De novo deletion of Xp22.2—pter in a female with linear skin lesions of the face and neck, microphthalmia, and anterior chamber eye anomalies

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Abstract
A female infant is described with an unusual combination of eye and skin findings. Raw linear skin lesions on the face and neck were present at birth, healing to leave pigmented streaks. In addition she had left sided microphthalmia and bilateral sclerocornea. Chromosome analysis showed a terminal deletion of the short arm of the X chromosome (Xp22.2—pter). Clinical findings were similar to three previously described children with translocations involving Xp22.3. The condition probably represents a new syndrome distinct from incontinentia pigmenti and Goltz syndrome.

We describe a patient with a deletion of Xp22.2—pter with skin and eye changes distinct from incontinentia pigmenti (IP) and Goltz syndrome. Three cases have been reported by Al-Gazali et al with a similar pattern of anomalies and disruption of Xp22.3. Our case provides further evidence of a new syndrome.

Case report
The female proband was the first child of non-consanguineous Nigerian parents aged 23 (mother) and 29 (father) years. She was born at term with a birth weight of 4500 g after a normal pregnancy and delivery. Raw linear lesions were noted at birth on the face and neck. They resembled scratches and did not

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blister. The rest of the skin was normal. In addition, both eyes were swollen but left sided microphthalmia was suspected. By two months the skin lesions on the face healed to leave pigmented scars.

The proband presented to our genetic clinic aged 20 months when her mother was 28 weeks pregnant. Examination at that time showed healed but hyperpigmented linear scars on the face and neck (fig 1). One lesion extended onto the left shoulder, but the skin was otherwise normal apart from a small café au lait lesion on the trunk. Ophthalmological examination confirmed left microphthalmia associated with bilateral sclerocornea. She had roving eye movements and vision was thought to be restricted to light perception only. Development appeared to be normal taking into consideration her visual handicap; she sat at six months, walked at 18 months, and had many single words.

Blood was taken for chromosome analysis. This showed a deletion of the terminal zone of the short arms of the X chromosome involving the region p22.2 to the telomere (46,X,del(X)(p22.2-pter) (fig 2). Parental chromosomes were normal. The patient left the country before it was possible to look at her pattern of X inactivation.

Discussion

The differential diagnosis of this combination of asymmetrical skin and eye anomalies in a female child include the syndromes of IP and Goltz. They are both multisystem disorders affecting structures of ectodermal and mesodermal origin and the inheritance is thought to be X linked dominant with lethality in males.

The skin lesions in IP classically progress from

### Comparison of features in our patient with those of IP and Goltz syndrome.

<table>
<thead>
<tr>
<th></th>
<th>Incontinentia pigmenti</th>
<th>Goltz syndrome</th>
<th>Xp22.2 deletion/disruption</th>
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</thead>
<tbody>
<tr>
<td>Skin</td>
<td>Bullous lesions at birth over any part of body. Later, verrucous eruptions and hyperpigmented whorls</td>
<td>Linear dermal hypoplasia with fat herniation</td>
<td>Acute weeping linear lesions on face and neck at birth. Later, hyperpigmented streaks.</td>
</tr>
<tr>
<td>Eye</td>
<td>Posterior chamber abnormalities, 'chorioretinitis', 'retrorenal fibroplasia'</td>
<td>Anterior chamber defects, microphthalmia, coloboma of iris</td>
<td>Microphthalmia, sclerocornea, orbital cysts</td>
</tr>
<tr>
<td>Teeth</td>
<td>Normal to anodontia</td>
<td>Hypoplasia and abnormal eruption pattern</td>
<td>Normal</td>
</tr>
<tr>
<td>Other</td>
<td>Cleft lip and palate, alopecia, syndactyly</td>
<td>Diaphragmatic hernia, exomphalos, syndactyly/oligodactyly, angiofibromata</td>
<td>—</td>
</tr>
<tr>
<td>Development</td>
<td>30% CNS abnormalities, seizures, MR, hydrocephalus</td>
<td>Occasional mental retardation. Usually normal development</td>
<td>Normal</td>
</tr>
<tr>
<td>Inheritance</td>
<td>X linked dominant mapped to Xq28</td>
<td>X linked dominant, not mapped.</td>
<td>? X linked dominant. Associated with deletion involving Xp22—pter.</td>
</tr>
</tbody>
</table>

Figure 2  Pairs of G banded X chromosomes showing the extent of the short arm deletion of the X (left) compared to the normal X (right). The deletion involves Xp22.2—pter.
bullae, present at birth or soon after, to verrucous eruptions and later hyperpigmented whorls. The abnormal eye findings usually involve the posterior chamber and resemble chorioretinitis or retrolental fibroplasia. Anterior chamber anomalies are rare in this condition.

The gene for IP has recently been assigned to Xq28 by linkage analysis of eight families with two or more affected females. This is contrary to four previous reports of X;autosome translocations involving Xp11. Goltz syndrome (focal dermal hypoplasia) was first reported as a separate condition in 1962 by Goltz et al. These affected girls have linear areas of dermal hypoplasia, which may be associated with areas of fat herniation and abnormal pigmentation. The eye is involved in 20% of reported cases and the lesions tend to involve the anterior segment. Iris colobomata and microphthalmia are common eye findings in this condition. The teeth, skeletal system, and nails can be involved but features are very variable. No linkage data have been reported.

Our patient has features in common with both IP and Goltz syndrome (table). However, lack of early bullous and verrucous skin findings and the anterior chamber eye anomalies make this unlikely to be IP. Her eye findings would be consistent with Goltz syndrome, but the skin changes are not typical of focal dermal hypoplasia, are only confined to the face, and are not associated with fat herniations. Also, she does not have any of the other associated features of this condition.

Al-Gazali et al described three cases with similar clinical features to our patient: irregular linear areas of erythematous skin hypoplasia involving the head and neck with microphthalmia, corneal opacities, and orbital cysts. All had normal development. Two of these patients were females with de novo X;Y translocations and the other case was a 46,XX male in whom DNA studies indicated exchange of material between the short arms of the X and Y chromosomes. Cytogenetic analysis put the breakpoint on the X chromosome at Xp22.3.

We consider that our patient has the same condition. We believe that it is probably distinct from IP and Goltz syndrome and represents a newly recognised combination of eye and skin findings owing to a disruption of a gene distal to Xp22.2.

References: