Functional Xp disomy and de novo t(X;13)(q10;q10) in a girl with hypomelanosis of Ito

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Abstract
We report on a 16 month old girl with hypomelanosis of Ito and a balanced de novo (X;13)(q10;q10) translocation in which the der(X;13q) had the X centromere (as assessed by FISH with the DXZ3 probe). A functional Xp disomy was shown in a small proportion of cultured lymphocytes by means of a BrdU terminal pulse. This observation supports the notion of a distinct form of hypomelanosis of Ito resulting from a functional Xp disomy. (J Med Genet 1997;34:161–163)

Keywords: hypomelanosis of Ito; X;autosome translocation; functional Xp disomy.

Hypomelanosis of Ito (HI) is characterised by hypopigmented whorls, streaks, and patches typically distributed along the lines of Blaschko; abnormalities of the central nervous, eye, and musculoskeletal systems have also been described. When looked for, chromosomal abnormalities have been documented in nearly half of the patients and can be classified into two main groups: (1) various mosaicisms in about 20 patients with no instance of X;autosome translocation and a few XX/XY chimerisms, and (2) non-mosaic balanced X;autosome translocations in eight female patients (table 1). In the first group the pigmentary dysplasia seemingly reflects the clonal origin of chromosomally abnormal melanoblasts from neural crest precursors, whereas in the second group mosaicism for Xp functional disomy, rather than a disruption of a X linked gene, appears to be the causative mechanism.

Case report
The patient was the fourth child born to a 35 year old mother and an unrelated 38 year old father. The mother had had one abortion. Both parents and their three older children were normal; in particular, there was no family history of pigmentary anomalies. She was delivered at term by caesarean section because of pre-eclampsia; weight was 3110 g, length was 48 cm, and Apgar scores were 8 and 9 at one and five minutes respectively. At birth, streaky hypopigmented lesions were noted on the body. From 3 months of age she developed generalised tonic-clonic convulsions, which were unsuccessfully treated with phenobarbital. An EEG showed an abnormal generalised activity whereas a CT scan showed isodense zones in the frontal lobes. Ophthalmoscopy at 4 months showed nystagmus and hypoplasia of the fovea more apparent in the left eye. Physical examination at 16 months of age (fig 1) showed weight 10.5 kg (90th-97th centile), length 85 cm (above the 97th centile), OFC 44 cm (10th-20th centile), hypotonia, no head control, nystagmus, synphrys, upward slanting palpebral fissures, dysplasia of the teeth, widely spaced nipples, and irregular streaky hypopigmented lesions with a mottled, streaked, or whorled appearance on both sides of the trunk and the extremities; vesicobullous or verrucous lesions were not noted.

Radiographs showed brachycephaly, dorso-lumbar scoliosis, hypoplastic ischia, and an asynchronous accelerated bone age (pelvis 4 years, hands between 2 and 3 years).

CYTOGENETIC STUDIES
A peripheral blood karyotype (G bands) was 46,X,t(X;13)(Xp13q;Xq13p) in all 50 cells examined (fig 2A). The patients' karyotype was normal. A replication X study with a BrdU terminal pulse (n=150 cells) showed an inactive normal X chromosome as the predominant pattern, but in two cells (1.3%) the der(Xq13p) was inactive whereas the normal X and the der(Xp13q) were early replicating. FISH was performed according to the manu-

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<th>Table 1</th>
<th>Hypomelanosis of Ito with X;autosome translocations</th>
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<td>Sex</td>
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<td>F</td>
<td>5 y</td>
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<td>21 y</td>
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L = lymphocytes, T = tumour cells (pexus papilloma), S = skin fibroblasts.
facturer's protocol (Oncor) with a digoxigenin labelled X centromere probe (DXZ3). The sites of hybridisation were visualised using either rhodamine labelled antidigoxigenin/DAPI or fluorescein labelled antidigoxigenin/propidium iodide. We observed the hybridisation signal on the normal X chromosome and on the der(Xp13q) chromosome (fig 2B). Therefore, the X breakpoint was assigned to q10. A cell line of this patient is not available.

Discussion

The balanced X:autosome translocations found in nine unrelated females with sporadic HI (table 1) exhibit some distinct characteristics, namely non-mosaic occurrence, a pericentromeric or centromeric X chromosome breakpoint, apparent randomness of the autosomes involved, and, apart from one case, a de novo origin. Therefore, these patients may constitute a distinct form of HI. The skewed X inactivation in the present case conforms to previous analyses in cultured cells but the alternate pattern documented in a small proportion of cells confirms previous observations in either cultured lymphocytes or uncultured skin cells. As for the causative mechanism, the work of Hatchwell et al points to a mosaic Xp functional disomy rather than to a gene disruption, a conclusion supported by our findings. In fact, the six other translocations in which the X chromosome functional disomy was not assessed are consistent with this view, that is, the X chromosome breakpoint in all of them allows for the occurrence of the alternate inactivation pattern in which the derivative with the X inactivation centre is inactivated and so these cells are functionally disomic for Xp sequences. Since a functional non-mosaic Xp disomy does not result in a HI phenotype (for example, Gustashaw et al), it may be inferred that Xp disomy, like other imbalances (triploidy, trisomy 18, and tetrasomy 12p), only causes such a phenotype when occurring in a mosaic state. Moreover, the absence of male patients with HI and a balanced X:autosome translocation, if not a random event, is in keeping with this, as X chromosome disomy cannot occur in them. The single inherited X:autosome translocation may be accounted for by a more stringent selection in the mother whose nearly normal phenotype facilitated the transmission of the translocation to her affected daughter.

According to the banding patterns, the present translocation was regarded as a whole arm exchange, yet FISH results indicated that the X chromosome breakpoint was outside the centromere. Hence, this rearrangement differs from the single true whole arm translocation associated with HI.

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1 Ito M. Studies on melanin. XI. Incontinentia pigmenti achromians: a singular case of nevus depigmentosus.


