Corpus Callosum Lipoma with Extracranial Extension Associated with Seizure

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Abstract
Intracranial lipoma is a congenital anomaly usually found incidentally. Pericallosal lipoma with extension into extracranial tissue through interhemispheric structures are not common and when this occurs, it is usually accompanied by other intracranial malformations. We discuss this rare case having corpus callosal lipoma with subcutaneous lipoma, a tubulonodular type, manifesting as a seizure in adulthood. Due to extra morbidity after surgery, we planned to follow up the patient’s with conservative management.

Keywords: Intracranial Lipoma; Extracranial Extension; Seizure

Introduction
Intracranial lipoma is a congenital benign lesion which constitute about 0.1% of all intracranial tumors. It is thought to arise either from lipoid cells of pia mater or fatty transformation of connective tissue or from embryonal remnants or from dedifferentiation of the meninx primitive to lipomatous tissue. Most of them are associated with congenital malformations. It is very rarely associated with extracranial extension [1-3]. We report a case of 31 year-old male who was referred from remote village hospital with the complaint of occasional seizures. We diagnosed the case as corpus callosal lipoma with extension into subcutaneous tissue. We will highlight here brief description of the rare case, image findings and our management strategy.

Case Report
A 31 year-old male farmer came to our hospital with the complaint of slow growing scalp swelling and on and off headache since birth and the occasional seizure for 2 months. Headache used to be mild and controls with analgesics so he didn’t consult the doctor before. During the last 2 months, he had tonic clonic seizure twice in a month. There was history of brief loss of consciousness associated with seizure. There was no history of congenital anomalies within the family. General physical examination of patient revealed a subcutaneous non mobile swelling of size 10 x 15 cm was present in the frontoparietal region it was in midline, rubbery and devoid of hairs. The swelling was also extended in the forehead. Neurological examination including funduscopic examination revealed normal findings. There was no sign of facial anomalies. Laboratory data were noncontributory.

We did MRI of brain which showed there was huge lesion in the callosal, pericallosal and supracallosal region extending into subgaleal region through a defect in the frontal bone. The lesion was hyperintense in both T1W and T2W images which was sufficient for diagnosis (Figure a-e). DWI showed no restriction of diffusion. Posterior part of body and splenium of corpus callosum were absent suggestive of agenesis of corpus callosum. Lateral ventricles were also invaded by the lesions.

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Figure a and b: T2WI image showing huge hyperintense lipoma with extracranial extension.

Figure c-e: Showing T1WI image which is hyper intense and absence of corpus callosum.

After our diagnosis was confirmed we called a meeting of all neurosurgeons in our hospital for individual views regarding the management. We concluded that the subcutaneous scalp swelling can be removed for cosmetic purposes. For intracranial part we decided not to touch as aggressive gross total removal may cause additional morbidity to patient and the seizure for which patient seek treatment may not subside seeing the location of lesion. So we decided to follow up the patient with anticonvulsants and occasional use of analgesics for headache. We clearly described the pros and cons of surgery to the patient. He didn’t agree to go through surgical interventions. Ultimately, we prescribed anticonvulsants and analgesics.

Discussion

Intracranial lipoma constitutes about 0.06 - 0.46% of intracranial lesions which is usually discovered incidentally. It is thought to be a congenital malformation and are generally asymptomatic. The most common manifestations are headache, seizures and raised intracranial pressure [1,2,4-6]. The most common locations are interhemispheric fissure, corpus callosum, quadrigeminal-ambient cistern, infundibular chiasmatic region, cerebellopontine angle and sylvian fissure [1,2]. Pericallosal lipoma constitute about half of all intracranial lipoma and is usually associated with agenesis of corpus callosum similar to our case. It may extend to interhemispheric fissure connecting to extracranial tissue, may extend into ventricles or may be associated with frontonasal dysplasia [6,7].

Interhemispheric lipoma has 2 types. Tubulonodular type is anteriorly located, larger, occurs before closure of anterior neuropore and is associated with agenesis of corpus callosum, calcifications and frontal bone defect whereas curvilinear type occurs posteriorly and is thin and not associated with cerebral anomalies [7,8].

Pericallosal interhemispheric lipoma may have extracranial connection through fibrous-lipomatous stalk or direct connection through cranium bifidum or mayn’t have connection between extracranial intracranial parts. It is thought that this connection may result from secondary dehiscence of cranium with invagination of small tuft to meninx primitive [5]. Some of the published cases having intracranial lipoma with extension into extracranial has been summarized in table 1.

<table>
<thead>
<tr>
<th>Case</th>
<th>Age</th>
<th>Sex</th>
<th>Symptoms</th>
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<tbody>
<tr>
<td>Kinal., et al.</td>
<td>3 months</td>
<td>M</td>
<td>Midfrontal tumor</td>
</tr>
<tr>
<td>Cant., et al.</td>
<td>5 years</td>
<td>M</td>
<td>Frontal swelling, seizures</td>
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<tr>
<td>Kuwabara., et al.</td>
<td>17 years</td>
<td>F</td>
<td>Ocular hypertelorism, forehead swelling</td>
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<tr>
<td>Addlestone., et al.</td>
<td>4 months</td>
<td>M</td>
<td>Forehead swelling</td>
</tr>
<tr>
<td>Kushnet., et al.</td>
<td>64 years</td>
<td>F</td>
<td>Convulsions, forehead swelling saddle nose deformity</td>
</tr>
<tr>
<td>Hyashi., et al.</td>
<td>3 yrs</td>
<td>F</td>
<td>Ocular hypertelorism</td>
</tr>
<tr>
<td>Kudoh., et al.</td>
<td>3 Months</td>
<td>F</td>
<td>Swelling in forehead, hypertelorism</td>
</tr>
<tr>
<td>a.sari., et al.</td>
<td>3 months</td>
<td>M</td>
<td>Swelling over anterior fontanelle</td>
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<tr>
<td>Chen., et al.</td>
<td>newborn</td>
<td>M</td>
<td>Swelling over vertex</td>
</tr>
<tr>
<td>Given., et al.</td>
<td>9 weeks</td>
<td>M</td>
<td>Anterior fontanelle mass</td>
</tr>
<tr>
<td>Ahmetoglu., et al.</td>
<td>5 months</td>
<td>F</td>
<td>Midline parietal mass</td>
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<td>Reddy., et al.</td>
<td>13 yrs</td>
<td>M</td>
<td>Occipital swelling</td>
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<td>Karabag., et al.</td>
<td>26 yrs</td>
<td>F</td>
<td>Vertex scalp swelling</td>
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<tr>
<td>Rai., et al.</td>
<td>40 yrs</td>
<td>M</td>
<td>Forehead swelling</td>
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<tr>
<td>Aggrawal., et al.</td>
<td>22 yrs</td>
<td>F</td>
<td>Scalp swelling</td>
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<tr>
<td>Chaurasia., et al.</td>
<td>31 yrs</td>
<td>M</td>
<td>seizure</td>
</tr>
</tbody>
</table>

Table 1: Several Cases of intracranial lipoma with extension into extracranial [3].
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MRI is superior to all imaging modalities over diagnosis of intracranial lipoma having characteristics hyper intensity in both T1W and T2W image and defining relationship to the surrounding structures [9].

Treatment is usually not requires as many patients are asymptomatic which can be followed with observation. Some has symptoms of raised intracranial pressure, seizures and cosmetic deformity which may necessitate interventions, surgical excision of scalp lesion may be achieved for cosmetic lesions but at that time the cranial gap should be obliterated with pericranium to avoid recurrence. Due to very dense adhesions of lipoma to surrounding structures intracranial parts should not be usually removed to minimize morbidity and mortality [4].

Conclusion

In conclusions seeing the nature of this huge lesion conservative management with anticonvulsants and analgesics are sufficient to satisfy the patient. It also avoids morbidity due to surgical damage to the intracranial vessels and surroundings vital structures which are tightly adhered to the lipoma. The rarity of this disease and the huge size (10 x 15 cm) compelled us to report this case for future learnings. None of the cases described in the literature report a larger size than our case.

Bibliography


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