Syndrome of polydactyly, cleft lip/palate or lingual lump, and psychomotor retardation in endogamic gypsies

V VÁRADI, L SZABÓ, AND Z PAPP

From the Departments of Paediatrics, Stomatology, Obstetrics, and Gynaecology, University Medical School, Debrecen, Hungary

SUMMARY  Six children in an inbred isolate (a gypsy colony) were found to have a syndrome of reduplication of the big toes, supernumerary fingers on the hands, cleft lip/palate or lingual nodule, and somatic and psychomotor retardation. Other features sometimes present were absence of olfactory bulbs and tracts, cryptorchidism, inguinal hernia, and congenital heart disease. The disorder has overlapping features with several previously delineated syndromes, but is most similar to the anomalies of trisomy 13 Mand ohr’s syndrome. Our patients had a normal karyotype. The mode of inheritance of this lethal genetic syndrome is probably autosomal recessive.

Since 1960 many cases of trisomy 13 syndrome have been reported, and its characteristic features are well defined.¹⁻⁴ There are also reports of similar phenotypes with normal karyotype.⁵⁻⁶ We have recently studied a male gypsy child with a combination of malformations similar to those of Patau’s syndrome and a normal male complement of 46 chromosomes in all cells cultured from peripheral blood. In this gypsy colony a further five cases have been found with similar features. The six cases are of interest from the nosological point of view and merit a detailed description.

Case report

Near Debrecen (the largest city in eastern Hungary) there are two large gypsy colonies 30 km from each other in Hajdúhadház and Hosszúpály with about 800 and 1500 residents, respectively. Within gypsy groups many children are born as a result of extramarital relationships, and because of the difficulties of tracing connections it has been possible to draw the pedigrees for only three generations (fig 1, 2, 3). The large number of admitted extramarital connections and the small number of surnames (only six) reveal a high degree of endogamy in these colonies. Hence, although consanguinity between families A, B, and C is not admitted, it is probable that it exists. Details of the six affected children are presented in the table, and additional features of the proband are given below.

CASE 1: F-K, III.22, FAMILY A
The parents are healthy and normally developed. The mother was aged 20 years and the father 21 years. The mother was aged 20 years and the father 21 years. The mother was aged 20 years and the father 21 years.
years at the child's birth. The mother gave no reliable history of illness or of having taken drugs during this, her first, pregnancy. Labour and delivery were normal and the child was born at term, birthweight 2300 g and length 47 cm. Most of the malformations were noted at birth (fig 4, 5). He had a right-sided cleft lip and palate, seven fingers on the left hand, six on the right hand, and six toes on each foot. The extra big toes were symmetrically placed. Additional features were elongated skull, ocular hypertelorism, epicanthic folds, strabismus, low-set malformed ears, equinovarus feet, inguinal hernia, and undescended testes. The karyotype on peripheral lymphocyte culture was normal, 46,XY. The mother's karyotype was 46,XX.

He had failed to thrive from birth, had continuous high temperature, and was retarded both physically and mentally, with only minimal vegetative functions. He had not been observed to have seizures. No protective or neonatal reflexes could be elicited. Psychomotor development was clearly retarded and at the age of 12 months corresponded to that of a 4-
 Syndrome of polydactyly, cleft lip/palate or lingual lump, and psychomotor retardation

FIG 4  Case I (F-K). Right-sided cleft lip and palate. Minor aberrations of face.

FIG 5  Case I (F-K). Reduplication of the big toes.

FIG 6  Case I (F-K). Ventral view of brain stem and cerebellum. Note absence of olfactory bulbs and tracts.

FIG 7  Case 2 (R-N). X-ray of polydactyl hands at the age of 5.

month-old child. He showed hypotrophy of the muscles of the trunk, shoulders, and upper arms, and general pronounced muscle hypotonia. Gross motor development was characterised by reduced head balance and mobility of the arms and legs. He could not speak and could neither sit up nor walk. His eyes did not follow objects which passed in his visual field.

At his death, aged 3 years, he weighed 5400 g with a length of 73 cm, fronto-occipital diameter of 16 cm, and biparietal diameter of 10 cm. The teeth were irregular with hypoplastic enamel, and some were missing. The direct cause of death was lung oedema, resulting from insufficiency of the left ventricle.

The necropsy uncovered further malformations including congenital heart disease (aortic stenosis), cholangiograms, absent olfactory bulbs and tracts, absent fossa interpeduncularis, and absent vermis in the brain (fig 6).

Discussion

The full complement of defects observed in these six children is unusual, although a number of them are characteristic of previously described syndromes. The features are most similar to the anomalies which have been found in trisomy 13 and in Mohr's syndrome. The anomalies which the proband had in common with the majority of established cases of trisomy 13 were harelip and cleft palate, feeding difficulty, failure to thrive, developmental and psychomotor retardation, polydactyly, absence of olfactory nerves, cardiac defect, ocular hypertelorism, epicanthic folds, strabismus, low-set malformed ears, equinovarus feet, undescended testes, and inguinal hernia. In all six cases a malformation
of the mouth was observed, either a cleft lip/palate (four cases), or a lingual nodule (two cases). Lingual lumps are associated with malformations of structures derived from the first branchial arch anlage, cleft lip, cleft palate, and dental anomalies, in Mohr's syndrome.9,10 However, they are not found in trisomy 13, and we have found no reported cases of Mohr's syndrome associated with absence of olfactory bulbs and tracts, which is a characteristic feature of Patau's syndrome.

A survey of published reports failed to reveal a report of familial occurrence with a similar combination of malformations to our case. We think that the findings in the proband and in other members of the gypsy families represent a nosological entity. The fact that the syndrome is confined to three families makes it unlikely that an environmental factor is responsible. G banding carried out in three cases showed no translocation or other structural rearrangement in any of the chromosomes, including autosome 13. The gene is lethal, with four of six dying within 2 weeks, one at 3 years, and one at 6 years. Examination of the pedigrees A, B, and C suggests that the mode of inheritance is most probably autosomal recessive and is unlikely to result from a dominant gene with incomplete penetrance.

We are grateful to Drs Sz Szakáll, E Martini, and P Molnár for permission to use data from their patients.

References


Requests for reprints to Dr V Váradi, Department of Pediatrics, University Medical School, H-4012 Debrecen, Hungary.

Note added in proof

Since submitting our manuscript we have confirmed a further case of this syndrome from the same gypsy colony. The mother of the patient has had a total of 11 children to date including, in addition to the patient, three who died within the first 3 months of life and seven who are alive and healthy. The patient, who is now 3 years old, is being cared for in a home for mentally retarded children, was small for dates and has polydactyly on each hand and foot, cleft lip and palate, and somatic and psychomotor retardation. The karyotype is normal. According to the parents one of the children that died also had extra fingers and toes and a hare lip.

This syndrome is detectable by fetoscopy and therefore provides an example of a type of mental retardation that is potentially preventable by selective abortion.
Syndrome of polydactyly, cleft lip/palate or lingual lump, and psychomotor retardation in endogamic gypsies.
V Váradi, L Szabó and Z Papp

doi: 10.1136/jmg.17.2.119

Updated information and services can be found at:
http://jmg.bmj.com/content/17/2/119

**Email alerting service**

*These include:*

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Notes**

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/