Closed Spinal Dysraphism: Analysis of Clinical, Radiological, and Surgical Findings in 104 Consecutive Patients

James H. Scatlliff, Brian E. Kendall, Derek P. E. Kingsley, Juliet Britton, D. Norman Grant and Richard D. Hayward

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Closed Spinal Dysraphism: Analysis of Clinical, Radiological, and Surgical Findings in 104 Consecutive Patients

We reviewed 104 consecutive cases of closed dysraphism in patients seen at one institution between December 1984 and June 1987. All patients had myelographic studies, and 43 had associated CT examinations. Clinical and surgical findings (64 patients) were correlated with myelographic information. Twenty-three patients (22%) with clinical or plain film findings compatible with dysraphism had normal-appearing cords on conventional myelography, movement between supine and prone positions, and no lesions in the spinal canal. Cerebellar tonsillar ectopia (majority of tonsils between foramen magnum and C1) was found in 17 patients (16%). Six patients (6%) exhibited varying degrees of hydromyelia. In the supine position, CT-myelography of meningoceles, meningomyeloceles, or lipomeningomyeloceles may limit demonstration of the neural placode and nerve roots because of compression of the CSF-containing sac. In the decubitus position, CT scans improved demonstration of neural tissue–CSF space relationships. CT scans were useful in demonstrating anomalous paraspinal bones, diastematomyelia spurs, and spinal and sacral bone deficiency. Axial CT-myelography of intradural lipomas showed apparent neural tissue extension into the lipomas.

We reviewed 104 cases of suspected or proved spinal dysraphism seen at the Hospital for Sick Children, Great Ormond Street, London, from December 1984 through June 1987. Surgical and pathologic data were correlated with clinical and diagnostic imaging findings. Although a considerable number of radiologic reports describing various expressions of spina bifida occulta have appeared in the last 20 years, the assessment of a large number of cases from one hospital is somewhat unique [1–5]. There were 39 postoperative myelograms in 32 patients. (The findings in these studies and their evaluation will be presented in another article.)

Materials and Methods

In most cases, imaging studies were performed while the patients were under general anesthesia. Only five patients over the age of 10 were given a local lumbar anesthetic. Plain films were made before or at the beginning of the myelographic procedure. Lumbar punctures with a 22-gauge needle were done off the midline to avoid a low cord or suspected intraspinal component of a lipoma. Iohexol,* 240 mg/ml, was injected in volumes of 5–15 ml. The great majority of injections were initially subarachnoid, although several mixed (subarachnoid-subdural) injections were made. There was one inadvertent injection into a lumbodorsal hydromyelia. Posteroanterior and lateral films made at right angles during the myelogram showed the needle to be within 2 or 3 mm of the cord or hemicord in most of the studies. In several patients the needle appeared to penetrate or distort the cord. In no study was there evidence of hematomyelia, and there was never any subsequent increase in neurologic findings.

After prone filming and withdrawal of the needle, supine anteroposterior and horizontal-beam lateral films were made in the lumbar region to assess cord movement. In the supine position it was assumed that the cord was not tethered when it floated anteriorly 5 mm or more in the contrast material. Supine filming produced more complete filling of the caudal

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* Nyegaard, Oslo, Norway.
Dermal

D 
edroid
tethered cord

Atretic

Diastematomyelia:

Normal Myelogram

Other patients.

TABLE 1: Diagnostic Studies for Spinal Dysraphism Between December 1984–June 1987 (n = 104)

<table>
<thead>
<tr>
<th>Finding</th>
<th>Initial Myelograms</th>
<th>Preoperative Myelograms-CT</th>
<th>Postoperative Myelograms</th>
<th>Postoperative Myelograms-CT</th>
<th>Tonsillar Ectopia</th>
<th>Hydromyelia</th>
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<tr>
<td>Normal</td>
<td>23</td>
<td>1</td>
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<td></td>
<td></td>
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<tr>
<td>Tethered cord</td>
<td>14</td>
<td>3</td>
<td>4</td>
<td>3</td>
<td>1</td>
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</tr>
<tr>
<td>Lipoma</td>
<td>10</td>
<td>8</td>
<td>7</td>
<td>2</td>
<td></td>
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<tr>
<td>Lipomyelomeningocele</td>
<td>16</td>
<td>10</td>
<td>6</td>
<td>2</td>
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<tr>
<td>Meningomyelomeningocele</td>
<td>7</td>
<td>3</td>
<td>4</td>
<td>1</td>
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<td></td>
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<tr>
<td>Meningocele</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>1</td>
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<tr>
<td>Atretic meningocele</td>
<td>2</td>
<td>2</td>
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<tr>
<td>Diastematomyelia:</td>
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<td></td>
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<tr>
<td>Bone spur</td>
<td>18</td>
<td>10</td>
<td>11</td>
<td>1</td>
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<td>Fibrous spur</td>
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<td>2</td>
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</tr>
<tr>
<td>No spur</td>
<td>5</td>
<td>1</td>
<td>3</td>
<td>1</td>
<td></td>
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</tr>
<tr>
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<tr>
<td>Local atrophy</td>
<td>2</td>
<td>1</td>
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<td></td>
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<td></td>
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<tr>
<td>Total</td>
<td>104</td>
<td>43</td>
<td>39</td>
<td>12</td>
<td>17</td>
<td>6</td>
</tr>
</tbody>
</table>
In all probability, specific neuropathologic changes in the cord were not seen. Only one myelographic-CT study was done in this group. It is possible that some cord clefts or other focal changes were not defined by conventional myelography. It is also uncertain that the demonstration of cord movement between supine and prone positions excludes myelodysplasia. Of interest is the finding in two patients included in the series that local atrophy, or hypoplasia, of the cord found on myelography was responsible for lower extremity changes. One patient had disparity of leg lengths from birth, and the other patient had an inverted foot.

Tethered Cord

Fourteen patients had tethered cord. Eight of these had lumbar skin abnormalities characteristic of dysraphia, and six had foot or leg inequality and/or deformity. There were no club feet. Bilateral hydronephrosis was seen in one patient with a conus in the upper sacrum. A neuropathic bladder with bilateral reflux was noted in a patient with sacral dysgenesis. Twelve of these patients exhibited an abnormally low conal position; nine had varying degrees of filum thickening. The transverse diameter of the filum at L5 ranged from 1.5 to 4.0 mm. In three of these the cord continued without narrowing down to the point of tethering. Hydromyelia was seen in a postoperative myelogram in one of three patients in whom CT was done. A tethered cord was found in another patient with an occipital encephalocele and lacunar skull. Tonsillar ectopia to C1 was present in a patient whose cord extended to S1 (Fig. 1). Asymmetric hemicords close together were noted only by CT in a patient with typical myelographic findings for a tethered cord (Fig. 2).

All the patients showed decreased movement of the cord as myelographic position changed from prone to supine. This was also true in the two patients whose conus was at a normal level. In one of these patients a thickened filum continued from the conus. In the other the conus was tethered posteriorly by a band evident in the myelogram. The tethering lesions seen at myelography and on CT in all the patients in this category were small lipomas, meningoceles, or fibrous bands.

Lipoma

Ten children had evident or palpable lipomas. Eight of these had urinary or fecal incontinence and six had disparity in leg length as well as foot deformities. Eight patients had preoperative myelograms, which

![Fig. 1.—9-year-old girl with wasted and weak right leg.](image)

A, Conus extends to L5. Filum is thickened and tethered (arrow).
B, Supine lateral cervical myelogram shows cerebellar tonsillar ectopia to C1.
showed significant intradural lipomas intimately involving the conus and/or nerve roots. The lipomas were in either the lumbar or sacral canal and extended to the subcutaneous tissues of the back. In all these patients the spinal cord was abnormally low, and two of these eight had hydromyelia (Fig. 3). Two of the 10 patients had had surgery in infancy for their lesions, and they were examined for new or continuing symptoms. One of these was found to have a lumbodorsal hydromyelia and the other a neuroenteric cyst adjacent to the upper lumbar cord.

In three patients the spinal cord was displaced anteriorly by the lipoma; in another three the cord was in a central position in the spinal canal as it fused with the lipoma, and in two patients the cord was displaced posteriorly. The initial preoperative studies in two patients were not available. Transverse CT sections in three patients with lipomas are shown in Figure 4. The patient in Figure 4A appeared to have a well-defined interface between the neural placode and lipoma. However, the neurosurgeon found the lipoma contained considerable disorganized tissue, including fibrous elements, which made the placode difficult to identify. The lipoma was also attached by thick bands of fat and fascia to the walls of the spinal canal, and its dissection was difficult.

The CT sections of patients shown in Figures 4B and 4C had well-defined strands of tissue that could be followed from the cord into the lipoma. At surgery, neural tissue was found in the tumors. Direct correlation of this observation with the CT scans of the lipomas could not be made. In three other patients CT showed relatively homogeneous fat density in the lipomas. The surgical dissection in these patients was difficult, however, because of nerve roots at the margins of the lipomas. These could not be identified in retrospect in the CT studies.

**Lipomyelomeningocele**

Sixteen patients exhibited varying degrees of lipomatous material closely associated with myelomeningoceles. One patient had previous surgery. All the patients had palpable lumbar or sacral masses, some quite large. Seven patients
Fig. 3.—4-year-old boy with right leg weakness and double incontinence.
A, Cord is enlarged and extends to S2.
B, Hydromyelia (arrow) is noted in a 5-hr delayed scan.
C, Tethering lipoma expands sacral canal.

Fig. 4.—Intracanalicular lipomas in three patients.
A, Placode and lipoma interface (arrow) is well defined without change in lipoma to suggest presence of neural tissue. Resection of lipoma was difficult because of fibrous attachments to walls of spinal canal.
B, Well-marginated shadow (white arrow) in lipoma suggests neural tissue associated with placode (black arrow).
C, Branching bands of probable neural tissue in lipoma (white arrow); placode is at most anterior aspect of spinal canal (black arrow).
had foot or leg abnormalities, including two with unilateral club feet and three with feet and legs of different size and length. Five patients had either urinary incontinence, infection, or reflux. Two patients with urinary tract signs were less than 6 months old. One 8-year-old had fecal incontinence.

The radiologic studies alerted the surgeon that the conus was abnormal in position and involved with the lipoma and the myelomeningocele in all the preoperative patients. In seven of 10 preoperative CT studies the specific relationship between the placode, lipoma, and meningomyelocele could be resolved satisfactorily. The placode–myelomeningocele interface was better defined in one patient as a result of lateral decubitus CT scanning (Fig. 5).

Nerves in the neural sac were seen on four conventional myelograms. CT demonstrated nerve roots in the neural sac in only one patient. Compression of the myelomeningocele in the supine position prevented nerve root visualization in the remaining CT examinations. Nerve roots in the neural canal and their associated placode were seen in CT scans of three patients. The failure to define this relationship with greater frequency was attributed to the complexity of the lesion and incorrect CT windowing.

Preoperative assessment of bone abnormalities associated with the lesion was important. In one patient (Fig. 6) an anomalous bone surrounded by lipoma and not recognized on CT made resection of the lipoma and access to the spinal canal more difficult. In another patient (Fig. 7) hemisacral deficiency was seen on CT with presumed absence of sacral nerve roots. Because of this observation only cosmetic resection of the large lipoma was planned. Of interest was the prevalence of tonsillar ectopia. Although minimal in three patients, the tonsils were between C1 and C2 in another three.

**Meningomyelocele, Meningocele, and Atretic Meningocele**

Of the 12 patients in this group, nine presented with palpable masses over the spine. In another patient an anterolateral meningomyelocele was mistaken for a cystic kidney in infancy (1977) and explored without prior myelography or CT. Two patients were studied postoperatively and a small remaining meningocele was seen in one. Two patients thought clinically to have cervical dermoids with associated sinuses were shown after histologic examination to have atretic meningoceles. Four patients displayed foot abnormalities, and another had urinary incontinence.

The placode and nerve roots were seen well with CT in four patients. In two of these, CT confirmed the presence of nerve roots that were faintly seen on myelograms. CT nicely outlined the complex relationship between a diastematomyelia bone spur and a meningocele with its subcutaneous placode (Fig. 8). In another infant, CT brought out a centrally placed placode subtending two winglike extensions of the meningocele (Fig. 9). Eight of the 13 patients had tonsillar ectopia. The majority of tonsils were between C1 and just below the foramen magnum. However, in one patient with a mid-thoracic meningocele, a Chiari malformation extended to the C5 level.

**Diastematomyelia**

Twenty-three of the 25 patients in this group had lower extremity signs including disparity of foot size, leg length, and girth. The smaller extremity was associated with the smaller hemi-cord in these patients. Of the two patients with normal extremities, one was 5 months old and the other was 10 years old. Two infants developed leg abnormalities in early childhood. One child had bilateral clubfeet. Unilateral clubfoot was noted in two patients. One patient had leg wasting and an ipsilateral dislocated hip. One patient had symmetric leg wasting.

Eight of 25 patients had hairy patches at the level of the bony or fibrous spurs. Two patients had meningoceles, which were partially resected at birth, and one extended from T12 to L4. Postoperative adhesions in this case produced prominent arachnoidal cysts in the thoracic spinal canal. No urinary or bowel abnormalities were noted in this patient group.

Preoperative CT was performed in 12 patients. Separate dural sheaths around each hemi-cord could be seen in six.
Fig. 6.—1½-year-old boy with lumbosacral lipoma and right club foot. There was previous resection of the lipoma. CT scan shows placode (white arrow) between myelomeningocele and lipoma. Black arrow shows anomalous bone and adjacent lipoma. Resection of lipoma was difficult because of attachment to anomalous bone.

Fig. 7.—8-year-old girl with lumbar lipoma and small deformed feet bilaterally. Lipoma is in left hemisacral region where bone is absent. Lipomatous mass extends to left buttock (solid arrows). Neural tissue in placode (upper open arrow) could be followed into lipoma. Thecal sac is adjacent to placode (lower open arrow).

Fig. 8.—2-year-old boy with lumbosacral hemimyelocele and bone spur. Left leg is shorter and thinner than right leg.
A, Prominent nerve root (upper arrow) spans myelocele and joins placode (lower arrow).
B, Bone spur and adjacent neural tissue (arrow) is seen in next more rostral scan.

Fig. 9.—6-month-old girl with compressible mass in lumbar region.
A, Myelogram shows apparent myelomeningocele extending into subcutaneous tissue of back.
B, CT clearly brings out relationship between placode (arrow), myelomeningocele, and nerve roots. A cursor was placed at posterior aspect of placode.
patients. Complete or almost complete bone spurs were present in these patients. Four patients had hemicords enclosed in a single dural sheath; two of these had small incomplete spurs, and two had none. One patient had an unusual finding of renal blastema attached to an incomplete bone spur (Fig. 10). CT showed seven bone spurs arising from posterior elements; three came from the posterior vertebral body margin.

**Dermoid**

One infant had a dermoid with cystic change and hair in the conus. The lesion was brought to clinical attention because of the child’s refusal to bear weight on one leg. A preoperative myelogram showed a mass expanding the conus obstructing cephalad flow of contrast material. The second patient presented with a discharging sinus tract connecting with the filum in the sacrum; these findings were confirmed at surgery.

**Discussion**

The radiology of spina bifida occulta has advanced remarkably over the last 25 years. Lesions in the spinal canal such as dermoids, lipomas, and bone spurs could be seen with Pantopaque (Myodil).\(^1\) Visualization of cord or thecal abnormalities was limited, however [11]. Air-myelography in the 1970s was a step forward, affording better definition of cord outline and movement [12]. However, sufficient detail, especially for conal position, was frequently lacking.

The major breakthrough for more precise resolution of the pathology of closed dysraphism has been the use of nonionic water-soluble contrast material, as well as the advent of CT. Iohexol [13] was used in all the myelograms in this series and there has been no complication from its use other than postexamination headache and nausea, some of which could be attributed to general anesthesia.

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1 Lafayette Pharmacal, Inc., Lafayette, IN.

CT was used preoperatively in one-half of the positive cases and probably would have added further information in the other half. But CT examination has to be carried out shortly after the conventional myelogram, before dilution of contrast material occurs, and this may not always be possible in a busy radiology department. CT studies also require careful monitoring to ensure cross-sectional imaging at appropriate levels of known pathology. Correct windowing and filter use is important to maximize the details of cord and nerve root relationships in thecal sacs to differentiate meningo(myelo)celes from meningoceles.

This is also true for lipomyelomeningoceles. The usual [14] asymmetric relationship of the lipoma, placode, and meningocele was seen in all the patients in this series. The placode in some lesions was difficult to define, and when seen appeared to be of irregular outline in both the adjacent lipoma and myelomeningocele.

The recognition by CT of neural tissue in lipomas and the lipomatous part of the lipomyelomeningocele in the reported cases was difficult. Although braidlike tissue in two lipomas (Figs. 4B and 4C) could be traced from the placode-lipoma interface, neurosurgical confirmation of this observation was not possible. The neurosurgical reports in several cases in this series indicated that nerve roots or fibrogliotic material were present in lipomatous tissue. This observation has been reported in surgical [15] and pathologic material [16].

Specific correlation of the tissues present in the lipoma could not be made in this series. This was attributed to fractional resection as well as to laser coagulation of the tumor mass. If definite patterns of fibrous and neural tissue can be established by CT or MR, considerable help would then be available for the surgeon in planning the debulking of the lipoma. This information would also be useful in assessing the possibility of untethering the cord and placing it in the spinal canal with only a limited remaining cuff of fatty material.

The surgery for lipomas, lipomyelomeningoceles, and meningo(myelo)celes is often formidable [14]. Myelography and CT can at present show the surgeon major structural points of the lesion and its relationship to the vertebra or sacrum.

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![Fig. 10.—5-month-old girl with L5/S1 diastematomyelia. Patient had sacral dimple and no other abnormality.](image-url)

A, Bone spur is denoted by upper arrow. Lower arrow shows soft tissue at tip of spur. The histologic report for the material was renal blastema and neural tissue.

B, Next rostral scan shows thin hemicords (solid arrows) enveloped by separate dural sheaths. Bone spur (black open arrow) and soft tissue component of septum (white open arrow) are seen. Note cleft in first sacral segment.
The same is true for various expressions of diastematomyelia. Finer details—such as exact position and number of nerve roots in relation to the placode-lipoma, the presence of fibrous bands or arachnoidal adhesions, or prominent blood vessels—were discovered only at the time of surgery in this series.

An important point in our study is the visualization in plain films and CT of anomalous or deficient bone formation associated with dysraphic lesions. Naidich et al. [14] and others [17, 18] have emphasized recognition of posterior bone masses. Surgery can be facilitated with the foreknowledge that these may need removal for better access to the dysraphic pathology. Noting that a hemisacrum is largely absent and that sacral nerves are probably absent or abnormal may obviate surgery. CT-based 3D reconstruction of bone has now advanced to a level where the skeletal framework of dysraphic lesions can be shown to good advantage. The pathology of diastematomyelia can be shown very well by CT-myelography, as documented by Naidich and Harwood-Nash [5]. Abnormal bones and soft tissues depicted in 3D would be helpful in presurgical assessment. For informative display, however, the increased number of thin CT sections would be time-consuming and could have prohibitive radiation dosage.

The clinical findings in this survey have been enumerated not only to catalogue their prevalence but also to emphasize what dysraphism pathology means to patients, their parents, and their primary physicians. The imaging focus has been strongly directed toward revealing the pathology, with the surgeon operating in the belief that “untethering” the cord will arrest the progress of clinical signs. Unfortunately, this does not always occur. It is evident also from the study that nearly a quarter of patients with suspected dysraphism have a normal-appearing cord without restriction of movement.

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REFERENCES