Hypoglossia-hypodactyly syndrome with hydrocephalus: a clue to the aetiology?

Y Gillerot, L Van Maldergem, R Chef, L Koulischer

Abstract
A stillborn female with Hanhart's syndrome in association with hydrocephalus owing to stenosis of the aqueduct of Sylvius is presented. Neuropathological findings are suggestive of an acquired pathophysiological mechanism.

The association of terminal transverse defects and orofacial abnormalities has been subdivided by Temtamy and McKusick1 into several clinical entities. Hanhart’s syndrome covers aglossia-adactyly, hypoglossia-hypodactyly, and peromelia with micrognathia.2 The aetiology of this rare syndrome remains unknown and environmental factors as well as genetic factors have been postulated.3 We report on a new case of this syndrome with central nervous system alterations suggestive of an acquired inflammatory origin.

Case report
This female infant was the third child of healthy, unrelated parents. The mother’s second pregnancy ended at 30 weeks with the birth of a baby with left heart hypoplasia who survived a few days. The current pregnancy was uncomplicated until the birth at 36 weeks of a stillborn female baby with multiple malformations (fig 1). Birth weight was 2400 g, length was 46 cm, and head circumference 32 cm. Clinical examination showed a proportionate large head with wide fontanelles. There were reduction anomalies involving both hands (severe hypoplasia of the second, third, and fourth rays owing to absence of the second and third phalanges) and feet (total absence of phalanges). Examination of the mouth showed marked hypoplasia of the tongue (fig 1), which was closely adherent to the palate. A cleft palate was seen on dissection.

Macroscopic examination of the brain confirmed, after section, a slightly dilated ventricular system. Macroscopic examination showed an intense fibroblastic proliferation of meningeal tissue around the brainstem (fig 2a). Originating from this thickened meningeal envelope were small vessels penetrating deeply into the parenchyma inducing an astroglial reactional proliferation and ending in a necrotic calcified zone (fig 2b). The aqueduct of Sylvius was abnormal and showed a number of small channels with an intact ependymal lining. The cerebral hemispheres were thin and the cerebellum was normal.

Peripheral lymphocyte culture showed a normal 46,XX karyotype (R and G banding).

Discussion
This stillborn female fulfils the criteria of aglossia-adactyly syndrome.1 Autosomal recessive inheritance has been postulated by Tuncbilek et al4 in view of three reports of children born to consanguineous parents. Delligrammaticas et al5 also suggested recessive inheritance, one of their two patients also being the product of a consanguineous marriage. Temtamy and McKusick1 suggested a dominant trait with variable expressivity since orofacial abnormalities, although less severe, were observed among the relatives of their reported cases. A dominant mutant gene was suggested by Nevin et al.5

However, Opitz,6 in a comment on the case of Delligrammaticas et al,3 considered that there is, at present, no convincing evidence for considering this condition as a Mendelian trait and suggested a negligible recurrence risk should be given after the birth of a child with Hanhart’s syndrome. Other reports implicate a secondary phenomenon for explaining this syndrome, such as environmental factors including drug intake during pregnancy,7–8 amniotic bands, and thromboembolic events.9 This last report was of three patients among whom were a discordant pair of monozygotic twins, an infant with an apple peel bowels, possibly secondary to an occlusion of the superior mesenteric artery, and a third child with a malformation complex also compatible with an intrauterine arterial occlusive event.
Hypoglossia-hypodactyly syndrome with hydrocephalus: a clue to the aetiology?

Figure 1  General appearance of the proband. Note hypoplastic tongue.

Our case is, to the best of our knowledge, the first observation of Hanhart's syndrome associated with hydrocephalus and neuropathological changes. One could hypothesise the coincidental occurrence of Hanhart's syndrome and acquired hydrocephalus or, conversely, the association of these two conditions as a consequence of a similar causative agent is another possibility. The reported cases with vascular involvement mentioned above support the last hypothesis in which limb buds, tongue, and brain would be subject to the same vascular changes, thus leading to the syndrome described here.

The authors would like to thank Professor G Lyon (UCL, Brussels) for neuropathological data on this case.