Spontaneous cerebrospinal fluid leak from an anomalous thoracic nerve root: case report

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The authors report the case of a 28-year-old woman with a spontaneous cerebrospinal fluid leak from the sleeve of a redundant thoracic nerve root. She presented with postural headaches and orthostatic symptoms indicative of intracranial hypotension. CT myelography revealed that the lesion was located at the T-11 nerve root. After failure of conservative management, including blood patches and thrombin glue injections, the patient was successfully treated with surgical decompression and ligation of the duplicate nerve, resulting in full resolution of her orthostatic symptoms.

http://thejns.org/doi/abs/10.3171/2016.4.SPINE151465

KEY WORDS cerebrospinal fluid; leak; anomalous; dura; spine; congenital; thoracic nerve root

Spontaneous intracranial hypotension most commonly affects adults and is more often encountered in women.2,3,7,10,18,19,22,23,25 Postural headache is frequently the only reproducible clinical symptom; however, associated nausea, vomiting, abducens nerve paresis, tinnitus, hyperacusis, and visual field defects have been reported.19,25

Cerebrospinal fluid (CSF) leaks underlie the majority of cases of spontaneous intracranial hypotension and may occur throughout the spine, with a predilection for the cervicothoracic junction and thoracic regions.19 Spontaneous CSF leaks have been associated with connective tissue disorders,19,21,22 leading to hypotheses that meningeal fragility results in dural perforation or the formation of diverticula following minor trauma.1,6,11,14,20 In untreated CSF leaks, the postural component of the headache may abate with chronicity; thus, undiagnosed CSF leaks may prove an occult cause of otherwise idiopathic headache.11,19

Spinal nerve root anomalies are rare but well-characterized entities in the lumbosacral spine. Anomalies have not been reported in the thoracic spine and have not been associated with CSF leaks. The authors present a unique case of intracranial hypotension secondary to a CSF leak at a congenital thoracic nerve root anomaly.

Case Report

History and Examination

A 28-year-old woman presented complaining of positional occipital headaches associated with tinnitus and nonspecific visual changes for 4 months with no history of recent trauma. Her symptoms were relieved in recumbence. No other neurological deficits were appreciated.

Imaging

MRI of the brain revealed leptomeningeal enhancement (Fig. 1), indicative of intracranial hypotension. The patient underwent CT myelography, which demonstrated extrathecal contrast with suspicion of a CSF leak at the left T-11 nerve root sleeve (Fig. 2).

Treatment

Nonspecific lumbar blood patches, lesion-directed blood patches, and thrombin glue injections provided temporary improvement, but all subsequently resulted in full return of symptoms. After failure of conservative therapies, surgical exploration by left foraminotomy was performed. A redundant left T-11 nerve root (Neidre and Macnab classification Type 2A13) was discovered, focally leaking CSF from the axilla. The nerves were then ligated proximal to the defect and a lumbar drain was placed (Fig. 3).

Postoperative Course

The patient was kept supine with flat bed parameters with a lumbar drain for CSF drainage (10–15 ml/hr) for 1 week postoperatively. She experienced complete reso-
olution of her positional symptoms with residual nonposi-
tional headaches of significantly reduced intensity. CT
myelography was subsequently performed to evaluate the
continued headaches and revealed no detectable CSF leak
(Fig. 4). MRI of the brain also revealed resolution of lep-
tomeningeal enhancement (Fig. 5). At the 1-year follow-
up visit, the patient reported resolution of all orthostatic
symptoms.

Discussion

Spontaneous CSF leaks have been associated with
connective tissue diseases including polycystic kidney
disease, Marfan and Ehlers-Danlos syndrome.19,21,22 The
patient had no diagnosis of connective tissue disease at
the time of presentation. The leak was discovered at the
axilla of a nerve root redundancy—suggesting an ana-
tomical or mechanical etiology. As such, we hypothesize
that the anomalous nerve root contributed to the failure
of the dura. This represents a rare and unusual etiology of
a CSF leak that has not previously been described in the
literature.

An embryological explanation for congenital nerve root
anomalies in humans has not been fully elucidated; how-
ever, animal studies implicate errors in axon guidance.
The process of an axon finding and attaching to its tar-
get muscle group relies on a complex series of chemical
gradients and receptor modulation.16 A dose-dependent
increase in spinal nerve–to-muscle mismatch (which pro-
duced aberrant nerve root anatomy) has been demonstrat-
ed in Class III semaphorin-deficient chick embryos.12,16,17

Spinal nerve root anomalies in humans are well de-
scribed in the lumbosacral spine but not in the thoracic
spine. Lumbosacral anomalies have been reported to oc-
cur in 8.5%9 to 30%4 of autopsy specimens. Clinically,
these have been largely implicated in the study of degener-
ative conditions of the spine due to the atypical nerve
conformation occupying a larger portion of the interven-
tebral foramen and also for placing the surgical patient at an
increased risk of iatrogenic injury during decompression.3

In a study of 46 cases, Postacchini et al.15 found that
21% of patients with an anomalous nerve root exhibited
concomitant lumbosacral bony abnormality, including
lumbarization of S-1, sacralization of L-5, and a case of
congenital absence of the facet nearest the anomaly. Pa-
tients with lumbosacral nerve root anomalies presented
with radicular symptoms if they had any symptoms due to
these lesions. No patients were reported to have symptoms
of a dural leak. While nerve root anomalies have not been
associated with congenital defects in the dura, the paucity
of available evidence allows for either a congenital or a
mechanical etiology of our patient’s leak. We posit 2 hy-
potheses: 1) that the dura was weakened by the bifurca-
tion of the nerve root or 2) that repeated microtrauma between
the redundant roots within a stenotic foramen eroded the
dura. A tethering effect on a single nerve root in an anom-
alous pair has been described as a cause of radicular pain5
and may also explain undue tension at the axilla of the
anomaly.

Definitive diagnosis and localization of a CSF leak may
be achieved using a variety of imaging techniques, includ-
ing CT myelography, MR myelography, and radioisotope
cisternography. Of these 3 techniques, CT myelography
has been found to be the most effective at identifying the
defect.19,21 In our patient, CT myelography localized the
leak but was unable to identify the nerve anomaly. These
findings are concordant with a study by Haijiao et al.,8
which showed MRI to be superior for diagnosing nerve
anomalies.

Surgery to repair a CSF leak is reserved for patients

FIG. 1. Axial contrast-enhanced T1-weighted MR image of the brain
revealing diffuse leptomeningeal enhancement.

FIG. 2. Coronal (A) and axial (B) CT myelograms revealing evidence of
CSF extravasation at the level of the left T-11 nerve root sleeve (arrows).
Figure is available in color online only.
who have not obtained satisfactory relief with more conservative management and have an identifiable anatomical abnormality on imaging studies. Diverticula are ligated and dural tears are repaired in the fashion most customary to the surgeon. Outcomes have been found to be positive with few complications. In our patient, posterior exposure of the thoracic nerve root allowed for ligation of the entire nerve root, leading to complete and lasting resolution of orthostatic symptoms. The headaches that persisted in the subacute postoperative period are likely unrelated to the CSF leak, as they were nonpositional, decreased in severity, and were of different quality from her presenting complaint.

**Conclusions**

Thoracic nerve root redundancy may represent an undiagnosed etiology of spontaneous CSF leak. CT myelography is insufficient for visualizing these anatomical anomalies. Treatment with surgical ligation proved a viable option for management when conservative measures failed.
References

Disclosures
The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

Author Contributions
Conception and design: Dahdaleh, Lopez, Arnaout, Curran. Acquisition of data: Dahdaleh, Arnaout, Shaihani. Analysis and interpretation of data: Dahdaleh, Lopez, Arnaout, Curran, Shaihani. Drafting the article: Lopez, Campbell. Critically revising the article: all authors. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Dahdaleh. Study supervision: Dahdaleh.

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