CURRENT TOPIC

Childhood hypertrichosis: diagnosis and management

F A M Baumeister, H P Schwarz, S Stengel-Rutkowski

There is wide variation in the normal pattern of hair growth. Hypertrichosis, which can be defined as excessive growth of hair compared with that in other subjects of the same age, sex, and race, must be distinguished from hirsutism, a term restricted to an androgen dependent hair pattern that is characterised by excessive hair growth on the upper lip, chin, chest, lineae albae, thighs, and axillae. Unlike hypertrichosis, unexplained hirsutism in childhood usually warrants investigation to exclude an endocrine cause for the virilisation.

In generalised hypertrichosis there is accentuation of facial hair in the frontal, temporal, and preauricular regions. The eyebrows may be bushy or confluent. On the back of the trunk the hair converges on the midline, often forming whorls over the spine. It may occur as part of a syndrome or metabolic disorder ('symptomatic hypertrichoses') as opposed to 'congenital hypertrichosis' where markedly excessive hair growth is the most prominent feature.

While this review will focus on generalised hypertrichosis in childhood, it should be noted that localised hypertrichosis can occur and may be related to naevi or spina bifida occulta, previous trauma or chemical irritations, and in a few inherited conditions such as hairy elbows, hairy ears, hairy nose tip, or hairy palms and soles.

Symptomatic hypertrichosis in childhood

ASSOCIATION WITH DYSMORPHIC SYNDROMES

A characteristic facial appearance in a child with hypertrichosis may lead to the recognition of one of a number of dysmorphic syndromes (table), for example Brachmann-de Lange syndrome, Coffin-Siris syndrome, Rubinstein-Taybi syndrome, Seckel's syndrome, cerebro-oculofacioskeletal syndrome, Gorlin's syndrome, Schinzel Giedion midface retraction syndrome, or Barber Say syndrome.

Association with acro-osteolysis may result in diagnosis of the Hajdu Cheney syndrome or postaxial polydactyly to diagnosis of the Weyers' acrofacial dysostosis syndrome. Hypertrichosis associated with osteochondrodysplasia or gingival fibromatosis may indicate the presence of other genetic entities. Association with photosphobia may indicate amaurosis congenita (cone-rod type) with congenital hypertrichosis.

In the newborn hypertrichosis associated with markedly reduced subcutaneous fat may indicate a diagnosis of leprechaunism, a lethal condition which is also associated with an unusual facial appearance, severe intrauterine and postnatal growth retardation, and hyperinsulinaemia with hyperplasia of the pancreatic

Symptoms associated with generalised hypertrichosis

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>MIM Number</th>
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<tbody>
<tr>
<td>Brachmann-de Lange syndrome</td>
<td>122470</td>
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<tr>
<td>Coffin-Siris syndrome</td>
<td>139500</td>
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<tr>
<td>Rubinstein-Taybi syndrome</td>
<td>268600</td>
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<tr>
<td>Seckel’s syndrome</td>
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<tr>
<td>Cerebro-oculofacioskeletal syndrome</td>
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<td>Gorlin’s syndrome</td>
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<tr>
<td>Schinzel Giedion midface retraction syndrome</td>
<td>269150</td>
</tr>
<tr>
<td>Barber Say syndrome</td>
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<tr>
<td>Hajdu Cheney syndrome</td>
<td>193530</td>
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<tr>
<td>Weyers' acrofacial-dysostosis syndrome</td>
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<tr>
<td>Osteochondrodysplasia with hypertrichosis</td>
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<td>Gingival fibromatosis with hypertrichosis</td>
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<tr>
<td>Amaurosis congenita (cone-rod type) with hypertrichosis</td>
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<tr>
<td>Leprechaunism</td>
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<tr>
<td>Patterson's syndrome</td>
<td>169170</td>
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<tr>
<td>Seip's syndrome</td>
<td>269700</td>
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<tr>
<td>Partial trisomy 3q syndrome</td>
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β cells as a result of an insulin receptor defect. Pseudoleprechaunism (Patterson's syndrome) is also associated with hypertrichosis but is distinguished from leprechaunism by a normal birth weight, large hands and feet, cutis gyrata, and skeletal anomalies. In Seip's syndrome hypertrichosis is associated with lipodystrophy, muscular hypertrophy, increased stature and non-ketotic insulin resistant diabetes.

ASSOCIATION WITH METABOLIC OR CHROMOSOMAL DISORDERS OR PRENATAL AND POSTNATAL DRUG EXPOSURE

Metabolic disorders associated with hypertrichosis include the mucopolysaccharidoses,4 GM1-gangliosidosis,5 and porphyria. Among the chromosomal disorders, hypertrichosis is most prominent in partial trisomy 3q.6 Hypertrichosis may also be a feature of patients with anorexia nervosa.2 Hypertrichosis may result from maternal alcohol abuse in pregnancy,4 as well as prenatal or postnatal exposure to hydantoins4 or minoxidil.7 Treatment with cyclosporin or diazoxide also leads to hypertrichosis, as does treatment with glucocorticoids.

Congenital hypertrichosis

Congenital hypertrichosis universalis is a very rare genetic condition in which the whole body is covered by excessive fine, light coloured hair that can reach a considerable length.8 9 The condition was erroneously thought to be associated with increased mortality because the first reported case of leprechaunism was described (and subsequently cited) as congenital hypertrichosis universalis.10-15 To confuse matters further, the terms congenital hypertrichosis universalis, congenital hypertrichosis lanuginosa, and hypertrichosis lanuginosa have been used synonymously,9 11 12 14 16-23 although they cover three different clinical entities. The three subtypes are believed to result from mutation of an autosomal dominant gene, but differ with regard to the persistence and pattern of hypertrichosis, and the associated anomalies.

In the Ambras' syndrome24 generalised hypertrichosis is present at birth (fig 1) and persists for life. The hair is most abundant on the face, ears, and shoulders (fig 2) and this becomes more accentuated with increasing age; this unique pattern allows differentiation from the other hypertrichosis syndromes. If not shaved the hair reaches a considerable length (fig 2). Abnormalities of the teeth, accessory nipples, and preauricular regions may be associated findings.

Transient congenital hypertrichosis universalis is present at birth but disappears during infancy and is characterised by generalised hypertrichosis which spares the face, hands and feet (fig 3).18 19 22 The condition was associated with a neonatal tooth in one report22 and with congenital glaucoma in another.19

In persistent hypertrichosis universalis affected subjects are only slightly hairy at birth but increasing hairiness occurs in infancy.9 12 14 16 20 In contrast to Ambras' syndrome, the facial hair is not uniformly distributed but is accentuated in the frontal, temporal, and preauricular regions. No other associated abnormalities have been reported.

Management of hypertrichosis

The need for treatment depends on the degree of hypertrichosis and the resulting psychosocial
problems. In an excessively hairy newborn early removal of hair may be necessary because of difficulties the family may have in accepting the child, leading to social isolation.21 23 24 Removal of hair is also needed to allow cleaning of the nappy area.22

There are several ways of removing hair. Chemical depilatories are effective but repeated use leads to irritation of the skin and may cause contact dermatitis. Wax epilation or depilatory plasters are painful and remove fine vellus hair which may induce transformation to coarse terminal hair, giving the impression of increased hairiness.2 Electrolytic destruction of individual hair papillae removes some hair permanently but up to 30% of the hair papillae treated in any one session regrow.2 There is also a risk of scarring after destruction of the deep dermal papillae. For these reasons electrolysis cannot be recommended in children with generalised hypertrichoses and repeated shaving remains the treatment of choice.9 11 21-24

We thank Dr J Sigalas for the photograph in fig 1 and Dr J W Partridge for permission to reproduce fig 3.

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