Muscular Dystrophy (Duchenne) in a Girl with Turner’s Syndrome*

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According to classical genetics, a recessive sex-linked or X-linked gene manifests its existence phenotypically only in the absence of its normal allele, that is, in the homozygous state or in the absence of a second X chromosome carrying the normal allele. The latter requirement is met, for instance, in males who have haemophilia as a sex-linked trait: they have only one X chromosome in their XY chromosome complement. In females with the usual XX sex-chromosome composition a gene such as the haemophilia gene is not able to manifest itself, as it is neutralized by the presence of its normal allele on the second X. The situation is different in XO females: as in XY males, all recessive genes carried by the single X chromosome can freely express themselves.

The case reported here is that of a girl with Turner’s syndrome and Duchenne type muscular dystrophy. Chromatinnegative Turner’s syndrome is usually the consequence of a complete or partial (mosaic) XO sex-chromosome complement. Duchenne type muscular dystrophy usually appears to be sex-linked (Morton, Chung, and Peters, 1963; Boyer and Fainer, 1963; Walton, 1964). The present case offers cytogenetical evidence in support of sex linkage and of a permissive effect of abnormal sex-chromosome complement with regard to muscular dystrophy.

Case Report

This 7½-year-old girl was the youngest of three sibs. The parents were unrelated and healthy. The mother was 37 and the father 38 years old at the time of birth of the patient. On the paternal side, a first cousin of the patient, a boy, had a palatoschisis, and another first male cousin suffered from haemophilia. The mother’s side of the family appeared to be normal, but most of these relatives were living in another part of the country. The 14-year-old brother of the patient had had progressive pseudohypertrophic muscular dystrophy of the Duchenne type since the age of 7 and had been confined to a wheelchair since the age of 12.

The patient’s birth weight was 2,040 g., length 46 cm. Physical and motor development were slow: she sat up by 8 months of age and walked alone at 20 months. The parents were disturbed by her lack of appetite and extremely poor weight gain in infancy. Growth had always been very slow and regular. The patient was not having major difficulties at school; she rated 86 on the Wechsler intelligence scale. At 6 years of age the parents began to notice that the girl had some difficulty in climbing stairs, tired more easily on exercise, and fell down more often than before. These troubles were mild but progressed slowly.

Examination at 7½ years revealed the following features: stature (102 cm.) and weight (13 kg.) were well below the third percentile for this age (Fig. 1a and b). The lower segment (pubis to sole) was 46 cm., and the span was 98 cm. Her narrow pelvis and broad shoulders and chest, her leanness and the relative muscular prominence further contributed to give to her body the appearance of a miniature male adolescent. Muscular hypertrophy was particularly obvious at the calves. Salient facial features were downward eye slanting, a few pigmented naevi, large external ears, and a high-arched palate. Permanent incisors were present. The thyroid was not palpable. There was no webbing of the neck. There was no evidence of cardiovascular malformation, and the arterial blood pressure was 105/50 mm. Hg. External genitalia were normal female and infantile. The extremities were normal except for the impaired radial adduction of both wrists (Fig. 2a). Neurological findings were limited to generalized but moderate muscular weakness, with conservation of the deep tendon reflexes. The gait was normal but the patient needed a support to

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Muscular Dystrophy (Duchenne) in a Girl with Turner's Syndrome

pull herself up when climbing stairs. She exhibited the classic 'body building' manoeuvre with hands taking support on the thighs when rising up from reclining to upright position. Colour vision, using the Ishihara tables, was found to be normal in the patient as well as in the other members of the family.

Radiographic and Laboratory Data. Radiographs showed bone maturation to be consistent with chronological age. The skull was normal. Both wrists showed the so-called Madelung's deformity (Madelung, 1879) (Fig. 2b), i.e. a pronounced angulation of the distal epiphysis of the radii. An intravenous pyelogram revealed a normal excretory system.

Urinalysis and blood counts were within normal limits.

Enzyme Determinations. Serum creatine phosphokinase activity was measured in Dr Richterich's laboratory in Berne and was found to be increased to 49·0 U. (normal: 0 to 2·0 U.). The same determination was repeated 6 months later (at the age of 8) and gave a value of 102·0 U. The serum of the affected brother was tested also and an activity of 25·5 U. was demonstrated, whereas the healthy sister, the mother, and the father had 2·3 to 4·0 U. and 2·1 U. respectively.

Muscle Biopsy. A biopsy from the right gastrocnemius muscle showed small foci of muscle fibre necrosis with round cell phagocytosis and proliferation of histiocytes.

Blood Group Determinations. Blood samples from the whole family were sent to Dr R. Sanger in London for blood group determinations, particularly regarding the sex-linked Xg group. All members of this family were Xg (a−).

Cytological Data. Buccal smears from the patient were interpreted as showing a negative chromatin pattern. However, in a very small percentage of the nuclei a closer inspection revealed the presence of Barr bodies of smaller than average size (Fig. 3).

Tissue cultures of skin fibroblasts disclosed a mosaic chromosomal complement of the XO/Xx type, x standing for what was interpreted as a centric fragment of an X chromosome (Fig. 4, 5, and 6). However, the possibility of a ring X chromosome could not be excluded with certainty: 59 cells with 45 chromosomes were counted, and 11 of them were karyotyped, showing the XO complement; 10 cells with 46 chromosomes were counted, and all 10 were karyotyped, showing the Xx complement.

Short-term cultures from peripheral blood leucocytes also revealed the existence of the two cell lines. 61 cells were counted: one had 44 chromosomes, 39 cells had 45 chromosomes, 20 cells had 46, and one cell had 45 or 46. Karyotyping was carried out on 24 metaphase plates: 9 cells were of the 45-XO type, and 13 of the 46-Xx (or XXRinge) type. Two cells showed a comple-
FIG. 2a.

The wrist deformity due to the abnormal shape of the distal end of the radii (Madelung).

FIG. 2b.

The wrist deformity due to the abnormal shape of the distal end of the radii (Madelung).

Discussion

Turner's Syndrome. Turner's syndrome was diagnosed in this patient on the basis of dwarfism, a wide shield-like thorax, various facial anomalies, and the karyotype of 45 chromosomes with the Xx set, but in each of them a different autosome was missing, indicating that the complement of 45 in these cells was probably due to an artefact.
anomalies (Haddad and Wilkins, 1959; Lemli and Smith, 1963), and a malformation of the wrist (Kosowicz, 1962; Finby and Archibald, 1963). Sex chromatin and the karyotype findings substantiated the diagnosis. Though an XO chromosome complement remains the basic cytological characteristic of Turner’s syndrome, a large number of mosaic cases have now been described (Jacobs, Harnden, Buckton, Court Brown, King, McBride, MacGregor, and Maclean, 1961; De la Chapelle, 1962; Lindsten, 1963; Jones, Ferguson-Smith, and Heller, 1963; Engel, 1964). In the present instance most of the cells had a 45-XO complement, but there were a few 46 chromosome cells possessing one X chromosome plus a small chromosome, equal or inferior in size to the smallest acrocentrics of the 21–22 group (Fig. 7 and 8). In some cells this small chromosome had no visible centromere, but in others it had one with two very short arms and it could be interpreted as a deleted X chromosome. This interpretation was suggested by the fact that in the majority of the cells one of the X chromosomes was missing. The same factor responsible for the complete disappearance of one X may have been at play for the partial deletion of this chromosome in another cellular stem-line in the zygote. Moreover, there are now reports of a certain number of examples of gonadal dysgenesis or Turner’s syndrome with an identical mosaic karyotype (Blank, Gordon, and Bishop, 1961; Ferrier, Gartler, Waxman, and Shepard, 1962; De la Chapelle, 1962; Jones et al., 1963; Lindsten, 1963; Quinodoz, Ferrier, Ferrier, Zahnd, and Prod’hom, 1964). These patients may or may not exhibit as many malformative stigmata as the purely XO females.

**Duchenne Type Muscular Dystrophy.** The diagnosis of Duchenne type muscular dystrophy rested upon clinical features such as muscular weakness with muscular pseudohypertrophy, particularly at the calves, the progressive nature of the dystrophy, the positive family history, the serum creatine kinase determinations, and the muscle biopsy. As shown by Ebashi, Toyokura, Momoi, and Sugita (1959), Dreyfus, Schapira, and Demos (1960), and Aebi, Richterich, Stillhart, Colombo, and Rossi (1961), serum creatine phosphokinase activity is increased in Duchenne muscular dystrophy, particularly in the early stages of the disease, when the patient is still able to walk.

**Inheritance of Duchenne Muscular Dystrophy.** In the majority of affected families, the study of pedigrees provides evidence in favour of a sex-linked recessive inheritance for this type of muscular dystrophy (Stevenson, 1953; Walton, 1955, 1956). Further evidence comes from the fact that affected boys born of the same mother may be fathered by different men (Walton, 1956),
and from the study of a family in which a crossing-over had possibly occurred between this type of muscular dystrophy and red-green colour blindness (Philip and Walton, 1956).

In some uncommon pedigrees, however, there are affected females as well as affected males, and it is thought that the disease may sometimes be produced by an autosomal recessive gene (Lamy and de Grouchy, 1954; Kloepfer and Talley, 1958; Dubowitz, 1960; Jackson and Carey, 1961).

Recent findings still further complicate the issue. Studies in female carriers of the sex-linked recessive gene responsible for most instances of the disease have shown that over 70% of them can be identified by their moderately increased serum creatine kinase activity (Schapira, Dreyfus, Schapira, and Demos, 1960; Aebi, Richterich, Colombo, and Rossi, 1962; Hughes, 1962; Richterich, Rosin, Aebi, and Rossi, 1963). Van den Bosch (1963) and Barwick (1963) have reported

Fig. 4. A 45/XX cell (skin fibroblast, right calf).
that minor electromyographic anomalies suggestive of a myopathy may also be observed in carriers. Emery (1963) and Pearson, Fowler, and Wright (1963) have shown that minor clinical manifestations such as minimal muscular wasting and pseudohypertrophy of the calves may be present in certain carrier females. Dubowitz (1963a, b) demonstrated that these manifestations were correlated with typical histological changes in the muscles. These observations are in agreement with the Lyon hypothesis (Lyon, 1961, 1962) on the random inactivation of one of the X chromosomes in females. In carrier females a certain proportion of the muscle cells are in fact dystrophic,
due to the inactivation of the X chromosome carrying the normal gene, but this proportion is usually too small to give rise to clinically detectable muscular weakness (Pearson et al., 1963).

In the present case the pedigree was too rudimentary to ascertain that the disease was transmitted according to a sex-linked mode of inheritance. However, its occurrence in the same sibship in a boy and a XO/Xx mosaic girl, while the normal sister was free of the disease, might be considered as suggestive evidence in favour of the sex-linked transmission. A similar association of Turner's syndrome and Duchenne muscular dystrophy has been observed by Walton (1956) in a girl with a negative sex chromatin pattern. Apparently the karyotype of this patient was not established, but it was assumed to be XO and this example is cited by Walton (1964) as evidence in favour of sex-linkage.

Other but less likely explanations may be put
Muscular Dystrophy (Duchenne) in a Girl with Turner’s Syndrome

forward to account for this association, as can be deduced from the above quoted studies. Recessive autosomal inheritance cannot be absolutely ruled out from the pedigree; the association would then lack any cause and effect relationship. The other alternative would be to consider the patient as a carrier with minor clinical manifestations. Her muscle biopsy would be compatible with this hypothesis. However, her serum creatine phosphokinase activity fell within the values obtained in patients with Duchenne muscular dystrophy in the same laboratory (Aebi et al., 1961), and well above those of the carrier mother (3.2 U) and possible carrier sister (2.3 to 4.0 U). Finally, her clinical manifestations were more important than the muscle changes without weakness reported in some female carriers (Emery, 1963; Pearson et al., 1963). For these reasons, the hypothesis of a carrier state ought to be discarded.

Origin of Non-disjunction of the X-chromosome in Turner’s Syndrome. In Turner’s syndrome, studies on the inheritance of associated sex-linked characteristics such as colour blindness (Polani, 1961) and Xg blood group (Lindsten, Fraccaro, Polani, Hamerton, Sanger, and Race, 1963; Lindsten, 1963) point to the maternal origin of the single X in the majority of cases. If one admits the sex-linked nature of muscular dystrophy in the present patient’s family, then the patient is another example of Turner’s syndrome with the single X deriving from the mother, as shown by the clinical and pathological findings. Meiotic non-disjunction in the father accounts for the XO complement in most cases of XO Turner’s syndrome. On the other hand, in mosaic cases mitotic non-disjunction in the zygote is more likely to have occurred. If such is the origin of the mosaic in the present patient, one must admit that in mitotic non-disjunction of the X, the paternal X is preferentially lost (XO cells) or inactivated (Xx cells). So whether the origin of the non-disjunction is meiotic or mitotic does not make much difference: the paternal X is lost preferentially in Turner’s syndrome.

Probability Calculations. The manifestation of a sex-linked trait in a subject with an XO chromosome complement is a statistically predictable event, if the independence of the two is accepted. Its incidence can be calculated to be equal to the product of the frequency of XO females in the general population (0.2 to 0.5/1,000: Bergemann, 1961; Maclean, Harnden, and Court Brown, 1961; Court Brown, 1962; Nakagome, Hibi, Konoshita, Nagao, and Aikawa, 1963) by the frequency of the X-linked mutant in the same population (279/million for Duchenne muscular dystrophy: Morton et al., 1963). The chance of occurrence of both disorders in the same person, therefore, is of approximately 1/10 millions.

Summary

The case reported is that of a 7½-year-old girl who suffered from muscular dystrophy of the Duchenne type associated with dwarfism and other characteristics of Turner’s syndrome. Her chromosome complement in both skin fibroblasts and peripheral leucocytes was found to be a mosaic of XO cells and Xx (= X and a deleted X) cells. The family pedigree, though not absolutely conclusive, suggested a sex-linked or X-linked inheritance of the muscular dystrophy. The view is expressed that the loss of genetic material, total or sub total, from the X chromosome was responsible for the clinical manifestation of the disease in this phenotypic female. This is the second recorded instance of such an association. Furthermore, this case is thought to represent another example of Turner’s syndrome in which the paternal X has been preferentially lost or deleted.

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REFERENCES


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