Radiculopathy in the setting of lumbar nerve root compression due to an extradural intraforaminal lipoma: a report of 3 cases

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A true adult spinal lipoma is an exceedingly rare cause of lumbar compression neuropathy. Only 5 cases of true extradural intraforaminal lipomas have been documented in the medical literature. The diagnostic criteria and treatment guidelines for this specific lipoma have yet to be established. This report features 3 histologically confirmed cases of extradural intraforaminal spinal lipomas that recently presented to the authors’ practice. In addition, the literature was surveyed to include the 5 previously reported cases of true adult extradural intraforaminal spinal lipomas. The consistency in presentation, response to surgical intervention, and postoperative recovery in these 8 cases supports surgical intervention at the time of diagnosis. The authors’ findings support elevated clinical suspicion, efficient diagnosis based on MRI, and early surgical intervention for this rare pathological entity. All cases presented in this report were symptomatic and occurred in the absence of other significant pathologies such as general spinal epidural lipomatosis, intradural lesions, tethering, or severe degenerative stenosis or herniated discs. The clinical, neuroradiological, and histological findings characteristic of a true adult extradural intraforaminal lipoma are emphasized to differentiate this lesion from the more common etiologies for lumbar compression neuropathy. Heightened awareness and clinical suspicion for the focal, foraminal spinal lipoma as a cause of radiculopathy symptoms will enable more efficient diagnosis and treatment.

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KEY WORDS spine; adult extradural intraforaminal spinal lipoma; radiculopathy; lumbar

AN extradural intraforaminal spinal lipoma in the absence of myelospinal dysraphism is an exceedingly rare pathological entity. Clinically, this lesion manifests as a unilateral radiculopathy refractory to conservative management. Although true adult spinal lipomas account for only 0.4%–0.8% of all intraspinal tumors, the associated constellation of lumbosacral radiculopathy symptoms are estimated to affect 3%–5% of the adult population in the US. The differential for common lumbar radiculopathy is vast and most frequently can be attributed to common pathologies such as disc herniation or spinal stenosis. Next to these highly prevalent etiologies, the possibility of compressing lipomatous tissue should not be excluded. Magnetic resonance imaging is the gold standard in diagnostic imaging for lumbar compression neuropathy due to a potential extradural intraforaminal lipoma. Magnetic resonance imaging of a lipomatous lesion, such as a true adult intraforaminal lipoma, angiolipoma, or spinal epidural lipomatosis (SEL), will reveal hyperintensity on T1-weighted sequences with decreased to intermediate intensity on T2-weighted sequences. Diagnosis is further complicated by the difficulty in differentiating a true lipoma from more prevalent lesions such as an angiolipoma or SEL. If surgical exploration is performed, careful histological examination of the excised mass is essential for accurate differentiation of a true lipoma from an angiolipoma.

The objective of our retrospective case series and survey of the literature is to increase clinical suspicion for this rare etiology of lumbar radiculopathy. We encourage the differentiation of true spinal lipomas from SEL, angiolipomas, and degenerative sources of lumbar radiculopathy via diagnostic MRI and histopathology.

Case Reports
Case 1
History and Examination
A 38-year-old nonobese man presented to our practice with a medical history significant for chronic left L-5 radiculopathy. The patient’s chief complaint was progres-
sively worsening weakness and “shooting” left-sided radicular pain. The radiculopathy had proven refractory to all attempted conservative treatment modalities over the preceding 6-year interval. Repeat MRI of the lumbar spine revealed a left L5–S1 lumbosacral mass that involved the entire left L-5 foramen with caudal displacement of the L-5 nerve root (Fig. 1).

Operation

Elective left laminectomy, medial facetectomy, foraminotomy, and resection of an encapsulated lesion were performed without complication. The lesion was excised, and interval decompression of the involved foramen and nerve root was achieved. Gross examination of the lesion revealed a lobulated, yellow fatty mass encapsulated by a thin layer of fibrous tissue. Histopathology showed no evidence of vascular atypia and was consistent with a true adult extradural intraforaminal lipoma.

Postoperative Course

Postoperatively, the patient recovered without complication. At his 3-month follow-up visit, he reported markedly attenuated radicular symptoms; there was a drastic reduction in his low-back pain, left leg pain, paresthesia, and weakness. Steady improvement was noted at the 6- and 10-month follow-up visits with continued alleviation of his left lower-extremity radiculopathic pain. By the 10-month follow-up, he was comfortable and ambulating normally with full motor strength and range of motion.

Case 2

History and Examination

A 40-year-old man presented to our practice with refractory left L-3 radiculopathy manifesting as persistent weakness, paresthesia, and severe radiating pain. The radiculopathy was refractory to 2 years of conservative medical management. Initial MRI in the patient suggested a radiculopathy caused by lumbar stenosis due to foraminal lumbar epidural lipomatosis rather than an organized L3–4 lipoma causing cranial displacement of the root within the foramen (Fig. 2).

Operation

Elective surgical intervention with L3–4 left laminectomy, medial facetectomy, foraminotomy, and resection of the mass was performed. Gross examination in surgery revealed a yellow, lobulated epidural mass within a fibrous capsule. Histology confirmed adipose tissue in the absence of vascular atypia, most consistent with a true adult lipoma.

Postoperative Course

The elective procedure was tolerated well by the patient, and there were no postoperative complications. At the 3-month follow-up, the patient showed marked improvement in his symptoms with alleviated pain and significantly improved strength in the left lower extremity. At the 6-month follow-up, he was ambulating at the baseline level with resolution of the lumbar radiculopathy. He had maintained these results without complication or symptomatic recurrence at his 12-month follow-up visit.

Case 3

History and Examination

A 44-year-old man presented with the chief complaint of chronic radicular pain in an L-4 distribution pattern. Conservative treatment efforts consisted of transcutaneous electrical nerve stimulation, epidural steroid injections, physical therapy, acupuncture, and analgesic pharmacotherapy. Magnetic resonance imaging revealed an L-4 intraforaminal lesion with mass effect on the lateral thecal sac and nerve root at the foramen (Fig. 3). There was no evidence of severe degenerative stenosis.

Operation

Elective surgical intervention with L4–5 left laminectomy, facetectomy, and foraminotomy for resection of the mass was performed without complication. An encapsulated lipomatous lesion was excised, and the mass was confirmed to contain true lipomatous tissue without vascular atypia on final histopathology.

Postoperative Course

The patient’s postoperative course was encouraging, without complication or relapse. At his 3-month follow-up assessment, he reported drastic improvement in his radiculopathic pain as well as significantly improved strength.
in his left lower extremity. At the 6-month follow-up, he
demonstrated continued improvement in strength and am-
bulation. He did have a minor trauma, with increased back
pain prompting lumbar MRI during an emergency room
visit. The imaging did show interval resection of the fo-
raminal lipoma without impingement on the lateral recess.
This progress was maintained at the 12-month follow-up.
The patient’s only complaint at this visit was mild back
pain, which was attributed to a traumatic incident.

Discussion
Spinal lipomas are unique pathological entities rarely
observed in the absence of myelospinal dysraphism.4 Given
the rarity and characteristically benign MRI findings of a
true spinal lipoma, this lesion can be easily overlooked or
misdiagnosed. Symptoms related to mass effect and sec-
ondary compressive myelopathy from a lipomatous mass
could be caused by a true lipoma, an angiolipoma, or an
SEL. True adult lipomas must be distinguished from both
an angiolipoma and an SEL.15 The process of differentiat-
ing between these lipomatous lesions begins with a survey
of patient risk factors for SEL. Six of the 8 patients con-
sidered in this report presented with none of the 3 primary
characteristics associated with SEL: chronic exogenous
steroid use, endocrinopathy, and obesity (Table 1).31 The
remaining 2 patients presented with only 1 of the estab-
lished SEL risk factors: obesity. Magnetic resonance imag-
ing can be used to reliably differentiate a true adult lipoma
from an SEL, which consists of a diffuse overgrowth of
epidural adipose tissue within the spinal canal. To differ-
entiate a true adult lipoma from an angiolipoma, however,

<table>
<thead>
<tr>
<th>Authors &amp; Year</th>
<th>Age (yrs), Sex</th>
<th>BMI (kg/m²)</th>
<th>Presentation</th>
<th>Imaging Findings</th>
<th>Postop Complications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Present Case 1</td>
<td>38, M</td>
<td>24.8</td>
<td>L-5 radiculopathy</td>
<td>L5–S1 L mass involving entire foramen; 1.1 × 1.1 × 0.3–cm epidural mass; no degenerative stenosis</td>
<td>None; significant improvement of symptoms by 6-mo FU; continued improvement in back &amp; leg pain by 10-mo FU</td>
</tr>
<tr>
<td>Present Case 2</td>
<td>40, M</td>
<td>40.0</td>
<td>L-4 radiculopathy</td>
<td>L3–4 Lt recess &amp; foraminal mass; 2.5 × 1.6 × 0.3–cm epidural mass; no degenerative stenosis</td>
<td>None; significant improvement of symptoms by 6-mo FU; complete resolution of leg pain by 12-mo FU</td>
</tr>
<tr>
<td>Present Case 3</td>
<td>44, M</td>
<td>29.0</td>
<td>L-4 radiculopathy</td>
<td>Lt L4–5 foraminal mass; 1.1 × 0.8 × 0.2–cm epidural mass; no degenerative stenosis</td>
<td>None; significant improvement of symptoms by 6-mo FU; complete resolution of symptoms w/ only mild back pain noted at 12-mo FU</td>
</tr>
<tr>
<td>Zevgaridis et al., 2008</td>
<td>62, F</td>
<td>&lt;30</td>
<td>L-5 radiculopathy</td>
<td>Compression of Lt L-5 nerve root by an extradural intramural mass; no degenerative stenosis</td>
<td>None</td>
</tr>
<tr>
<td>Kim et al., 2012</td>
<td>55, M</td>
<td>31.6</td>
<td>L-5 radiculopathy</td>
<td>Epidural mass posterior to L-5 vertebral body</td>
<td>None; complete resolution of symptoms by 2-yr FU</td>
</tr>
<tr>
<td>Dierckx et al., 1987</td>
<td>43, M</td>
<td>&lt;30</td>
<td>S-1 radiculopathy</td>
<td>Extrudal filling defect at L5 vertebra on myelogram</td>
<td>None; complete resolution by 3-mo FU</td>
</tr>
<tr>
<td></td>
<td>62, M</td>
<td>&lt;30</td>
<td>L-5 radiculopathy</td>
<td>Extrudal filling defect on myelogram w/ minor L5–S1 disc prolapse</td>
<td>None; complete resolution by 10-mo FU</td>
</tr>
<tr>
<td></td>
<td>56, M</td>
<td>&lt;30</td>
<td>L-4 radiculopathy</td>
<td>Extrudal filling defect in L3–4 on myelogram w/ some degeneration</td>
<td>None; symptomatic improvement noted post-operatively</td>
</tr>
</tbody>
</table>

BMI = body mass index; FU = follow-up.  
* There was no preoperative endocrinopathy in any of the listed cases, and steroid therapy was not used in any of the listed cases.
histopathological study is required. The high prevalence of radiculopathy due to common degenerative pathologies, such as disc herniation or degenerative foraminal stenosis, can also limit consideration of the rare extradural intraforaminal lipoma.\textsuperscript{7,12,14} As a result of its challenging diagnosis and low clinical suspicion, a true adult spinal lipoma is very likely underrepresented in the medical literature.\textsuperscript{8,15}

The 3 patients who presented to our practice had endured years of equivocal MRI studies, failed conservative therapies, and inconclusive assessments by multiple spine specialists and surgeons. For all 3 patients, we confirmed our diagnosis of extradural intraforaminal lipoma via macroscopic and histological inspection after total resection. Contrasting the findings in our 3 patients with those in the 5 cases from the literature (Table 1), we were able to demonstrate high consistency in clinical presentation, diagnostic MRI, and responsiveness to surgical intervention. On postoperative follow-up, all patients with a confirmed extradural intraforaminal lipoma demonstrated significant alleviation of radicular pain and reversal of preoperative neurological deficits by the 3- to 6-month interval.\textsuperscript{2,6,15}

Overall, we found the MRI findings to be predictable across all cases. All 3 of our patients had a high-signal-intensity epidural lesion on T1- and T2-weighted imaging that was located within a foramen ipsilateral to the side of symptoms, causing possible impingement of the lateral thecal sac (Figs. 1–3). Magnetic resonance imaging consistently showed displacement of the nerve root cranially or caudally, leading to the “empty foramen sign.” This “missing root” on axial images is identifiable on sagittal imaging in which it is possible to see the root displaced up or down. Although diagnosis of a true adult spinal lipoma is challenging given the low clinical suspicion and subtle neuroradiological findings, the highly favorable prognosis and early recovery achievable through surgical intervention merits screening for this lesion in patients who suffer from refractory radiculopathic pain in the absence of degenerative stenosis or herniated disc.

Conclusions

In summary, although MRI offers a means for early identification and treatment of an extradural intraforaminal lipoma, the neuroradiological diagnosis for a true spinal lipoma is a challenging task. The subtle MRI findings for an extradural intraforaminal lipoma can be easily overlooked or dismissed within the realm of normal anatomical variation; therefore, clinical suspicion for these lesions must increase in the presence of a high T1 and T2 signal occupying the lateral recess and foramen with unilateral displacement of a nerve root cranially or caudally in the absence of severe degenerative findings. The empty foramen sign, in which the nerve root on the side of radiculopathy is not easily identifiable, is helpful in diagnosing the extradural intraforaminal lipoma (Figs. 1–3). Ultimately, histopathology is essential for accurate diagnosis. The 8 cases considered in this report support early surgical intervention for a symptomatic extradural lipoma. The prognosis following resection of this lesion has been favorable. Because of its rarity, the extradural intraforaminal lipoma continues to lack definitive treatment guidelines. Our findings in this small case series and retrospective analysis strongly support surgical intervention for symptomatic lesions. Although we report symptomatic improvement at 10–12 months after surgery, longer-term follow-up is needed to determine if patients are at risk for delayed recurrence.

References


Author Contributions

Acquisition of data: Gottfried, Loriaux. Drafting the article: Loriaux. Critically revising the article: all authors. Reviewed submitted version of manuscript: Loriaux. Approved the final version of the manuscript on behalf of all authors: Gottfried. Study supervision: Gottfried.

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