Intracranial dermoid cysts are rare dysembryonic tumors of benign nature. These are uncommon in adults. If present, they are usually located in the midline or along the lines of embryonic fusion. The posterior fossa region is an infrequent site. Extrudal or interdural locations are even more rare. In this case report, the authors report a laterally located large posterior fossa right cerebellar convexity interdural and extradural dermoid cyst over the sigmoid sinus. It was managed by totally extradural maximum possible safe decompression with microneurosurgical technique. The authors share their experience of addressing this rare pathology at the rarest location with unusual imaging findings.

Introduction

Intracranial dermoid cysts (DCs) are rare dysembryonic tumors of benign nature, accounting for 0.04 to 0.7% of all intracranial space-occupying lesions. Because of their congenital origin, they are mostly seen in infants and young adults, and comparatively rarely in older age groups. They have a predilection for midline or lines of embryonic fusion. The common locations are cisternal spaces, mainly in the cerebellopontine angle and the parasellar cisterns. DCs of the suboccipital and occipital regions are uncommon. If present, they are remarkably in midline within the vermis or in the fourth ventricle. Laterally located intracerebellar dermoids are also reported. DCs of interdural space or extradural space are infrequent and only few cases have been reported. Herein, the authors present an interesting case of a large cerebellar convexity DC with both interdural and extradural components, compromising venous drainage of the ipsilateral transverse and sigmoid sinus. The patient was managed successfully by microneurosurgical decompression.
Case Report

A 34-year-old lady presented to our department with a 4-month history of headache with on and off vomiting. On clinical examination, she had left cerebellar signs. Her magnetic resonance imaging (MRI) revealed a 35 × 48 × 46 mm broad dural-based extra-axial right cerebellar mass lesion. It had mixed signal intensity on T1 (►Fig. 1) and was hyperintense on T2 (►Fig. 2). There were areas of peripheral enhancement on contrast administration (►Fig. 3A–C) with diffusion restriction on apparent diffusion-weighted (ADW) imaging (►Fig. 4). There was no obvious surrounding edema.

The right transverse sinus was not appreciable, the fourth ventricle was chinked with mild upstream hydrocephalus, and there was cerebellar midline shift to the left. Magnetic resonance venogram (MRV) showed nonvisualization of the right transverse and sigmoid sinus (►Fig. 5).

Right paramedian suboccipital craniotomy was contemplated under general anesthesia. Postcraniotomy grayish-white adipose tissues with desquamated debris and hairs were found in the extradural space extending interdurally between the leaves of the tentorium cerebelli (►Figs. 6 and 7). Maximum possible safe decompression without injuring the dura mater was done (►Fig. 8). Histopathology
was consistent with DC. Postoperative MRI of the cranium had minimal residual disease adhered at the dural surface of the sinus (Fig. 9). The postoperative course was uneventful.

Discussion

The history of intracranial dermoid and epidermoid cysts dates back to 1829 when Cruveilhier coined the term “tumeur perles” for them. These tumors possess both ectodermal and mesodermal components. They are supposed to originate from inclusion of the ectodermal elements within the neural tube during its closure between the third and fifth week of embryonic development. This explains their preponderance in midline, in the diploe of the fontanel extradurally, and in the parasellar region intradurally. Traumatic implantation has also been an alternate hypothesis in the genesis of dermoids. Lateral extradural dermoids have been postulated to develop from dural multipotent...
embryonic cells or from translocation of the epithelial cells of developing neovasculature. These rare tumors account for 0.04 to 0.7% of all intracranial lesions. These are mostly intradural lesions located at the frontobasal, suprasellar, parasellar, etc., regions. Extradural DCs are much rarer and have been reported in the midline of the posterior fossa, anterior fontanel, and orbital region in children. Interdural DC in the lateral wall of cavernous sinus has also been cited. But DC in the region of cerebellar convexity at an unusual lateral location with both interdural and extradural components holds its paucity. Esaki et al have described an atomically similar case. These are commonly diagnosed in infants to young adults owing to their congenital origin and association with sinuses and sinus tract infections. Thus, they are relatively rare in the middle-aged or elderly population.

There is a wide array of presenting symptoms of these slow-growing tumors, depending upon the size and location, namely headache, dizziness, seizure, blurring of vision, progressive hearing loss, facial deviation, lower cranial nerve palsy, etc. Rupture may lead to severe aseptic meningitis, ventriculitis, and death.

MRI of the cranium is the investigation of choice. Dermoids show variable relaxation time depending upon fat content. Typically, these are hyperintense on T1- and T2-weighted images due to partially liquified fat content. However, low-/mixed-intensity T1-weighted images can also be found, as in the indexed case, because of solid crystalline cholesterol and keratin. Usually, these tumors do not enhance on contrast administration, but there was peripheral enhancement in our case. It can be attributed to peripheral granulation with or without previous infection. These show diffusion restriction in ADW images.

These are benign slow-growing radio-resistant tumors with rare chance of malignant transformation in long-standing cases. Tsugu et al reported a case of squamous cell carcinoma arising in an intracranial DC. Complete surgical excision renders effective treatment for prevention of recurrence and complications. But in cases of extensive adhesion between the cyst wall and the surrounding vital neurovascular structures, subtotal resection can be contemplated.

**Conclusion**
The authors have reported an extremely rare case of laterally located cerebellar convexity DC in a middle-aged lady with both interdural and extradural components with unusual MRI findings in T1-weighted and postcontrast sequence images. Maximum safe resection should be the goal of surgery considering the low risk of recurrence.

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References