LETTERS TO THE EDITOR

The frequency of mental retardation in hypochondroplasia

Hypochondroplasia is a relatively mild skeletal dysplasia usually with less growth retardation than achondroplasia. It is clearly autosomal dominant and many familial cases have been reported.

One difficulty in genetic counselling is the association between hypochondroplasia and mental retardation. Langer1 has suggested that 20% of cases of hypochondroplasia suffer from mild mental retardation. Hall and Spranger2 reviewed 32 patients in whom there was both clinical and radiological data, and found three out of 32 (9.4%) with mental retardation. The mental retardation appears to be found in both sporadic and familial cases, as Frydman et al3 reported development delay in the son of an affected but intellectually normal mother.

In order to obtain a more accurate figure for counselling, we reviewed the data on hypochondroplasia from the Skeletal Dysplasia Group’s Register. There were 27 cases of hypochondroplasia (15 female, 12 male) of whom 11 were familial cases. They had been ascertained through orthopaedic genetic clinics. The age range was 2 to 69 years with a mean of 27 years. Mental handicap was not a feature in any of the 27 cases. This suggests to us that the figures of 10 to 20% frequency of mental handicap may be overestimating the true frequency of mental handicap in this disorder.

RUTH WYNNE-DAVIES
2 Dale Close,
St Ebbe’s,
Oxford OX1 1TU.

MICHAEL A PATTON
Department of Clinical Genetics,
St George’s Hospital Medical School,
Cransmer Terrace,
London SW17 ORE.


Unusual inheritance of Becker type muscular dystrophy

Tommerup1 has recently pointed out that a family published by us in 1978,2 in which we suggested an unusual inheritance of Becker type muscular dystrophy (BMD), was compatible with the presence of a familial reciprocal X;autosome translocation. At that time there were three generations containing affected subjects including the mother and grandmother of the index case, who also had two brothers with the disease. The affected females had fairly severe symptoms and we proposed that, by chance, the normal X chromosome was inactivated in more cells than in the usual non-manifesting heterozygote. Tommerup1 suggested that a balanced X;autosome translocation would explain this type of non-random X inactivation, and he further stated that this abnormality had been found in at least 20 females with BMD or Duchenne muscular dystrophy.3

Chromosome studies were not done in the initial study of the family; however, after exchanging letters with Dr Tommerup, we decided to perform them in order to test the above hypothesis. When the family was contacted again, it was found that the proband now has a male child who is 12 years old and is very likely also affected, as indicated by bilateral calf hypertrophy and abnormally raised serum CK levels found by another physician several months before we examined him.

In spite of this probable male to male transmission of the disease, which eliminates the possibility of it being X linked, we went ahead with the chromosome studies to rule out rare situations, such as the son being a 46,XX male. We studied the proband, his wife, and their affected male child. Peripheral blood samples were obtained and cultured in the usual manner with PHA stimulation. After harvesting, the slides were stained to obtain G bands. Thirty metaphases from each subject were analysed and all three subjects were found to be cytogenetically normal.

We now believe that our family represents an unusual, dominantly transmitted form of muscular dystrophy, rather than an unusual inheritance of BMD, as we could find no similar published case.

R LISKER
O MUTCHINICK
L RUZ
Department of Genetics, Instituto Nacional de la Nutrición, Fasco de Queraga No 15, Tlalpan 14000, DF, Mexico.


Haematometra in the Langer–Giedion syndrome

Recently we reviewed a 25 year old woman with the Langer–Giedion syndrome whose case had been diagnosed and reported at the age of 8 years by Koslowski et al4 in 1977. She was mildly intellectually handicapped and lived at home with her mother. Her general health was fair. She had short stature (132 cm) accentuated by a moderate thoracic kyphosis. Her facies was characteristic of the syndrome but with some overall coarsening (figure). She had multiple exostoses visible and palpable at the wrists and ankles; one on her scapula had required surgical excision. Over the years her mild deafness had not increased but she had two major medical problems. The first was ureteric reflux (noted when she was aged 8 years) with recurrent urinary infections; in the past year she had had her ureters reimplanted in the bladder.

The second problem had been episodic abdominal pain starting about the age of 15 years. At that time she had had some blood loss from the vulva which was interpreted as scanty menstruation but, in retrospect, was more likely associated with haemor-
The patient aged 25 years.

Congenital hydrometrocolpos. Clearly hydrometrocolpos with this problem for what it was. Externally the vagina was represented by a dimple and internally it was atretic. The left fallopian tube, uterus, and cervix were distended with old blood. The ovaries were normal. After full discussion with the patient and her parents a hysterectomy was carried out with subsequent relief of symptoms.

A chromosome study showed an interstitial deletion of bands 8q24.11 and q24.12.

In their definitive review of 36 patients with this syndrome Langer et al. found ureteral reflux or urinary infections in three patients (including ours). Congenital hydrometrocolpos was reported in one patient by Fryns et al. This was discovered at birth and treated surgically without complications. Clearly hydrometrocolpos and haematometra are complications of this syndrome which deserve attention.

M W PARTINGTON Regional Medical Genetics Unit, Western Suburbs Hospital, Newcastle, New South Wales 2298, Australia.

J RAE Community Health Centre, 180 Peel Street, Tamworth, New South Wales 2340, Australia.

M J PAYNE Tamworth Base Hospital, Tamworth, New South Wales 2340, Australia.


Two additional patients representing the possible human homologue for the mouse mutant disorganisation (Ds)

Winter and Donnai reported a patient with striking congenital defects (short upper and lower segments of the right leg with a popliteal web and nine toes on that side, a finger-like structure arising from the abdo-

men, and absent right kidney). They reviewed 13 similar reported patients with an extra lower or duplicated limb who had additional features such as skin tags, appendages, or papilae, lipomas or hamartomas, and genito-urinary abnormalities. The authors suggested that these patients represent the human homologue of mice heterozygous for the mutant disorganisation (Ds) semidominant gene. They also postulated that certain fetuses and infants reported to have amniotic rupture with malformations inadequately explained by band disruption may be additional examples of the human homologue of disorganisation. A new patient is presented here as a possible example of this fascinating malformation complex.

This boy was the second child born to an unrelated, healthy 29 year old mother and 27 year old father. The mother had had a first trimester miscarriage which was not examined. There was minimal first trimester bleeding, normal fetal activity, and no exposure history. An ultrasound done at the time of spontaneous premature labour at 35 weeks' gestation showed oligohydramnios and a poorly defined limb abnormality. The baby was delivered by caesarean section because of breech position. There was no heart rate at delivery and he lived one hour.

Birth weight was 2000 g (50th centile for 34 weeks). On external examination there was a box shaped cranium, wrinkled Potter’s facies (fig 1) with apparently low set ears and compressed pinnae, flat profile, epicantthic folds, mild telecanthus, flat and wide nasal bridge, normal palate, thin, downcurved lips, and mild micrognathia. The thorax and umbilical cord were small. The genital region was markedly abnormal (fig 2). The phallus-like appendage measured 2 x 0.5 cm and had rugation along the entire length, but lacked a urethral orifice, glans, palpable gonads, or scrotum. The rectum was imperforate. There was also a presacral phallus-like appendage measuring 1 x 3 cm with rugation of a labioscroital mound, a smooth glans, and no palpable gonads or urethral orifice (fig 3). There was right radial and thumb aplasia, right shoulder dimple with subluxed humerus, hypoplastic and subluxed left thumb, hypoplastic palmar creases, and bilateral transverse palmar creases. The lower extremities were tightly adducted, with short