Case Report

Solitary juvenile xanthogranuloma of the cervical spine in a child: A case report and review of literature

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ABSTRACT
The authors present a case of 15 years male child who presented with neck pain and progressive ascending quadriparesis. Magnetic resonance imaging showed lytic mass involving C5 and C6 vertebra with soft tissue extension. Surgical excision of mass done using anterior cervical approach. Postoperatively, patient showed improvement in spasticity and power. Histopathological examination of mass was suggestive of juvenile xanthogranuloma (JXG). At 6 months follow-up, patient was improving without any evidence of recurrence. Only 12 cases of JXG of spine have been reported till date including only four cases involving the cervical spine and among these four cases only two were of pediatric age group.

Key words: Cervical vertebra, histology, myelopathy, surgery, xanthogranuloma

Introduction
JXG is a rare proliferative histiocytic disorder that usually occurs in childhood and is regarded as a non-Langerhans cell histiocytosis.[1] It is not a true neoplasm, but it is thought to be a reactive proliferation of histiocytes. Etiology and pathogenesis of JXG remains unknown. It usually presents as a cutaneous and self-limited lesion in the first two decades of life. Extracutaneous lesions, especially in disease involving the spine are rare.[2] We report a case of a 15-year-old child who presented to us with compressive myelopathy at C5–C6 level with a mass lesion involving C5–C6 vertebra and was treated with surgical excision of the mass. On histopathological examination, the mass was found to be xanthogranuloma. To the best of our knowledge, this is 3rd reported case of JXG cervical spine in pediatric age group.

Case Report

History and examination
A 15-year-old male presented with pain in the nape of the neck and gradually progressive ascending weakness of all four limbs along with bladder and bowel involvement for 3 years. On examination, tone was increased in all four limbs with 4/5 power in bilateral upper limbs and 2/5 power in both lower limbs, along with exaggerated deep tendon reflexes in all four limbs. Superficial abdominal reflexes were absent and, bilateral planters were extensor. He had 70–80% sensory loss below T4 level. His routine laboratory studies including hematological and biochemical parameters were within normal limits.

Imaging studies
Computerised tomography scan of the cervical spine revealed a large osseous-destructive lesion involving body of C5 and C6 vertebra which was causing spinal canal compromise [Figure 1a and b]. The lesion was involving surrounding structures with extension into the submuscular plane anteriorly. On magnetic resonance imaging (MRI) evaluation, the mass was isointense on T1 and T2 weighted sequences with minimal contrast enhancement. It was causing distraction of C5 and C6 vertebra body with compression of thecal sac [Figure 2a-d].

Operative procedure
Using anterior cervical approach, the tumor resection was performed. Intra-operatively, the tumor involved whole of the C5 and C6 vertebral body. There was a soft tissue component of the tumor, which was extending into the submuscular plane. Inside the capsule, the tumor consisted of cheesy material...
that was minimally vascular. Total tumor excision was done with tricorticate iliac crest bone grafting and plate and screw fixation between C4 and C7 vertebral bodies.

**Pathological examination**

Microscopic examination showed spindle cells disposed of in intersecting fascicles intermingled with mononuclear xanthomatosus cells displaying round nuclei with dense chromatin inconspicuous nucleoli and pale eosinophilic finely vacuolated chromatin. Spindle cells display bland appearing nuclear chromatin and eosinophilic cytoplasm. Occasional cells displayed nuclear atypia and enlargement. Scattered neutrophils were seen admixed with these cells. Focal areas of calcification and occasional osteoclasts like giant cells were also seen. Interspersed areas showed degenerated bony trabeculae and cartilage [Figure 3a and b].

**Postoperative course**

Postoperatively, patient had significant improvement in neck pain and spasticity in all four limbs. At follow-up of 6 months, he was doing well.

**Discussion**

JXG was first reported by Adamson in 1905 and was termed as a congenital xanthoma multiplex. JXG is not a true tumor, but rather a reactive proliferation of histiocytes. JXG is the most common non-Langerhans histiocytosis. Most tumors present early in life, and male predominance has been noted in childhood. Despite the term juvenile, JXG can present during adulthood and is most commonly seen during the third and fourth decade, but no sex predilection is seen in adulthood. In pediatric patients, it is more common in males. Central nervous system involvement in cutaneous and extracutanous JXG is rare and can be classified according to the location. (1) intraparenchymal (2) dural (3) cranial or spinal column.

Solitary JXG especially involving the spine is very rare. Until date, only 12 cases have been described in English literature with solitary JXG of the spine. Table 1 summarizes the details of all the 12 cases. Four of these were in the cervical spine, 3 in the thoracic spine and 5 in the lumbosacral spine. JXG is a slow growing tumor and presents with features according to location of the tumor.

On MRI, JXG is isointense to hyperintense on T1-weighted sequences and T2-weighted sequences with no or minimal contrast enhancement. According to Schultz et al., spinal JXG does not enhance whereas enhancement can be seen in cerebral JXG. Spinal JXG should be distinguished from tumors of neural origin such as schwannoma, meningioma, lymphoma, etc.,. On gross examination, JXG usually are encapsulated with or without cystic component that is yellowish to gray in color. Definitive diagnosis can be made by pathological and immunohistochemical studies. Microscopically, JXG has foamy histiocytic cells with or without Touton giant cells in the background of mononuclear cell infiltrates. In immunohistochemistry studies, JXG has mononuclear cells,
giant cells and spindle cells, which are positive for lysosomal stain and CD68, but negative for CD1a and S-100 proteins, which are reactive markers for Langerhans cells.

No standard treatment protocol for treatment of JXG of spine has been described as these lesions are extremely rare. These lesions are slow growing benign lesion without regression, unlike cutaneous lesions. Total excision of the tumor should be done with preservation of normal structures and cervical motion. Recurrence has not been reported after gross total excision. Of the 12 cases, two patients underwent partial excision after which they underwent radiotherapy, no recurrence was reported in these patients in follow-up. Despite these results, total excision of the tumor appears to be curative due to the benign nature of the tumor. As the natural course of solitary JXG is not known, so patient should be kept in the long term follow-up.

Our patient was treated with near total excision of the tumor with C5 and C6 corpectomy using anterior cervical approach with bone grafting and spinal stabilization. Postoperatively, patient had significant improvement in spasticity and power in all four limbs.

Cutaneous JXG is commonly found in pediatrics patients but till date only 12 cases of solitary JXG of spine have been reported. To the best of our knowledge, our case is 13th case of solitary xanthogranuloma of spine and 5th in the cervical spine. This is the third case in the pediatric age group affecting the cervical spine.

**Conclusion**

We present a histologically confirmed case of solitary JXG of the cervical spine in 15-year-old child. JXG should be considered to be a differential diagnosis for such tumors of the spine. Total excision of the tumor should be attempted with preservation of normal neural structures. As these lesions are benign, total excision appears to be curative. These patients should be kept in the long term follow-up as the natural course of these tumors is not known.

**References**


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**Table 1: Summary of the reported cases of JXG in the literature**

<table>
<thead>
<tr>
<th>Authors/year</th>
<th>Age/sex</th>
<th>Clinical features</th>
<th>Location</th>
<th>Surgery</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shimosawa et al.[10] 1993</td>
<td>13 months/female</td>
<td>Spastic paraparesis</td>
<td>T6–9 IDEM</td>
<td>T6–T9 laminectomy with excision of tumor</td>
<td>Improved</td>
</tr>
<tr>
<td>Kitchen et al.[11] 1995</td>
<td>15 years/female</td>
<td>Low back pain</td>
<td>S2 nerve root</td>
<td>L5 laminectomy with total excision</td>
<td>Symptoms improved</td>
</tr>
<tr>
<td>Kim et al.[12] 1996</td>
<td>16 months/female</td>
<td>Paraparesis</td>
<td>T1–2 IDEM</td>
<td>C7–T3 laminectomy with total excision of tumor</td>
<td>Resolution by 4 months</td>
</tr>
<tr>
<td>Iwasaki et al.[13] 2001</td>
<td>41 years/female</td>
<td>Weakness and numbness in both lower limbs</td>
<td>Cauda equina</td>
<td>L1–L5 laminectomy with excision of tumor</td>
<td>Not mentioned</td>
</tr>
<tr>
<td>Rampini et al.[14] 2001</td>
<td>34 years/female</td>
<td>Spastic quadriparesis</td>
<td>C5–C7 IDEM</td>
<td>C5–T1 laminectomy with total excision of tumor</td>
<td>Improved</td>
</tr>
<tr>
<td>Dehner et al.[15] 2003</td>
<td>14 years/female</td>
<td>Back pain</td>
<td>L3 vertebral body</td>
<td>Not mentioned</td>
<td>Not mentioned</td>
</tr>
<tr>
<td>Cao et al.[16] 2008</td>
<td>18 years/female</td>
<td>Neck pain</td>
<td>C2 root</td>
<td>C1–C2 laminectomy with total excision of tumor</td>
<td>No recurrence at 2 years</td>
</tr>
<tr>
<td>Castro-gago et al.[17] 2009</td>
<td>14 years/male</td>
<td>Paraparesis</td>
<td>Cauda equina</td>
<td>L2–L3 laminectomy with partial excision</td>
<td>Postoperative RT, no recurrence at 2 years</td>
</tr>
<tr>
<td>Lee et al.[18] 2011</td>
<td>29 years/male</td>
<td>Hemiparesis with contralateral hypothesia below C2</td>
<td>C1–C2</td>
<td>C1–C2 laminectomy with total excision of tumor</td>
<td>Improved</td>
</tr>
<tr>
<td>Jain et al.[19] 2011</td>
<td>22 years/female</td>
<td>Progressive back pain</td>
<td>T7 vertebral body</td>
<td>T7 laminectomy with excision of tumor</td>
<td>Improved</td>
</tr>
<tr>
<td>Agabegi et al.[20] 2011</td>
<td>47 years/female</td>
<td>Bowel, bladder involvement and back pain</td>
<td>L2 vertebral body</td>
<td>T12–L3 laminectomy with partial excision</td>
<td>Postoperative RT, improved at 8 months</td>
</tr>
<tr>
<td>Inoue et al.[21] 2011</td>
<td>38 years/male</td>
<td>Numbness and weakness of right hand</td>
<td>C8 nerve root</td>
<td>C7–T1 laminectomy</td>
<td>No recurrence at 2 years</td>
</tr>
<tr>
<td>Present case/2013</td>
<td>16 years/male</td>
<td>Spastic quadriparesis with neck pain</td>
<td>C5–C6</td>
<td>Anterior cervical approach with excision of tumor</td>
<td>Improved no recurrence at 6 months</td>
</tr>
</tbody>
</table>

RT – Radiotherapy; IDEM – Intradural extramedullary
Case report. J Neurosurg 2000;93:335-41


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