Dorsal Intradural Extramedullary Epidermoid in a One Year Old Child—A Case Report and Review of Literature

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Abstract

Epidermoids in the spinal canal are rare. They usually occur as a consequence of trauma or in the presence of an underlying dysraphism. Epidermoids are also usually seen in the lumbar region with an associated dorsal tract or sinus connecting them to the skin. We present an unusual case of a dorsal spinal epidermoid without any dysraphism, or history of trauma. The lesion was diagnosed on MRI and the lesion was completely excised surgically through a laminectomy and durotomy. Histopathological assessment confirmed the diagnosis of an epidermoid. To our knowledge this is the eighth such report in literature, making this an unusual and interesting report. A review on relevant literature is discussed in conjunction to this case as well.

Keywords: Epidermoid; Paediatric; Spine

Key Messages: There are only 8 reported cases of intradural extramedullary epidermoids without associated trauma or spinal dysraphism so far. Thus this occurrence along with its successful surgical excision presents a unique representation of spinal epidermoids and their natural history, along with a review of relevant literature.

Introduction

Spinal epidermoids are rare, and encompass less than 1% of the lesions seen in the spine [1]. They are seen frequently in association with dysraphism, sinuses and other congenital anomaly of the spine. Aquired epidermoids are seen in conjuncture with repeated lumbar punctures and trauma. The presence of an epidermoid without any of the above mentioned associated lesions or trauma is very rare [1,2]. We present a case of a de novo epidermoid of the dorsal spine without dysraphism, trauma or syndromic association which was successfully identified and removed surgically with no deficits for the patient. We present a review of relevant literature along with this case report.

Case Report

A 1 year old female child presented to the outpatient clinic with a brownish discoloured skin patch on the back with a slight swelling observed since birth. The child had no deficits and was healthy, feeding and playing well. On consultation with a General Practitioner (GP) in West Bengal (where the parents hailed from), they were told about possible underlying spine deformities that could exist. Hence they came to us for treatment. On examination, no other neurocutaneous markers were seen. There were no deficits, or deformities perceptible. No evidence of dysraphism existed. On Imaging of the child’s neuroaxis using a Magnetic Resonance Imaging (MRI), an intradural extramedullary lesion was detected at the level of D10 posterior to the cord.

Figure 1 showed a hyperintense intradural extramedullary lesion in right side of spinal canal from D9-D10 disc level to D10 vertebral level. The lesion was seen to be causing leftward displacement and compression on spinal cord. Figure 2 showed a T1W sagittal images of the lesion where it appeared hypointense. Figure 3 were the Apparent Diffusion Coefficient (ADC) and Diffusion weighted imaging (DWI) sequences respectively which show diffusion restriction within the lesion. This demonstrated that the cyst present was not a solid tumor. Figure 4 which was a T1W fat Suppressed sagittal postcontrast sequence showed mild enhancement along the superior margin of the lesion, along with demonstrating some enhancement along the sinus tract. Figure 5 consisted of T2W sagittal and axial images respectively which showed an oblique sinus tract extending from the left paramidline region of skin surface at D12 vertebral level up to the posterior dura at D10 vertebral level. The sinus tract appeared to be predominantly hypointense on T2W sequence (Figures 1-5).

No spinal dysraphism was seen. No tethering of cord, split cord or abnormal cystic dilatation of the spinal canal was seen. No bony defects

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were detected on X-ray of the spine. Thus based on the data revealed by the imaging modalities employed, an epidermoid was considered as the primary diagnosis along with the possibility of differentials such as a dermoid cyst with tract/sinus.

After explaining the situation to the parents and discussing the case with the in house paediatric neurologist, she was taken up for surgery. We planned a dorsal D9, D10 laminectomy with a midline durotomy and excision of the tumor (Figures 6-9).

The child was positioned prone on bolsters after induction of GA. Motor Evoked Potential (MEP) monitoring electrodes were connected to the different muscle groups and the circuit tested before draping the patient. The incision was marked with fluoroscopic guidance over D10. Laminectomy of D10 and a part of D9 was done in a standard fashion. The dural tube was seen with a tract extending dorsally into the muscular and subcutaneous plane (Figure 6). Dura was opened to reveal the tumor which was seen posterior to the cord, which was flattened and pushed anteriorly (Figure 7). The tumor capsule was opened dorsally and the tumor was decompressed before the capsule was dissected off the adjacent nerve roots and cord and removed completely (Figure 8). The contents of the lesion were a putinaceous yellowish white substance devoid of vascularity. At the end of the dissection, the cord and nerve
roots inside a clean dural tube devoid of tumor spillover was seen (Figure 9).

Closure was done in a standard fashion. The child was extubated post surgery and after a brief spell at the recovery room was shifted to the mother's side at the room. No deficits were present after surgery. No drain was placed. The child was discharged in 2 days with no complaints and was followed up regularly. No CSF leak, wound complications or weakness of limbs were seen after surgery. Urinary incontinence was absent after removal of the indwelling foley's catheter on Postoperative Day (POD) 1.

The histopathological examination of the specimen revealed it to be an epidermoid cyst. Histologically, epidermoid and dermoid cysts are lined by stratified squamous epithelium supported by an outer layer of collagenous tissue, progressive desquamation of keratin from epithelial lining toward the interior of the cyst produces a soft white material. Differentiation between the two forms is based on the presence of skin adnexa and adipose tissue presence in the wall of dermoid cyst, which is absent in the epidermoid. As this differentiation is clearly made Immuno-histochemistry (IHC) confirmation was deemed unnecessary by the pathologist. Figure 10 shows the Cyst wall lined by atrophic stratified squamous epithelium, lumen filled with lamellated keratin material (Figure 10). No skin appendages are seen. (H&E 20X)

The child continues to be on regular follow up every 6 months and has made a complete recovery devoid of deficits or other complications associated with such maladies.

**Discussion**

“Epidermoids” are uncommon benign lesions with an incidence of less than 1% in the spine. They commonly present in the fourth decade
with a slight male preponderance [1]. Described by Cruveilhier as tumeurs perlées (pearly white tumors) due to their gross appearance, they consist of soft, whitish, keratin material without any element of skin appendage [1]. Congenital epidermoids are frequently found in association with spinal dysraphic conditions such as tethered cord, low lying conus, dermal sinus, or spina bifida, while acquired ECs occur, following repeated lumbar punctures or trauma [2]. Hence a lesion seen in the absence of dysraphism, syndromic association or trauma makes this a rare entity indeed.

Among the spinal epidermoids the thoracic region is the commonest site of occurrence, followed by the sacral and cervical regions [2]. Epidermoids may extend into the spinal canal or laterally through the neural foramina. They may extend ventrally as well, breaching the anterior or posterior vertebral body. Urinary dysfunction with or without back pain may be the presenting symptom. Others may present with cutaneous manifestations of spinal dysraphism or meningitis, as was in our case [3-5].

MRI is the imaging modality of choice for the diagnosis of Spinal Epidermoids. The tumor is classically a nonenhancing lesion of “cerebrospinal fluid intensity” that is isointense or hypointense on T1-weighted sequences and hyperintense on T2-weighted sequences. Discrepancy in the intensity characteristics occur due to varying lipid and protein components. Abnormalities in vertebral bodies or posterior elements may be evident in long-standing lesions. DW imaging demonstrates restricted diffusion such as in ECs elsewhere, and may be of benefit while evaluating a cystic sacral lesion [5-7].

Total excision remains the treatment of choice in symptomatic lesions compressing the thecal sac [5,6]. However, a densely adherent capsule may preclude total excision resulting in early relapses. Residual cystic contents predispose to the occurrence of aseptic meningitis in the postoperative period, a major complication that can result in the development of normal pressure hydrocephalus [6-9].

Conclusion

Spinal epidermoids are rare and progress along a benign course. As in the majority of CNS tumors, symptoms secondary to pressure effects manifest frequently. The presence of factors such as dysraphim, syndromic predilection and trauma must be taken into account, when considering such a diagnosis. Imaging remains extremely important to the diagnosis and follow up of patients. Surgical excision remains the best and perhaps only satisfactory treatment modality. Complete excision is curative and offers the prospect of a disease free life devoid of deficits if done well.

References