FAMILIAL SPONDYLOLISTHESIS OF THE AXIS VERTEBRA


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This report describes a nine-year-old girl with a spondylolisthesis of the C2 vertebra allowing 14 mm of slip. Her father had very similar vertebral anomalies.

In contrast to the lumbar spine, spondylolysis and spondylolisthesis involving the cervical spine is very uncommon, only 27 cases of spondylolysis and 22 of spondylolisthesis having been reported (Dawley 1971; Fardon and Fielding 1981). Only three cases involving the axis vertebra have been described and only one of these had spondylolisthesis of the second on the third vertebra. We report a girl with spondylolisthesis of the axis and other vertebral and minor skeletal anomalies. The family history suggests an autosomal dominant inheritance of this condition.

CASE REPORT

A nine-year-old girl, the only child of the family, was referred after she had fallen on her back. She complained of mild local neck pain with no radiation and no neurological abnormality and had no previous history of spinal injury. She was slender with disproportionately long extremities and a slightly elongated neck. Her height was 140 cm and her span 152 cm, suggesting some marfanoid disproportion. Her fingers were long and narrow with broader distal segments and nails. Her neck movements were normal as were her cardiovascular and respiratory systems. She had slight hypertelorism but no visual impairment or lens dislocation.

Radiographs showed each pars interarticularis of the axis to be about twice the normal length with elongated inferior articular surfaces, and bilateral anteriorly placed spondylolysis (Figs 1, 2 and 3). In full extension the posterior surfaces of C2 and C3 were in line with the other vertebrae but in full flexion C2 slipped 14 mm anteriorly on C3, with tilting at this level from −12° to +13°. Tilt also occurred at the spondylyotic defect, changing the angle between the spinous process and the inferior surface of the body from 29° to 56°. The body of C6 was dysplastic and showed bilateral spondylolysis with minimal slip. Radiographs showed spina bifida of C2, C6, T12, L1, L5, S1 and S2 vertebrae. The metacarpal index of her hand was 10.2, clearly exceeding the normal value of 7.9 at this age (Eldridge 1964; Rand et al. 1980). The second and the third metacarpals were elongated (Fig. 4) as were the second and the third metatarsal bones. Radiographs of the skull and the chest showed no abnormalities.

In view of the unusual nature of the case, the patient was treated with skull traction for six days and then wore a neck support for six weeks. At no time did she show any abnormal neurological signs, despite the 14 mm amplitude of instability. No operative treatment was offered and she was allowed to mobilise her neck freely.

Fig. 1

Lateral radiograph of the upper cervical spine taken soon after the episode of minor trauma to show the chronic defect and long pars interarticularis of the axis vertebra.
Familial Spondylolisthesis of the Axis Vertebra

Figures 2 and 3—Flexion and extension radiographs of the cervical spine taken 19 months later. In flexion the superior articular surfaces of C3 lie against the posterior parts of the elongated inferior articular surfaces of C2. In extension they articulate with the anterior part of the articular surfaces. Note the angular movement of the lamina of C2 in flexion and extension, and the spondylolysis of C6 vertebra. Figure 4—Radiograph of the right hand to show the long second and third metacarpals with slightly short phalanges in these fingers.

At review one year and seven months after the minor injury she had no symptoms that could be referred to the spondylolisthesis and had led a normal life, taking part in sport activities at school.

Family history. The patients’ 37-year-old father mentioned that he had suffered mild transient low back pain 10 years previously. On examination he also showed disproportionately long extremities, but had a short neck and a slight thoracocervical kyphosis. His fingers were long and slightly hyperextensible with short and broad distal phalanges and flattened nails like those of his daughter. He also had elongated second and third metacarpals and metatarsals, with a metacarpal index of 8.2. His height was 183 cm and his span 202 cm. He had normal cardiovascular and respiratory systems and normal vision.

Radiographs of his cervical spine (Fig. 5) showed a bilateral corticated spondylolysis of C2 with a 2 mm forward slip of C2 on C3 in full flexion which was fully corrected in extension. His C6 vertebra showed dysplasia and spondylolysis. This was remarkably similar to the situation in his daughter except that he had only slight elongation of the pars interarticularis of the axis. The girl’s father has one sister and six brothers, but neither his parents nor any of his siblings had a history of spinal problems.

DISCUSSION

A number of studies have shown that the dysplastic type of lumbosacral spondylolysis and spondylolisthesis has a monogenic aetiology, and that the mode of inheritance of the condition is autosomal dominant with reduced penetrance (Amuso and Mankin 1967; Taillard 1976; Wiltse, Newman and Macnab 1976; Haukipuro et al. 1978; Albanese and Pizzutillo 1982). Associated anomalies of the lumbar vertebrae, such as spina bifida, have been frequently reported, but other regions of the spine had not been examined in these cases.

Our case suggests the need for careful clinical and possibly radiographic evaluation of patients with cervical spondylolisthesis or spondylolysis for other vertebral or skeletal anomalies. Furthermore, in view of the autosomal dominant mode of inheritance other members of

Fig. 5

Lateral radiograph of the cervical spine of the patient’s father to show spondylolysis of C2 with a forward slipping of 2 mm on C3 and dysplasia of C6 with spondylolysis.
the family should also be examined. It would appear that
the clinical expression of the condition may be variable,
and even non-penetration may be encountered.

REFERENCES
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BRIEF REPORT
SCIATICA CAUSED BY PERIFIBROSIS OF THE SCITATIC NERVE
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Case report. A 39-year-old man was admitted with a
history of right sciatica without pain in his back. He had
never had symptoms from his back or legs until six
months earlier when he fell two metres into a pit, landing
on his right buttock. Since then he had had right sciatica
extending down to the outer aspect of the ankle after sit-
ing for more than 10 minutes. Conservative treatment
had no effect. On admission he had reduced sensation to
touch and to pain on the lateral aspect of his right calf
and in the first dorsal web space of his foot, slight weak-
ness of dorsiflexion of his ankle and hallux, a positive
Lasègue's sign at 60°, and a positive Tinel sign over the
sciatic nerve in the mid-thigh. The tendon reflexes were
normal. Radiographs of his lumbosacral spine showed
spondylolisthesis at L4/5, with only slight forward slip.
Myelography demonstrated slight shortening of the right
fifth lumbar nerve root.

At exploration through a right partial hemilaminecto-
ymy no intervertebral laxity was demonstrated and
there was no disc herniation. Three weeks after this
operation his symptoms recurred and once again con-
servative treatment had no effect. When he was readmit-
ted three months later, his signs, including the positive
Tinel sign, were unchanged. A CT scan of the lumbar
spine showed no disc herniation. Electromyography sup-
ported the clinical suspicion of damage to the sciatic
nerve itself, with a partial block to conduction. At opera-
tion the nerve was found to be surrounded by tight fibro-
sis in the gluteal region and was decompressed.

Six months later the patient had resumed his job as a
policeman. His signs and symptoms had subsided, leav-
ing only slight wasting of the thigh; the Tinel sign and the
Lasègue sign were both negative.

Discussion. Sciatic nerve damage causing pain radiat-
ing distally is unusual (see references). The following charac-
teristics should arouse suspicion: a history of direct
trauma to the nerve; aggravation of symptoms when
pressure is applied, for example, by sitting; a positive
Tinel sign; exclusion of a herniated disc by myelography
or CT scan; and an electromyogram compatible with a
sciatic nerve block. None of these findings is specific and
a definitive diagnosis can only be achieved if, at opera-
tion, the nerve is found to be compressed and if signs and
symptoms subside after decompression. Electromyogra-
phy is probably the most useful diagnostic aid.

REFERENCES
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