A Unique Case Of Ecchordosis Physaliphora With Intratumoral Haemorrhage

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INTRODUCTION

EP is a rare intracranial extra-axial mass derived from notochordal tissue that is typically located intradurally in the pre-pontine cistern and attached to the dorsal wall of clivus (1). Fetal notochordal tissues play a major role in the genesis of the axial skeleton and persist in the adult at the nucleus pulposus of the intervertebral disc (6). Due to a common origin from the fetal notochordal rests, EP and chordomas share common histological and ultrastructural characteristics (3, 7-11). Ecchordosis physaliphora is an extremely rare lesion. It is found incidentally in as few as 0.5-2% of autopsies (3, 8, 10, 11). The natural pathogenesis of the disease is characterized by slow subclinical progression with only rare ostensible manifestations. Symptoms, if occur, are the direct result of involvement of surrounding neurovascular structures and depend on the location of primary mass. EP should be differentiated from retroclival chordoma as accurate identification of nature of these masses has relevance in the determination of patient prognosis and in the planning of therapy.

CASE REPORT

A 31 yr old otherwise healthy woman presented to us with frequent headaches but no associated fever or vomiting. There was no history of blurring of vision, rhinorrhoea or ear discharge. Physical examination findings were normal and there were no neurologic deficits. Magnetic resonance (MR) images of the brain and computed tomographic (CT) scans of the head were obtained.

Figure 1

Initial MR imaging of the brain revealed an intradural, well circumscribed, extra-axial retroclival mass located in the prepotine cistern. The mass appeared hyperintense on T2 and T1-weighted images, which likely represented haemorrhage, and restricted diffusion on diffusion-weighted images (Fig 1A, B, C). Contrast enhanced T1 weighted MR images were obtained and showed no enhancement (Fig 2A). On the T2-weighted image, the homogenously hyperintense signal intensity may have been less due to pulsation artifact in the lesion than in the pre-pontine CSF (Fig 2B). A possible diagnosis of Ecchordosis physaliphora was considered and sagittal CT scan with bone window settings was obtained. An osseous stalk arising from the clivus was well appreciated on sagittal CT scan with bone window settings (Fig 5). No bone destruction was noted.
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**DISCUSSION**

EP is a rare, congenital, benign hamartomatous lesion arising from an ectopic notochordal remnant. This tissue is located in the midline of the craniospinal axis, reaching from the dorsum sellae to the sacrococcygeal region (1-4). The notochord is the primitive skeleton of vertebrates that forms the nucleus pulposus of the intervertebral disk at maturity. Ectopic notochordal rests can be seen outside the nucleus pulposus anywhere along the axial skeleton (1, 12).

Intracranial EP is typically found intradurally in the prepontine cistern, where it is attached to the dorsal wall of the clivus via a small pedicle (1, 2, 5). Recognition of the imaging features of EP is helpful in suggesting the diagnosis and differentiating from other retroclival lesions. CT is limited in detecting the EP due to its small size and beam hardening artefacts in the posterior fossa (20). However, the osseous stalk at the dorsal wall of the clivus on thin-section CT images is considered as the morphological hallmark of EP and does not occur in other retroclival lesions (5). MR is the best imaging modality for the radiological detection of EP, and the characteristic appearance of EP in MR is that of a well circumscribed, round, extra-axial mass in the prepontine cistern. EP demonstrated high signal intensity on T2-weighted images, low signal intensity on T1-weighted images, and no contrast enhancement.

The differential diagnosis of retroclival intradural lesions in the prepontine cistern includes chordoma, dermoid and epidermoid cysts, arachnoid cyst and Ecchordosis physaliphora (1, 13). A diagnosis of chordoma is less likely in our patient as these tumours usually produce symptoms that suggest brainstem involvement or cranial nerve palsies, that show bone destruction, that enhance with contrast material administration (1, 13, 14). On MR images, dermoid and epidermoid cysts may have similar signal intensity to that of EP without contrast enhancement (15). However our lesion didn’t suppress on fat suppressed sequences and restricted diffusion on diffusion-weighted images. The key imaging feature in this patient was the osseous stalk arising from the dorsal clivus. The absence of symptoms apart from headaches, midline location along the craniospinal axis, lack of contrast enhancement, hyperintense on both T1 and T2-weighted images with restricted diffusion on diffusion-weighted imaging, with no obvious bone destruction also favoured EP with intratumoral haemorrhage as the most likely diagnosis in this patient (5).

EP is a midline intradural lesion that shows no contrast enhancement, likely because of the paucity of vascular structures (1, 5, 17). In contrast, chordomas usually are seen in the extraosseous intradural portion of the clivus, causing local bone destruction (13, 16 radio). A purely intradural chordoma is rare and requires differentiation from EP (1, 13, 16-radio). Both the typical extradural chordoma and rare intradural chordoma show contrast enhancement, which is key feature in differentiating them from EP (1, 5, 13, 14, 16, 18, 19).

In conclusion, EP is a congenital malformation that needs to be considered in the differential diagnosis of retroclival lesions. In rare cases, it may be haemorrhagic or may be symptomatic due to tumor compression of the brain stem. When a retroclival mass is established on MRI, it should be differentiated from other retroclival lesions by using imaging features, especially an osseous stalk, as this is relevant in the patient prognosis and planning of therapy.

**References**

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