Dominant transmission of Sprengel's shoulder and cleft palate

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SUMMARY  We present a family showing apparently autosomal dominant transmission of Sprengel's shoulder with cleft palate. Some, but not all, of the kindred showed features of the Klippel-Feil syndrome. The association between these abnormalities is discussed.

The isolated abnormality of cleft palate is considered to be inherited in a multifactorial manner, although when associated with lip pits, cleft palate is inherited as a Mendelian dominant condition (Van der Woude syndrome). Sprengel's shoulder (congenital raising of the scapula) usually occurs as a sporadic event, but there are six cases of isolated Sprengel's shoulder showing Mendelian inheritance. We report a family in which these two abnormalities are associated and

FIG 1 X-rays of cervical spine and shoulders of the proband.
inherited together in a dominant manner. This has not to our knowledge been described before. One member of the family had a Klippel-Feil anomaly of the neck, which has been reported to be associated with each of these other anomalies separately.

Case reports

A 26-year-old woman attended our clinic for genetic advice because she had a cleft palate. On examination she had a repaired congenital cleft palate without cleft lip. There were no lip pits or abnormalities of the teeth and the palate was not arched. She also had bilateral Sprengel's shoulder with large wide scapulae rotated medially and placed high on the thoracic cage. She could not raise her arms above the horizontal, and there was some limitation of neck movement, but the neck was not short. There were no other orthopaedic abnormalities and her hearing and intelligence were normal. X-ray of the cervical spine was entirely normal (fig 1).

FIG 2 Pedigree of family.

FIG 3 X-rays of cervical spine and shoulders of the proband's father.
Both her father and paternal grandmother had cleft palate and bilateral Sprengel’s shoulder, and her affected grandmother’s father had bilateral Sprengel’s shoulder but no cleft palate (fig 2). The father of the proband was a twin, but the other twin had died at birth and was said to have been deformed.

The father was examined. He had a cleft palate and bilateral Sprengel’s shoulder, more pronounced on the right than on the left. He also had a short neck with limitation of movement, particularly rotation. The arm movements were also restricted. The palate was not arched and the palatal cleft had been repaired. His lips were normal and he had normal dentition.

X-ray of his cervical spine showed a fusion of the arches and bodies of the fourth and fifth vertebrae. The other vertebrae showed spondylotic change at C5/6 and there was degenerative disease of the posterior intervertebral joints at C5/6 (fig 3).

Discussion

There are two published reports which present firm evidence of Mendelian inheritance of Sprengel’s shoulder. In both cases this was dominantly inherited through several generations. Four other families have been reported with small pedigrees which present possible evidence of Mendelian inheritance, but the pedigrees are too limited for certainty. In none of these pedigrees was cleft palate reported.

Sprengel’s shoulder deformity is associated with congenital defects of the cervical and thoracic vertebrae and ribs in about 50% of cases. Those associated with cervical vertebra anomalies merge into the Klippel-Feil syndrome, characterised by fusion of two or more cervical vertebrae, giving the clinical triad of short neck, low posterior hairline, and limitation of head and neck movements. The frequency of Sprengel’s shoulder in Klippel-Feil syndrome has been given as 30%.

Klippel-Feil syndrome is probably heterogeneous aetiology and is usually sporadic, although familial cases with possible dominant inheritance have been reported.

Cleft palate has been described in 5 to 20% of cases of Klippel-Feil syndrome. One explanation is that there is a developmental link between cervical fusion and cleft palate, with vertebral fusion causing compression of the mandible against the chest, forcing the tongue between the palatal shelves and preventing their closure in the midline. With the family reported here, however, Klippel-Feil syndrome was absent in the proband with cleft palate although present in her father, indicating that this mechanical explanation cannot be invoked in this kindred. The association of these anomalies within one kindred with autosomal dominant inheritance of cleft palate and Sprengel’s shoulder, each of which has separately been reported with the Klippel-Feil syndrome, suggests that these three anomalies are very closely linked.

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References


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