Discordant, non-syndromic, congenital diaphragmatic defects in sibs

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SUMMARY We report an Arab sibship of two brothers with non-syndromic, congenital diaphragmatic defects (CDD). The first had an extensive, left, Bochdalek-type hernia and the second hemidiaphragmatic agenesis; these were verified by surgical exploration and necropsy respectively. The parents are healthy second cousins. Other reported discordant cases of CDD are briefly reviewed.

Congenital diaphragmatic defects have been classified into (1) posterolateral (Bochdalek) hernia, (2) retrosternal (Morgagni) hernia, and (3) the rare extensive defects involving most or all of the hemidiaphragm (agenesis). They are generally considered to be one entity, that is, malformation or developmental defects with an incidence somewhere between 1 in 800 and 1 in 10,000. Makela reported the first family with affected sibs in 1916. Mertins, in 1952, was the first to suspect that CDD could occur in a familial aggregation. In Kuwait, we recently observed an Arab family with discordant, non-syndromic CDD in sibs.

Case reports

CASE 1
A male infant was born in 1984 after an uneventful, term pregnancy, weighing 3600 g. He was the second child of phenotypically normal, second cousin Jordanian parents. At birth he looked normal. His Apgar scores were 8 and 9 at one and five minutes. He rapidly developed severe respiratory distress and metabolic acidosis. An extensive, left sided diaphragmatic hernia was diagnosed clinically and by chest radiography. The diagnosis was verified by surgical exploration. The hernia sac contained the stomach, the intestines extending to the splenic flexure, the spleen, the left lobe of the liver, and the left testis. It was repaired at the age of six hours. The baby died four hours later from cardiac arrest. The mother, when pregnant again, was given a low risk of recurrence on the basis of the multifactorial inheritance hypothesis.

CASE 2
This male sib of case 1 was born in 1985 at 38 weeks' gestation, weighing 2800 g, after apparently normal ultrasonography findings in utero at 37 and 38 weeks. His Apgar scores were 1 and 3 at one and five minutes. He had progressive respiratory distress and acidosis, and he eventually died. Necropsy showed left hemidiaphragmatic aplasia. The left lobe of the liver, the spleen, the stomach, and most of the intestines were herniated into the thoracic cavity.

After two affected children, a high risk figure (25%) was quoted to the parents. The mother had an early spontaneous abortion in 1986, a healthy daughter in 1988, and is now pregnant again, in her first trimester.

Discussion

The occurrence of discordant, non-syndromic CDD in sibs is well known. Most reported families had one baby with the Bochdalek type of hernia and the second with hemi- or bilateral diaphragmatic agenesis, indicating that both malformations may have common causes. In Arabs, Arad et al reported the first affected sibs in Jerusalem. In Kuwait, with a unique mosaic population, we have to consider high recurrence risk estimates, especially in CDD with extensive diaphragmatic defects or agenesis. The present family offers further evidence that among the frequent multifactorial cases of non-syndromic CDD are a few cases of autosomal recessive inheritance.

References

Porencephalic cyst in pycnodysostosis

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SUMMARY We describe a case of pycnodysostosis with porencephaly and suggest an explanation for the porencephaly by a mechanism of imbalance between brain growth and its vascular supply and a normal but unopposed cerebrospinal fluid pressure.

In 1962 Maroteaux and Lamy1 described a skeletal dysplasia the main features of which are generalised osteosclerosis and moderate dwarfism. They named it pycnodysostosis meaning ‘thick bones’.

Other features of pycnodysostosis include underdevelopment of the cranial sinuses; wormian bones; delayed closure of the fontanelles with wide, unossified cranial sutures; a beaked nose; a high, grooved palate; multiple dental abnormalities; hypoplastic
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