of clinical presentation.

2. Presentation of cases

2.1. Case 1

Male, 42, with a history of smoking and drug use—methamphetamines. Presents occipital headache with 3 months evolution that radiates to the frontal region, pulsatile, which increases with Valsalva maneuvers with a frequency initially described as sporadic until evolving to daily, in the week prior to admission; accompanied by vomiting, on the day of admission, with intensified headache intensity. On physical examination, the patient exhibits intracranial hypertension symptoms (papilledema) without rostrocaudal deterioration, which is accompanied by indifferent lateropulsion; in imaging studies, extra-axial (Fig. 1) lesion is observed. Study protocol is completed, and resection of the lesion is performed by means of a right lateral suboccipital approach. Among the findings an important erosion of
the occipital squama was observed due to extra-axial injury, eroding the bone, avascular, not infiltrating with surrounding tissues, of pearly aspect with fatty content inside (Fig. 2). Histopathological studies, reports us soft lesion with granular areas and firm membranous areas with calcifications, reporting an epidermal cyst. The patient progressed favorably, without neurological deficits with one-year follow-up (Fig. 3).

2.2. Case 2

Male 46 years old, without major importance history. Since 2 months ago with palpebral edema, which lasted four days, and was treated at another health facility as cellulitis, with antibiotics and anti-inflammatory therapy, resolving favorably; due to increasing deformity of the right frontal region, consults with private doctor who diagnosed intracranial lesion with respective imaging studies and subsequently refers to our hospital, by the mass effect is admitted (Fig. 4); at the time of admission the patient develops severe holocranial cephalgia, accompanied by vomiting, papilledema and stupor, therefore is decided to take to the OR suite for resection of the lesion, through a centered approach; during surgery important right frontal bone erosion was
Fig. 3. Images of post-surgery studies tomography (a and b) and MRI (c–g), where we can observe the absence of lesion and how the structures compressed by it, return to their place.

Fig. 4. Simple skull tomography images, where lesion with important extra-axial intracranial mass effect is observed, with significant bone erosion with heterogeneous content.
Fig. 5. Intraoperative photos, where the lesion characteristic and the significant erosion of the right frontal bone is observed.

Fig. 6. Postoperative tomography images where the resolution of the mass effect that produced the lesion (a, b) is observed; also the bone defect that was observed (c) and subsequent cranioplasty images with methyl methacrylate (d, e) are observed.
observed due to extra-axial injury, not vascularized, not infiltrating with neighboring tissues, with fatty content inside (Fig. 5); histopathological result is epidermoid cyst; during the immediate post-surgery period, the patient develops mixed delirium, which is solved in three days; is discharged without any complications; cranioplasty is later performed at 3 months after resection surgery of intradiploic epidermoid cyst (Fig. 6).

3. Discussion

Intracranial epidermoid cysts are generally located in the cerebellopontine angle or parasellar region [13]. Extradural lesions manifest with local mass effect, with or without cephalae [14]. The incidence of intracranial epidermoid cysts is higher in the intradural location, as reported in their work Love and Kernohan—70% and Gormley et al.—72.7% [2,14].

Akar et al. reported 28 cases of intradural epidermoid cysts, of which the majority are found in the cerebellopontine angle—17; other locations were in the fourth ventricle, into the cisterna magna, in the fissure of Silvio, in the occipital lobe, in the lateral ventricles and diploic space [4]. The intradiploic extradural epidermoid cysts are located in the cranial vault, destroy the internal table and thin the external table [15]. The most common presentation of IEIC is the prominence at the level of the soft tissues and then presenting less frequently local pain and cephalae; rarely the size of the lesion can cause focal neurological signs [16,17].

They can be located in the paranasal sinuses, in the temporal and sphenoid bone, however it occurs most frequently at the level of the occipital, frontal and parietal bones [18].

The differential diagnosis of an intradiploic cranial lesion includes the aneurysmal bone cyst, dermoid cyst, cavernous hemangioma, Langerhans cells histiocytosis, fibrous dysplasia and eosinophilic granuloma [16,19]. Atypical lesions may be larger than 5 cm and irregular [1].

Histologically, it is covered by fibrous tissue capsule, in which are included keratin, cholesterol, proteins and detritus crystals [20]. Malignant transformation of epidermoid cysts is very rare; the possibility of squamous cell carcinoma originated from a IEIC is described with poor prognosis even after radio or chemotherapy [21,22].

Resection of these lesions are done to protect the patient from mass effect on intracranial mass, abscess formation and to avoid potential complications such as bleeding and malignant transformation of IEIC [1,23,24].

4. Conclusion

These benign lesions although they are of low incidence, are seen very rarely in intradiploic locations and above all, of significant size, may produce significant mass effect in patients, which was initially tolerated because of its slow growth, however, they may become decompensate and cause intracranial hypertension syndrome.

Dates of congress

This paper has not been presented in any congress.

Conflict of interest

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Ethical approval

The name of the ethics committee is “collegial session neurosurgery”, with number 213892 for case 1 and 214450 for case 2, of the Neurology and Neurosurgery National Institute.

Consent

Written informed consent was obtained from the patient for publication of this case report and accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal on request.

Author contribution

Moreira-Holguín Juan Carlos: writing the paper. Medélez-Borbonio Rafael: data collection. Quintero-López Eduardo: traducing. García-González Ulises: data collection. Gómez-Amador Juan Luís: concept of paper and interpretation.

Guarantor

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