Spinal neurenteric cysts and their relation to more common aspects of occult spinal dysraphism

MICHAEL J. RAUZZINO, M.D., R. SHANE TUBBS, M.S., P.A.-C., EBEN ALEXANDER, III, M.D., PAUL A. GRABB, M.D., AND W. JERRY OAKES, M.D.

Peninsula Neurosurgical Associates, Hampton, Virginia; Brigham and Women's Hospital, Boston, Massachusetts; and Section of Pediatric Neurosurgery, Division of Neurological Surgery, Department of Surgery, University of Alabama at Birmingham, Alabama

Object. Neurenteric cysts are infrequently reported congenital abnormalities believed to be derived from an abnormal connection between the primitive endoderm and ectoderm. The authors report a series of 13 patients treated over a 50-year period.

Methods. Of the 13 patients, seven were female and six were male. Their ages at presentation ranged widely from 5 weeks to 52 years of age. Children presented more commonly with cutaneous stigmata of occult spinal dysraphism (OSD) whereas adults presented primarily with pain. Neurological deficit as a presenting symptom was less common in our series, a finding that reflects the slow growth of these lesions. In all but one patient some form of vertebral anomaly was associated with the cystic lesions, including two patients with Klippel–Feil abnormalities. There was a high incidence of associated forms of OSD including split cord malformation, lipoma, dermal sinus tract, and tethered spinal cord. In previous reports the authors have suggested that neurenteric cysts are more common in the cervical region and in a position ventral to the cord. In the present series these cysts most commonly occurred as intradural, extramedullary masses in the thoracolumbar region, situated dorsal to the spinal cord. The median follow-up period was 7.5 years, and postoperative outcome reflected a patient’s preoperative neurological status; in no patient was outcome worsened due to surgery.

Conclusions. Complete excision of the neurenteric cyst remains the treatment of choice, as subtotal excision is associated with recurrence.

KEY WORDS • neurenteric cyst • occult spinal dysraphism

Spinal neurenteric cysts are infrequently reported congenital abnormalities believed to be derived from an abnormal connection between the primitive endoderm and ectoderm during the 3rd week of life. Neurenteric cysts are not confined to the spinal column but may be found within the brain, mediastinum, abdomen, pelvis, or even in a subcutaneous location.1,3,16,17,28,48,50 These lesions are rare. Intraspinal neurenteric cysts represent 0.3 to 0.5% of all spinal “tumors.”35 They are, in fact, not tumors, which differentiates them from teratomas; instead, they are more similar to harmatomas—displaced nests of endodermally derived tissue. The terminology for these lesions is problematic, as they have been reported as a neurenteric cyst, enterogenous cyst, enteric cyst, gastrocytoma, dorsal enteric fistula, split notochord syndrome, and teratoid cyst.6,12,20,26,27,37,49 Part of the confusion in naming these cysts is that they are not uniform; likely they exist as a spectrum of lesions. The presentation may vary from a simple isolated intraspinal cyst with no other abnormalities to that of the so-called split notochord syndrome, as proposed by Bentley and Smith6 in which there may be multiple visceral and vertebral anomalies.

Neurenteric cysts are considered to be a form of OSD, as are the following entities: lipoma, lipomyelomeningocele, SCMs, meningocele manqué, dermal sinus tract inclusion cysts (dermoids and epidermoids), terminal syringohydromyelia, and myelocystoceles. A consistent association between neurenteric cysts and these other forms of OSD has not been reported. In analyzing our 13 cases, we were struck by both their fascinating case histories as well as the incredible diversity of associated findings (typically either other forms of OSD or the various cutaneous and orthopedic stigmata of OSD). We present these patients in detail, compare them with other large reported series, and discuss possible causes of intraspinal neurenteric cysts.

CASE MATERIAL AND METHODS

The patients in this series were treated over a 50-year period in large part by the senior author (W.J.O.). Patients for whom either operative data or histological results were incomplete were excluded from this study. Complete records for 13 patients formed the basis for this study.
RESULTS

Patient Demographics

The patients ranged in age from 5 weeks to 52 years. Patients most commonly presented either early in life (first two decades) or well into adulthood (fourth or fifth decades). The youngest patients typically presented with cutaneous stigmata indicative of OSD whereas the older patients presented with pain or neurological dysfunction. There was a slight female preponderance (1.2:1) which is in contrast to previous reports in which a male preponderance has been reported (Table 1).

Cyst Location

The cysts in this series were found in a variety of locations, all of which were consistent with the proposed classification of developmental posterior enteric remnants proposed by Bentley and Smith (Fig. 1). In relation to the spinal cord, the most common location was dorsal to the cord (either as an intradural extramedullary mass [four cases] or as an extradural dorsal mass [two cases]). The second most common location was intramedullary (four cases), followed by an anterior relationship to the cord (three cases). The cysts appeared in all regions of the spine, ranging from C-2 to L-5 (Table 1). We found no cysts at the craniocervical junction or in an intracranial location. The most common location was at the thoracolumbar junction (six cases), followed by the upper-thoracic region (four cases), cervical region (two cases), and, least commonly, the lower-lumbar spine (one case). In our series the location and position of the neurenteric cysts differed from the preferred location (that is, in the cervical spine with an anterior relationship to the cord) that has been previously reported by many authors.15,30,31,38,54

Neuroimaging Findings

Many of the patients presented prior to the advent of MR imaging. Computerized tomography myelography

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Age (yrs), Sex</th>
<th>Presentation</th>
<th>Cyst Location</th>
<th>Relation of Cyst to Spinal Cord</th>
<th>Level</th>
<th>Bone Abnormalities</th>
<th>Cutaneous Stigmata</th>
<th>Other Forms of OSD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>52 yrs, F</td>
<td>neck pain, rt arm numbness &amp; weakness</td>
<td>intradural, extramedullary</td>
<td>anterior</td>
<td>C2–C3</td>
<td>Klippel–Feil</td>
<td>none</td>
<td>none</td>
</tr>
<tr>
<td>2</td>
<td>17 yrs, M</td>
<td>pain, leg length discrepancy</td>
<td>intradural extramedullary</td>
<td>dorsal</td>
<td>T12–L2</td>
<td>scoliosis, spina bifida, T-11 hernivertebrae hole in C-3 VB, scoliosis</td>
<td>hairy patch</td>
<td>SCM, tethered cord</td>
</tr>
<tr>
<td>3</td>
<td>9 yrs, F</td>
<td>back pain, lt pain &amp; numbness</td>
<td>intradural, extramedullary</td>
<td>anterolateral</td>
<td>C2–3</td>
<td>none</td>
<td>none</td>
<td>none</td>
</tr>
<tr>
<td>4</td>
<td>23 yrs, M</td>
<td>back pain, progressive leg weakness</td>
<td>intradural, extramedullary</td>
<td>dorsal</td>
<td>T1–2</td>
<td>spina bifida</td>
<td>pedunculated mass</td>
<td>dorsal meningoclele</td>
</tr>
<tr>
<td>5</td>
<td>10 wks, M</td>
<td>LUQ mass, postmediastinal mass</td>
<td>intradural, extramedullary</td>
<td>anterior</td>
<td>T2–3</td>
<td>thoracic hemi &amp; butterfly vertebrae, scoliosis</td>
<td>none</td>
<td>none</td>
</tr>
<tr>
<td>6</td>
<td>4 yrs, M</td>
<td>back pain</td>
<td>intradural, extramedullary</td>
<td>dorsal</td>
<td>L1–3</td>
<td>spina bifida</td>
<td>cutaneous hemangioma</td>
<td>none</td>
</tr>
<tr>
<td>7</td>
<td>5 wks, F</td>
<td>lumbar mass</td>
<td>extradural (subcutaneous)</td>
<td>dorsal (subcutaneous)</td>
<td>L1–2</td>
<td>multiple spina bifida (L1–S1)</td>
<td>hairy patch, sacral mass &amp; dimple</td>
<td>adjacent intradural lipoma, nonadjacent subcutaneous lipoma tethered cord w/thickened filum, lipoma</td>
</tr>
<tr>
<td>8</td>
<td>8 yrs, M</td>
<td>back &amp; rt leg pain</td>
<td>intradural, intramedullary</td>
<td>intramedullary</td>
<td>L–5</td>
<td>spina bifida</td>
<td>dermal sinus tract, hairy patch, capillary hemangioma subcutaneous mass</td>
<td>intradural lipoma</td>
</tr>
<tr>
<td>9</td>
<td>1 yr, F</td>
<td>lumbar subcutaneous mass</td>
<td>subcutaneous</td>
<td>dorsal (subcutaneous)</td>
<td>T11–L2</td>
<td>spina bifida</td>
<td>subcutaneous mass</td>
<td>intradural lipoma</td>
</tr>
<tr>
<td>10</td>
<td>51 yrs, M</td>
<td>progressive pain &amp; numbness in legs</td>
<td>intradural, intramedullary</td>
<td>intramedullary</td>
<td>T2–3</td>
<td>spina bifida</td>
<td>subcutaneous mass</td>
<td>syringomyelia, dorsal tethering, SCM, intramedullary lipoma, tethered cord from thickened filum, meningoclele manqué</td>
</tr>
<tr>
<td>11</td>
<td>45 yrs, F</td>
<td>progressive paraparesis</td>
<td>intradural, extramedullary</td>
<td>dorsal</td>
<td>T4–7</td>
<td>spina bifida</td>
<td>hairy patch</td>
<td>SCM, intradural lipoma</td>
</tr>
<tr>
<td>12</td>
<td>19 yrs, F</td>
<td>mirror movements</td>
<td>intradural, intramedullary</td>
<td>intramedullary</td>
<td>T12–L2</td>
<td>Klippel–Feil</td>
<td>none</td>
<td>SCM, intradural lipoma</td>
</tr>
<tr>
<td>13</td>
<td>6 wks, F</td>
<td>hairy patch</td>
<td>intradural, intramedullary</td>
<td>intramedullary</td>
<td>T12–L2</td>
<td>none</td>
<td>hairy patch</td>
<td>SCM</td>
</tr>
</tbody>
</table>

* LUQ = left upper quadrant.
demonstrated the lesions typically as a complete or partial block at the appropriate levels, but often the diagnosis of neurenteric cyst was an unexpected finding at surgery. Other bone abnormalities such as diastematomyelia were well demonstrated on these imaging studies. In our later patients, MR imaging revealed the neurenteric cysts to be typically isointense to hyperintense relative to cerebrospinal fluid on long–relaxation time sequences. On T₂-weighted MR imaging they appeared isointense or slightly hyperintense to cerebrospinal fluid. Based on reports in the literature, these signal characteristics are typically and correlate with the high–protein content fluid within the cysts. Using MR imaging, were able to make a presumptive diagnosis of neurenteric cyst in most of these patients preoperatively.

**Histopathological Examination**

Neurenteric cysts are classified by the World Health Organization under the heading of “other malformative tumors and tumor-like lesions” and are described as cysts “lined by mucin secreting epithelium resembling that of the gastrointestinal tract.” Wilkins and Odum have classified these cysts into three types based on the histological features of the cyst wall and its contents. The walls of Type A cysts mimic gastrointestinal or respiratory epithelium with a basement membrane supporting single or pseudostratified cuboidal or columnar cells, which may be ciliated. Type B cysts also contain glandular organization, usually producing mucin or serous fluid. Type C cysts are the most complex containing ependymal or glial tissue within the cyst. In our series, one patient (8%) harbored a simple Type A cyst, 10 patients (77%) harbored the moderately complex Type B cyst, and two patients (15%) harbored the most complex Type C cyst (Table 2). It has been reported that neurenteric cysts with associated abnormalities are of greater histological complexity as compared with those cysts without associated dysraphic features. Our findings support this association. Paleologos, et al., compared the pathological features of 136 reported cases of neurenteric cysts and divided them into two groups: 80 patients with neurenteric cysts without associated dysraphism and 56 patients harboring neurenteric cysts with associated dysraphism. In the nondysraphic group 84% of

**TABLE 2**

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Histopathological Finding</th>
<th>Wilkins &amp; Odum Classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>single-layer columnar epithelial cells w/ basement membrane &amp; microvilli</td>
<td>A</td>
</tr>
<tr>
<td>2</td>
<td>single-layer columnar cells w/ mucin droplets; smooth-muscle cells &amp; glial tissue</td>
<td>C</td>
</tr>
<tr>
<td>3</td>
<td>columnar epithelium w/ mucin glands (PAS positive)</td>
<td>B</td>
</tr>
<tr>
<td>4</td>
<td>columnar epithelium w/ mucin glands</td>
<td>B</td>
</tr>
<tr>
<td>5</td>
<td>columnar &amp; squamous epithelium w/ mucus glands</td>
<td>B</td>
</tr>
<tr>
<td>6</td>
<td>ciliated columnar to flattened epithelium w/ mucin droplets &amp; mucin glands</td>
<td>B</td>
</tr>
<tr>
<td>7</td>
<td>columnar epithelium w/ mucin glands similar to respiratory glands</td>
<td>B</td>
</tr>
<tr>
<td>8</td>
<td>columnar epithelium w/ smooth-muscle cells, cartilage (PAS positive)</td>
<td>B</td>
</tr>
<tr>
<td>9</td>
<td>single-layer columnar epithelial cells w/ basement membrane; PAS-positive glands resembling GI tissue</td>
<td>B</td>
</tr>
<tr>
<td>10</td>
<td>cuboidal epithelium w/ mucin glands &amp; goblet cells</td>
<td>B</td>
</tr>
<tr>
<td>11</td>
<td>columnar epithelium w/ mucin glands resembling GI tissue</td>
<td>B</td>
</tr>
<tr>
<td>12</td>
<td>columnar epithelium w/ mucin glands</td>
<td>B</td>
</tr>
<tr>
<td>13</td>
<td>single-layer columnar epithelial cells resembling GI &amp; respiratory-like mucosa; mucin glands, smooth-muscle cells, cartilage, &amp; glial tissue also noted</td>
<td>C</td>
</tr>
</tbody>
</table>

*GI = gastrointestinal; PAS = periodic-acid Schiff.
patients harbored the simplest Type A cyst, and in the dysraphic group only 39% had a similar lesion type.

**Patient Outcome**

The relatively benign nature of these slow-growing lesions is evident by the good outcome experienced by most patients in this series (Table 3). Postoperative neurological status typically reflected the preoperative status, and no patients experienced worsened symptoms after undergoing surgery. There was a cyst recurrence rate of 27% in our series after the cyst had been incompletely resected (either by us or by a previous surgeon). These recurrences occurred early in our series because, in the later part of the series, a significant attempt was made to resect the entire cyst wall if the intraoperative diagnosis of neurenteric cyst was considered and if this was technically possible without causing significant neurological deficit.

**Associated Abnormalities**

**Vertebral Abnormalities.** All but one of these patients (92%) presented with some form of vertebral abnormality, often at multiple levels. Anterior vertebral column abnormalities were seen in five patients (39%) including two patients (15%) with Klippel–Feil abnormalities. Posterior vertebral column abnormalities were found in eight patients (62%), most typically as some form of spina bifida. Scoliosis-related spinal deformity was present in two patients (15%) with the scoliosis resulting from thoracic hemivertebrae.

**Cutaneous Stigmata.** A number of cutaneous stigmata have been associated with OSD, but rarely has this association been made with neurenteric cysts. Nine (69%) of our 13 patients had some form of cutaneous stigmata, and some had multiple stigmata. The cutaneous stigmata included focal hirsutism (five cases), subcutaneous mass (three cases), flat capillary hemangioma (two cases), dermal sinus tract (two cases), and pedunculated mass (one case). In the three youngest patients in this series, it was the presence of this cutaneous stigmata that led to the early diagnosis of the neurenteric cyst.

**Associated Forms of Spinal Dysraphism**

Nine patients (69%) harbored a neurenteric cyst accompanied by another form of spinal dysraphism in which three were intraspinal abnormalities. Four patients (31%) harbored a cyst in which there were no other intraspinal abnormalities. The most common associated findings were SCM (31%) or associated intradural lipoma (31%). A tethered spinal cord caused by a thickened terminal filum was found in three patients (23%). There was also one case each of a dorsal meningocele, syringomyelia, and meningocele manqué (8%, respectively).

**ILLUSTRATIVE CASES**

**Case 1**

This 52-year-old woman presented with a 4-year history of neck and right arm pain, which progressed over several months to include numbness and weakness. Cervical myelography demonstrated an anterior intradural extramedullary mass at the C2–3 level. Plain radiography revealed a congenital fusion of the C-2 and C-3 VBs. A cervical laminectomy was performed, and at surgery, the spinal cord was shown to be displaced posteriorly by a cystic mass from which clear fluid was aspirated. The wall of the cyst consisted primarily of a single layer of columnar epithelium with some ciliation resting on a thick basement membrane. Postoperatively, electron microscopy showed signs of mucinous granules, microvilli, and nuclei consistent with an intestinal origin. The patient awoke without deficit and had relief of her neck and arm pain.

**Case 2**

This 17-year-old man presented with a 2-year history of worsening low-back pain. He was noted to have scoliosis
Spinal neurenteric cysts

Case 3
This 9-year-old child with a progressive congenital cervicothoracic scoliosis presented with a 1-year history of neck pain that progressed to torticollis. Left leg pain and weakness developed and a cervical myelogram revealed an anterior intradural extramedullary mass. A cervical laminectomy was performed, and at surgery the spinal cord was found to be displaced posteriorly by a cystic mass, which was intradural and anterolateral to the cord on the right. The cyst was opened, and a thick mucous component was noted. A circular opening in the anterior dura was observed just to the right of the midline at C-3 with tissue continuing into the VB. The tissue was ligated. Histopathological examination showed columnar epithelium with mucinous glands. Postoperatively the patient experienced resolution of left leg weakness. At the 5-year follow-up review, the patient was asymptomatic.

Case 4
This 23-year-old man presented with spina bifida of the upper-thoracic spine. At age 15 years he had developed neck and shoulder pain, which was aggravated by coughing. Physical examination showed a small pedunculated mass in the midline at T-2. He was then believed to have a small dorsal bony spina bifida with a meningocele and the cord fixated dorsally. Two years later he developed left arm pain and atactic gait with bilateral leg weakness. He declined surgery at the time. He presented at age 30 years with a progressive upper-thoracic scoliosis, increased leg weakness, and urinary hesitancy. He underwent a T1–2 laminectomy in which a large intradural dermoid cyst was removed and a dorsal meningocele was excised and repaired. The stalk of the meningocele contained clear fluid that passed through a bified T-1 lamina. A dorsally located, encapsulated mass containing grayish gelatinoid material extended from C-6 to T-2 and was partially excised. The patient presented 10 years later with a 3-day history of progressive paraparesis and a C-8 sensory level. A myelogram demonstrated a complete block at T-2, and he underwent a C7–T3 laminectomy. A thin-walled extramedullary cyst containing mucus was totally excised. The cyst wall was lined by columnar epithelium with underlying mucous glands. The patient experienced improvement of his lower-extremity strength, but a spastic gait, hyperreflexia, and diminished position sense in his legs remained.

Case 5
This 10-week-old boy was noted to have a left upper-quadrant mass that enlarged in the first few weeks of life. His neurological status was normal, and CT scanning demonstrated a smooth cystic mass in the left upper quadrant and a smaller paravertebral mass in the posterior mediastinum. Plain radiographs of the chest demonstrated multiple thoracic hemi- and butterfly vertebrae. Exploratory laparotomy revealed an enteric duplication cyst. The posterior mediastinal cyst was removed by thoracotomy, and a connection was seen attached to the T-4 VB. A thoracic laminectomy was performed, and an intradural extramedullary cyst that displaced the cord dorsally and laterally, was removed. The cyst wall was noted to have columnar and squamous epithelium with mucinous glands. The patient remained neurologically intact.

Case 6
This 4-year-old boy presented with a 1-year history of back pain. Neurological examination demonstrated normal findings, but the patient was noted to have a capillary hemangioma over the lumbar spine. Plain radiography demonstrated a bifikd L-5 vertebra and an expanded lumbar canal. Computerized tomography myelography revealed an intradural extramedullary tumor at L-1 and a mass posterior to the cord. A T12–L4 laminectomy was performed, and a midline intradural cystic tumor was found that seemed to arise from the cord and blend with it laterally. Histologically, the cyst consisted of ciliated columnar to flattened epithelium with mucin droplets as well as mucous glands. The patient underwent a repeated subtotal resection 1 year later and eventually at age 10 years required a third operation in which complete excision of the cyst was achieved. He has remained neurologically intact and free from recurrence at 30 years of age.

Case 7
This 5 1/2-week-old girl was noted at birth to have a sacral mass and dimple with a hairy patch (Fig. 3 left). Plain radiography demonstrated multiple posterior arch defects, and CT myelography demonstrated two areas of malfor-
The first was at the L1–3 level where, in the subcutaneous tissue, a water-density cyst was observed that had a stalk extending through the lumbodorsal fascia. Additionally, at the lumbodorsal fascia two posterior fat density masses were noted to compress the cord. The second area was at L5–S1 where the neural arches were disrupted and bone compression of the thecal sac had developed. An extradural fatty tumor was also noted at this level. At surgery a subcutaneous bluish cystic structure, yielding milky mucinous fluid, narrowed to a thickened fibrous structure that pierced the dura and attached dorsally to the cord at L1–2. An intradural exploration revealed two lipomatous masses lying on either side of the cord. The dermal pit and hair tuft at L5–S1 was found to overlie a subcutaneous lipoma that entered the dura at the L-5 level. Histological examination of the subcutaneous cyst showed columnar epithelium with glandular structures containing apical mucin similar to respiratory tract epithelium. At the 7-year follow-up examination, the patient remained neurologically intact and experienced no recurrence of lesions.

Case 8
This 8-year-old boy presented with a 2-year history of neck, back, and right leg pain. At birth he had been noted to have a dermal sinus tract in the lumbar region surrounded by a capillary hemangioma and a hair tuft. On plain x-ray films obtained at that time, an L-5 spina bifida was observed, and focal swelling of the conus medullaris was demonstrated on air myelography. He underwent surgical sectioning of the filum and was then lost to follow up. He presented at 8 years of age with recurrent back pain and incontinence precipitated by coughing. Repeated CT myelography demonstrated focal swelling in the conus medullaris extending over three vertebral segments. An L1–4 laminectomy was performed, and on opening the dura, the dorsal surface of the conus medullaris was replaced by a cystic swelling (Fig. 4). During removal of the cyst, thick mucinous fluid was spilled. Histological examination of the cyst wall showed a columnar epithelium and evidence of smooth-muscle cells and cartilage. The tissue sample stained periodic-acid Schiff positive, and the lesion was diagnosed as a neurenteric cyst. At the 12-year follow-up examination, the patient was asymptomatic except for occasional pains shooting down the right leg.

Case 9
This 1-year-old girl had undergone an attempted excision of a subcutaneous lumbar mass; however, the general surgeon had aborted the procedure when the cyst was discovered to have a stalk that pierced the lumbodorsal fascia. The tract was tagged and the wound closed. Subsequent MR imaging revealing a thoracic spina bifida and what was believed to be an intradural, extramedullary mass in association with the extradural neurenteric cyst. At the second surgery, the remainder of the subcutaneous cyst was seen to pass through the L-1 spina bifida where a sizable portion of the cyst extradurally compressed the thecal sac. On opening the dura, a sizable lipomatous mass was observed to be adherent to the dorsal surface of the spinal cord. Histopathological examination of the cyst wall showed single-layer columnar epithelial cells with a basement membrane and periodic-acid Schiff–positive glands resembling gastrointestinal tissue. The patient awoke neurologically intact.

Case 10
This 51-year-old man presented with a 3-month history of neck pain, and a burning sensation and numbness in his lower extremities. He had an interesting history. As an infant spina bifida had been diagnosed, and a mass was excised. The patient underwent an attempted excision of a subcutaneous lumbar mass; however, the general surgeon had aborted the procedure when the cyst was discovered to have a stalk that pierced the lumbodorsal fascia. The tract was tagged and the wound closed. Subsequent MR imaging revealed a thoracic spina bifida and what was believed to be an intradural, extramedullary mass in association with the extradural neurenteric cyst. At the second surgery, the remainder of the subcutaneous cyst was seen to pass through the L-1 spina bifida where a sizable portion of the cyst extradurally compressed the thecal sac. On opening the dura, a sizable lipomatous mass was observed to be adherent to the dorsal surface of the spinal cord. Histopathological examination of the cyst wall showed single-layer columnar epithelial cells with a basement membrane and periodic-acid Schiff–positive glands resembling gastrointestinal tissue. The patient awoke neurologically intact.
Spinal neurenteric cysts

cised from his interscapular region. At presentation his neurological status was normal except for subtle long tract signs and mildly ataxic gait. Preoperative MR imaging revealed a previous thoracic laminectomy with a dermal sinus tract leading to an intramedullary cystic mass and dorsal adhesion of the cord (Fig. 5). Additionally a non-contiguous syrinx was observed in the upper-cervical region. At surgery, the dura was reopened, and in the midline, a large intramedullary cystic mass within the central portion of the cord was easily visualized. Mucous was aspirated from the cyst and an attempt was made to excise the cyst wall. The spinal cord was untethered from the dorsal adhesions, and the syrinx above was shunted into the subarachnoid space. Histopathological examination of the cyst wall showed cuboidal epithelium with mucin glands and goblet-shaped cells. The patient awoke without additional deficit and experienced relief of his pain. His bilateral lower-extremity spasticity persisted. Within 2 months his dysesthetic pain had returned.

Case 11

This 45-year-old woman presented with increasing paraparesis. She had undergone a spine operation as a 6-week-old infant, but the operative report and pathological results were not available for review. She did not present again until age 37 years when she developed an acute paraparesis after a minor traumatic injury. She underwent a second operation for removal of what was described as an intramedullary teratoma. She recovered but over the next 9 years developed increasingly severe back pain, lower-extremity weakness, and bladder dysfunction. Preoperative MR imaging demonstrated two areas of SCM (one bone septum and one fibrous septum), an intramedullary cyst, an intramedullary mass consistent with lipoma, and a tethered cord from a thickened terminal filum. At surgery the neurenteric cyst was noted to arise dorsal to the two hemicords, and it contained characteristic mucous material. A number of dorsal tethering bands (meningocele manqué) were found and sectioned in addition to the filum. The intramedullary lipoma above the cyst was partially debulked. Histopathological examination showed columnar epithelium with mucin glands that resembled gastrointestinal tissue. The patient awoke without new deficit. Her leg weakness and bladder dysfunction were improved but not normalized.

Case 12

This 19-year-old woman presented with neck pain that was found to be associated with an extensive fusion process of her cervical spine. At presentation, she was noted to have extensive mirror movements as part of her Klippel–Feil syndrome. A brain and cervical spine MR imaging study demonstrated medullary diaschisis and the aforementioned bone changes but no cervical spinal cord disease. As part of her workup, a total spine MR imaging study was ordered, and unexpectedly, an incidental finding was made of a major abnormality of the conus medullaris. This included an SCM with a cystic mass distorting the left hemicord, fatty tissue between the two hemicords, and tethering of the cord at this level (Fig. 6). After a long discussion with the family, it was thought that the natural history of this lesion could represent leg weakness, and the family desired the patient undergo excision of the mass. At surgery via a posterior laminectomy, the cystic mass in the left hemicord yielded thick mucinous fluid. The intramedullary cyst wall was only partially excised. There was a calcified spur separating the two hemicords and a dorsally located intradural lipoma, which was resected. Histopathological examination of the cyst wall showed a single layer of columnar epithelium with a large number of mucin glands. The patient awoke without deficit and remains well now 6 years postoperatively. The cyst has gradually refilled as documented by MR imaging.

Case 13

This 6-week-old girl presented with a hairy patch over the thoracolumbar junction. Her family history was remarkable for having a sibling and a cousin born with a myelomeningocele. An MR imaging study demonstrated an intradural dumbbell-shaped mass that appeared to arise from the dorsum of the cord. A posterior laminectomy was performed, and the dura was opened in the midline to reveal a cystic lesion. The cyst was aspirated, yielding

Fig. 5. Case 10. Midsagittal T2-weighted MR image of intramedullary cyst with a dermal sinus tract.

Fig. 6. Case 12. Axial T2-weighted MR image of neurenteric cyst arising from a hemicord.
mucinous material, and unexpectedly, it became obvious that there were two distinct hemicords between which the tumor arose. Histopathological examination showed a complex cystic lesion in which there were single-layer columnar epithelial cells with a basement membrane. Both gastrointestinal- and respiratory-like mucosa were noted with mucin glands and some smooth-muscle cells and cartilage. The patient awoke without deficit and is healthy now at 7 years of age.

**DISCUSSION**

**Embryological Development**

A number of theories have been proposed to explain the occurrence of endodermal tissue in the central nervous system. A knowledge of early embryological development is necessary to understand these theories. Any theory would have to allow not only for the formation of the cyst but also account for the myriad abnormalities that often accompany the cyst. As bone defects are commonly associated with these cysts, it would seem likely that the primary event would occur within the first 3 weeks of gestation during gastrulation when these endodermal and mesenchymal layers are intimately associated with each other. Initially, the fertilized embryo consists of a bilaminar disc containing layers of endoderm and ectoderm. A groove, termed the primitive streak, then forms on the caudal portion of the ectodermal surface and terminates as an elevation known as the Hensen node. Mesodermal cells migrate through the primitive streak and continue in lateral and cephalic directions to form the notochord of the trilaminar germ disc by the end of the 3 week of gestation. There is a connection called the neuromeric canal that is usually transient and connects the amniotic cavity and the yolk sac through the primitive pit. A tube of mesoderm condenses to form the notochord and paraxial mesoderm cells migrate around the notochord as precursors to the future vertebral column. The notochord also induces the overlying ectoderm to form the neural plate, which will eventually fold on itself, and close during the 4th gestational week to form the neural tube.

The main theories for neurenteric cyst formation include: 1) a primary adhesion of endoderm anterior to the notochord, 2) incomplete excalation of the notochord, 3) persistence of the neurenteric canal or formation of an accessory neurenteric canal with a split notochord, and 4) displacement of endodermal cells (Fig. 7).

Beardmore and Wigglesworth proposed that during the outgrowth of the notochord an adhesion could form between the endoderm and ectoderm that would become a barrier to the elongating notochord and thereby splitting it. The paired neuroepithelial precursors could then develop independently, forming two hemicords, and the endodermal remnants of the adhesion could then be found anywhere between the gut and the surface ectoderm.

Rhaney and Barclay suggested that aberrant intercalation or excalation of the notochord could lead to strands of endodermal remnants that could traverse a secondarily split notochord.

Bremer postulated that neurenteric cysts and diastematomyelia could be explained by the persistence of a neurenteric canal. He noted that because most neurenteric cysts are located more rostrally in the spine and the neurenteric canal is ultimately destined for the caudal end of the spine, an accessory canal is more likely to be the cau-
Spinal neurenteric cysts

ative factor than persistence of the original neurenteric canal. As part of a unified theory of SCMs, Pang, et al. have proposed that a secondary neurenteric tract may develop and become invested with mesenchymal tissue that splits the notochord and neural plate. Depending on the timing of such an event, two hemicords might develop with a cleft between them that could go on to form a bone spur with the potential to tether the cord. This theory would explain the association of SCMs with neurenteric cysts demonstrated in our series.

Bentley and Smith expanded Bremer’s theory, postulating that the splitting of the notochord is the primary event. The subsequent deficiency in the overlying neural plate could allow for an endodermal diverticulum to herniate through the spinal column and make contact with the surface ectoderm. In such a situation the persistence of the neurenteric connection which occurs may be transient or permanent, partial or complete (Fig. 1).

A number of scenarios may potentially result from this scheme. 1) If only the ventral portion remains, then a duplication of the alimentary canal without associated spinal abnormalities can occur. 2) If, on the other hand, only the terminal dorsal portion persists, then a cutaneous abnormality possibly associated with a sinus might result. 3) If the midportion only persists and if the cleft in the VB is incompletely obliterated, then anterior spina bifida and possibly diastematomyelia may result. If the cleft in the VB is transient and incomplete, then an intraspinal cyst can occur in the absence of permanent anterior bone anomalies. 4) Any combination of the aforementioned three scenarios may occur. This split notochord theory is attractive because it provides an easily understandable mechanism that can account for the entire spectrum of lesions in our series, including the presence of dorsally located lesions. However, this theory does not explain the formation of neurenteric cysts in their entirety. There have been well-documented cases of neurenteric cysts in an intracranial location, and these lesions would not be predicted by this theory. It is possible, then, that neurenteric cysts may occur by all or none of the previously mentioned theories.

Comparison With Previous Reports

A neurenteric cyst is classically reported as a solitary lesion in the cervical region, located anterior or anterolateral to the cord. Bone abnormalities, if present, are likely to involve the anterior column. Wilkins and Odum initially reviewed 47 patients with reported neurenteric cysts and later Wilkins and Rossitch added another 66 patients to this group to yield a review of 113 reported cases of neurenteric cysts. The most common reported location was cervical (63 cases), followed by thoracic (53 cases) and lumbosacral (27 cases). Junctional cysts were reported in both locations. There was a slight male predominance of 1.8:1 with the most common location of the cyst being intradural, extramedullary, and anterior or anterolateral to the cord. Although they found some case reports in their literature review of neurenteric cysts in association with other forms of OSD, the combination was infrequent.

Our series of patients clearly differs from this profile. The cysts were most commonly located dorsal to the cord in the thoracolumbar region. In our series the presence of a neurenteric cyst was associated with another form of OSD in 69% of the cases. At least some form of abnormality accompanied the cyst in 92% of our patients. In other reviewed series, neurenteric cysts were associated with abnormalities in 12% (Arai, et al.), 30% (Agnoli, et al.), 43% (Wilkins and Odum), and 54% (Holmes, et al.) of cases. In no previously reported series has the association between neurenteric cysts and other forms of OSD been so marked. The series most similar to ours was reported by Holmes, et al., in which they reviewed 26 pediatric patients in whom they noted a high percentage of associated abnormalities. Interestingly, they also reported that 70% of their cysts occupied a dorsal location to the cord, similar to our series. It is possible that the association of neurenteric cysts with spinal dysraphism may simply be underreported. Pang has noted that in his unreported series of neurenteric cysts, occurrence of associated SCMs are presented in 60% of the cases. Hoffman has similarly noted that in his unreported series of neurenteric cysts there is a nearly universal association with spinal dysraphism.

Treatment Recommendations

The surgical management of neurenteric cysts follows the general guidelines for other forms of OSD. Magnetic resonance imaging is helpful in predicting the complex anatomy of the cysts and their associated lesions. Anteriorly located cysts were well managed in this series via a posterior approach, but when feasible, an anterior approach to these lesions should be considered. Our 27% recurrence rate would suggest that the natural history of incompletely resected cysts is unfavorable and every attempt should be made to achieve a complete resection of the cyst and its wall at initial operation when technically feasible.

References


Manuscript received December 4, 2000. Accepted in final form December 20, 2000. Address reprint requests to: Michael J. Rauzzino, M.D., Peninsular Neurosurgical Associates, 2102 Executive Drive, Hampton, Virginia 23666. email: mrauzzino@aol.com.