A tethered cord (TC) can be caused by an abnormally tight filum terminale and result in traction on the spinal cord and clinical symptomatology of TC syndrome, including back or lower-extremity pain, sensory and motor abnormalities, musculoskeletal deformity, and voiding dysfunction. Radiological hallmarks can include a low-lying conus medullaris and/or an abnormally thickened or fatty filum. The definitive treatment for symptomatic patients with filum tethering is surgical transection.

Cutaneous findings, including large raised hemangioma and sacral dimples, are associated with a higher risk for the presence of a TC. The incidence of a TC in patients with a neurocutaneous syndrome is not known. Neurofibromatosis type 1 (NF1) and NF2 are autosomal dominant neurocutaneous syndromes. NF1 is associated with spinal skeletal deformity but not with occult spinal dysraphism or spinal cord tethering. Based on observations of multiple patients with NF, abnormalities of the filum terminale, and cord tethering, we hypothesized that NF might be associated with an increased incidence of TC. The Division of Pediatric Neurosurgery at Oregon Health & Science University (OHSU) serves as a major tertiary referral center at Oregon’s only academic health center. A broad range of pediatric neurosurgical procedures are performed in this division (which currently employs 3 pediatric neurosurgeons); the volume in 2017 was 780 major operating room–based procedures and 800 clinic-based new patient referrals.

Incidence of symptomatic tethered spinal cord in pediatric patients presenting with neurofibromatosis types 1 and 2

Carolyn S. Quinsey, MD, Katie Krause, MD, PhD, Lissa C. Baird, MD, Christina M. Sayama, MD, MPH, and Nathan R. Selden, MD, PhD

Department of Neurological Surgery and Doernbecher Children’s Hospital, Oregon Health & Science University, Portland, Oregon

OBJECTIVE The relationship between a tethered cord (TC) and neurofibromatosis type 1 (NF1) and NF2 is not known. The purpose of this study was to define the incidence of TC in pediatric neurosurgical patients who present with NF.

METHODS The authors performed a single-institution (tertiary care pediatric hospital) 10-year retrospective analysis of patients who were diagnosed with or who underwent surgery for a TC and/or NF. Clinical and radiological characteristics were analyzed, as was histopathology.

RESULTS A total of 424 patients underwent surgery for a TC during the study period, and 67 patients with NF were seen in the pediatric neurosurgery clinic. Of these 67 patients, 9 (13%) were diagnosed with a TC, and filum lysis surgery was recommended. Among the 9 patients with NF recommended for TC-release surgery, 4 (44%) were female, the mean age was 8 years (range 4–14 years), the conus position ranged from L1–2 to L-3, and 3 (33%) had a filum lipoma, defined as high signal intensity on T1-weighted MR images. All 9 of these patients presented with neuroromotor, skeletal, voiding, and/or pain-related symptoms. Histopathological examination consistently revealed dense fibroconnective tissue and blood vessels.

CONCLUSIONS Despite the lack of any known pathophysiological relationship between NF and TC, the incidence of a symptomatic TC in patients with NF1 and NF2 who presented for any reason to this tertiary care pediatric neurosurgery clinic was 13%. Counseling patients and families regarding TC symptomatology might be indicated in this patient population.
Here, we report our institutional (tertiary care pediatric hospital) experience and review the clinical presentation, radiological findings, and histopathological features of pediatric patients with NF and a symptomatic TC, and we discuss the association between NF and symptomatic TC that we found. We also define the incidence of TC in pediatric patients with NF to better inform patients and their families and clinicians who treat such patients.

Methods

In accordance with an institutional review board–approved protocol, with a waiver of consent, electronic medical records were reviewed retrospectively to determine the occurrence of TC surgery over a 10-year span (2006–2015). We also searched electronic medical records over the same time period to identify pediatric patients (< 18 years of age) with an International Classification of Diseases, Ninth Revision (ICD-9) code for NF1 or NF2 who were seen at any OHSU facility. Then, we cross-referenced the records of patients who underwent surgical treatment for a TC, NF1, and/or NF2 with the records of patient visits in the pediatric neurosurgery outpatient clinic during the same time period.

For identified patients with NF, the following information was collected: age; sex; NF type; presenting complaints; results of musculoskeletal, urological, neurological, and gait examinations; pain status; presence of scoliosis; conus medullaris level; presence of a syrinx; presence of a filum lipoma or other tethering element; and operations performed for detethering, along with operative and pathological findings, complications (if relevant), and postoperative course. Additional data were collected for each patient with a diagnosis of NF1 or NF2 who underwent TC-release surgery and/or who had an ICD-9 diagnosis of TC, including the results of urodynamic testing if performed.

The exact position of the conus medullaris was determined by reviewing radiology reports and through direct review of axial and sagittal T1- and T2-weighted MR images by a board-certified pediatric neurosurgeon (N.R.S., L.C.B., or C.M.S.) (Fig. 1). The presence of a filum lipoma was determined by high signal intensity on multiple T1-weighted axial MR images through the filum terminale and confirmation on the sagittal midline image. The presence of a syrinx was determined by the finding of T2-weighted hyperintensity within the central spinal cord parenchyma on axial and sagittal images.

For the purposes of this investigation, we focused on lower-extremity motor abnormalities and toe walking or other forms of abnormal gait or a regression in lower-extremity coordination over time. The musculoskeletal evaluations included an assessment of Achilles tendon tension, leg-length, and calf and foot-size discrepancy. Urological evaluations included obtaining a history of any secondary or age-inappropriate incontinence and, in some cases, performing formal urodynamic studies.

An English-language PubMed literature search was undertaken to find any previous reports of the co-occurrence of spinal cord tethering and NF1 or NF2 (search terms included “tethered cord neurofibromatosis” and “neurofibromatosis lipoma”).

Results

During the study time period, 447 pediatric patients with a diagnosis of NF1 or NF2 were evaluated within an OHSU clinic; 67 of these patients were seen in the pediatric neurosurgery clinic and are the subject of this study. A total of 424 pediatric patients underwent transection of the filum terminale for spinal cord detethering during this period. Eight patients with NF underwent surgery for a TC. We also included 1 additional patient with NF with a symptomatic TC who was recommended to undergo surgery but instead elected expectant management with clinical follow-up. Thus, 9 (13%) of the 67 patients with NF seen in the pediatric neurosurgery clinic over a 10-year period presented with a symptomatic TC, and 8 (12%) underwent TC-release surgery. All the patients diagnosed with both NF and a TC were seen in the pediatric neurosurgery outpatient clinic.

Four (44%) of the 9 patients with NF and a sympt-
had both NF1 and NF2 according to review by OHSU ge
was 8 years (range 4–14 years); 8 patients had NF1 and 1
sue was found in 1 specimen.

Histopathological examination of the filum terminale
- fibroadipose tissue; fibroadipose tissue

All 9 patients presented with neuromotor, skeletal,
- voiding, and/or pain-related symptoms. Five patients were
- noted to have bladder symptoms, 3 had refractory pain, 2
- had a toe-walking gait, 3 had repressive neuromotor function
- and coordination, 1 had tight Achilles tendons, and 1
- had new ankle clonus and hyperreflexia. Two patients under
- underwent formal urodynamic testing; one had documented
detrusor dysfunction with abnormal electromyographic
activity, and the other had a large bladder capacity without
contraction.

Histopathological examination of the filum terminale
for the 8 patients who underwent surgery revealed dense
fibrous connective tissue and blood vessels; fibroadipose tissue
was found in 1 specimen.

A PubMed literature search revealed no previous re
ports of the co-occurrence of spinal cord tethering and NF1 or NF2.

In the 8 patients who underwent surgery, TC symptoms
improved to various degrees. Three patients experienced
complete resolution of their back pain and lower-extremity
symptoms, and 5 patients reported improvement in their
symptoms. The 1 patient who presented with new ankle
clonus and hyperreflexia experienced resolution of these
symptoms. Of the 5 patients who presented with urinary
dysfunction, 3 experienced complete resolution of their
symptoms, and 1 patient reported improved symptoms.
The patient who had a large bladder capacity without
contraction based on urodynamic testing did not report
improvement in urinary symptoms. One patient initially
experienced improvement in coordination and bladder
dysfunction but developed new bilateral lower-extremity
numbness 7 years later. Subsequent MR images revealed
retethering, and the patient underwent a repeat TC-release
surgery, which resolved his lower-extremity numbness.

Discussion

Although no pathophysiological relationship between
them is known, the results presented here suggest an as
sociation between spinal cord tethering and NF in the pedi
atric patients who presented to our tertiary care pediat
ric hospital. This represents the first report in the English
literature of an association between tethered spinal cord
and NF. We propose that providers consider a diagnosis of TC when evaluating patients with NF. We recommend
clinical screening by obtaining a history and performing
a physical examination for patients with NF to detect any
obvious symptoms and findings of a TC that, if present,
should prompt MRI of the total spine. It should be noted
that spinal MRI in patients with NF and spinal cord–re
lated symptomatology is already advisable for ruling in or
out a diagnosis of compressive neoplastic pathology.

Although 1 patient in this study with a TC was diag
osed with NF2, in addition to NF1, no patient with an
isolated diagnosis of NF2 had a TC, which implies that the
association reported here might be specific to NF1.

Although no pathophysiological mechanism linking
NF and TC is known, we presume that the neural crest
abnormalities caused by defects in the NF1 gene also in
crease the risk of filar abnormalities that result in a TC.3
NF1 is expressed variably, which suggests that undisclosed
specifics of the NF-related genetic error determine the risk
of a TC in each patient.

Limitations to this study include its retrospective na
ture and performance at a single institution. Given their
propensity to form spinal masses, patients with NF are

<table>
<thead>
<tr>
<th>Pt.</th>
<th>Age (yrs), Sex</th>
<th>Op Level</th>
<th>Additional Imaging Finding(s)</th>
<th>Clinical Symptom(s)</th>
<th>NF Type(s)</th>
<th>Histopathological Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5, F</td>
<td>L1–2</td>
<td>Syrinx, fattyofilum</td>
<td>Toe walking</td>
<td>1</td>
<td>Dense fibrous connective tissue</td>
</tr>
<tr>
<td>2</td>
<td>8, M</td>
<td>L2–3</td>
<td>None</td>
<td>Toe walking, bowel &amp; bladder dysfunction</td>
<td>1 &amp; 2</td>
<td>Dense fibrous connective tissue</td>
</tr>
<tr>
<td>3</td>
<td>4, M</td>
<td>L-2</td>
<td>None</td>
<td>Bladder dysfunction, progressive in-turning toes</td>
<td>1</td>
<td>Dense fibrous connective tissue</td>
</tr>
<tr>
<td>4</td>
<td>13, F</td>
<td>L-3</td>
<td>None</td>
<td>Bladder dysfunction, refractory pain</td>
<td>1</td>
<td>Dense fibrous connective tissue</td>
</tr>
<tr>
<td>5</td>
<td>7, M</td>
<td>L1–2</td>
<td>Fattyfilum</td>
<td>Tight Achilles tendons, regressive neuromotor function &amp; coordination</td>
<td>1</td>
<td>Dense fibrous connective tissue, fibroadipose tissue</td>
</tr>
<tr>
<td>6</td>
<td>12, F</td>
<td>Low-L2</td>
<td>Syrinx, scoliosis</td>
<td>Refractory pain &amp; numbness, bladder dysfunction</td>
<td>1</td>
<td>Dense fibrous connective tissue</td>
</tr>
<tr>
<td>7</td>
<td>5, M</td>
<td>L2–3</td>
<td>Syrinx, fatty filum</td>
<td>Regressive neuromotor function &amp; coordination</td>
<td>1</td>
<td>Not available</td>
</tr>
<tr>
<td>8</td>
<td>14, F</td>
<td>L2–3</td>
<td>Syrinx, scoliosis</td>
<td>Regressive neuromotor function &amp; coordination, bladder dysfunction, refractory pain</td>
<td>1</td>
<td>Dense fibrous connective tissue</td>
</tr>
<tr>
<td>9</td>
<td>7, M</td>
<td>L-3</td>
<td>None</td>
<td>Clonus &amp; in-turning toes, hyperreflexia</td>
<td>1</td>
<td>Dense fibrous connective tissue</td>
</tr>
</tbody>
</table>

Pt = patient.
more likely than those in the general population to undergo complete neuraxis imaging. For these reasons, spinal imaging in asymptomatic patients might not be beneficial. Although it is possible also that an increased incidence of TC is a generalized feature of a broader range of neurodevelopmental disorders, we did not seek relevant data to answer that question.

We considered the alternative hypothesis that TC and NF randomly co-occurred in these patients, who presented to a pediatric neurosurgical practice that is attuned to diagnosing both conditions. However, the literature suggests that the rate of such incidental discovery of radiological spinal cord tethering is less than 1.5%.4

Conclusions

Despite the lack of any known pathophysiological relationship between NF and TC, the incidence of a symptomatic TC in patients with NF presenting for any reason to a single tertiary care pediatric neurosurgery clinic was 13%. Counseling patients and families regarding TC symptomatology might be indicated in this patient population, as would total-spine MRI in symptomatic patients.

Acknowledgments

We thank Shirley McCartney, PhD, for editorial assistance.

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: Selden, Baird. Acquisition of data: Quinsey, Krause. Analysis and interpretation of data: all authors. Drafting the article: all authors. 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: Selden.

Correspondence

Nathan R. Selden: Oregon Health & Science University, Portland, OR. seldenn@ohsu.edu.