Conotruncal anomaly face syndrome is associated with a deletion within chromosome 22q11

J Burn, A Takao, D Wilson, I Cross, K Momma, R Wadey, P Scambler, J Goodship

Abstract
The conotruncal anomaly face syndrome was described in a Japanese publication in 1976 and comprises dysmorphic facial appearance and outflow tract defects of the heart. The authors subsequently noted similarities to Shprintzen syndrome and DiGeorge syndrome. Chromosome analysis in five cases did not show a deletion at high resolution, but fluorescent in situ hybridisation using probe DO832 showed a deletion within chromosome 22q11 in all cases.

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Kinouchi et al.1 in 1976 reported a recognisable phenotype, the conotruncal anomaly face (CTAF) syndrome, comprising a variety of cardiac outflow tract defects and characteristic facies with hypertelorism, lateral displacement of the inner canthi, a flat nasal bridge, narrow palpebral fissures, a nasal voice, and minor ear anomalies. In subsequent analysis2 it was suggested that this was a frequent phenotype among patients with outflow tract defects such as tetralogy of Fallot. The authors subsequently noted the heterogeneity of this syndrome and its association with thymic involution.3 They noted the overlap with the syndrome later described by Shprintzen et al.4 and subsequent reviews noted the likelihood that these syndromes were differing manifestations of the same phenotype.5 Following the demonstration of 22q11 deletion in almost all cases of DiGeorge syndrome6 and Shprintzen syndrome7,8 and in familial heart defects9 it was decided to investigate children with the CTAF syndrome for this chromosome abnormality. We report the investigation of five affected subjects by high resolution karyotype analysis and chromosome fluorescent in situ hybridisation (FISH).

Methods
Five patients, two males aged 3 and 24 years and three females aged 7, 9, and 10 years, with CTAF syndrome were reviewed and a peripheral venous blood sample obtained. All five had tetralogy of Fallot with pulmonary atresia and a major aortopulmonary collateral artery, typical features of CTAF syndrome.12 The facial appearances of the two older girls are shown in fig 1. The one adult examined (fig 2A) had mild intellectual impairment with an intelligence quotient 2 SD below the mean. There was some uncertainty over whether this subject fulfilled the dysmorphic diagnosis, but subsequent review of a picture taken at 4 years (fig 2B) showed facial features consistent with the diagnosis.

High resolution cytogenetic analysis was attempted in each case on cell cultures established within 24 hours from the blood sample using a technique previously described.13 High resolution banding was achieved if band 22q11.22 could be visualised in one of the pair of chromosomes 22 (Standing Committee on Human Cytogenetic Nomenclature).

FISH was performed by a method previously described. Briefly, probe DO832 was labelled with digoxigenin by nick translation. This was hybridised to denatured chromosome preparations. The hybridised probe was then detected using fluorescein labelled antidigoxigenin antibody. The chromosomes were stained with propidium iodide and then visualised using a Nikon fluorescent microscope.

FISH analysis was performed on each patient with CTAF and also two normal subjects. Twenty metaphases from each patient and normal subject were analysed.

Results
High resolution chromosome banding was achieved in two of the five CTAF cases and both had a normal karyotype. In the remaining three cases the resolution was insufficient to identify band 22q11.22 and therefore the presence of a deletion could not be excluded. FISH analysis of the two normal subjects detected fluorescent signal on both chromosomes 22 in

Figure 1 Facial appearance of the two older girls aged 9 and 10 years.
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Discussion
We have shown that CTAF syndrome is associated with a deletion within chromosome 22q11. Similar deletions have been identified in subjects with DiGeorge and Shprintzen syndromes. The precise nature of the deletions within this region of chromosome 22 have not been shown to be identical in each of these syndromes. These results support the clinical suspicion, however, of a phenotypic overlap indicating a common aetiology. The facial features summarised by Takao et al (table) correspond closely to those in DiGeorge syndrome reviewed in this issue.14

Deletions within chromosome 22q11 may be responsible for a significant proportion of cases of cardiac defects either within a syndromic pattern or in apparent isolation. This has important implications for estimating the risk of recurrence in patients with cardiac defects.

The characteristics of conotruncal anomaly face syndrome of 50 cases.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Small mouth</td>
<td>44/50</td>
</tr>
<tr>
<td>Ocular hypertelorism</td>
<td>43/50</td>
</tr>
<tr>
<td>Mild lateral displacement of inner canthi</td>
<td>42/50</td>
</tr>
<tr>
<td>Short palpebral fissures</td>
<td>26/50</td>
</tr>
<tr>
<td>Boated eye lids</td>
<td>41/50</td>
</tr>
<tr>
<td>Low nasal bridge</td>
<td>43/50</td>
</tr>
<tr>
<td>Strabismus</td>
<td>5/50</td>
</tr>
<tr>
<td>Ptosis of eyelid</td>
<td>2/50</td>
</tr>
<tr>
<td>Nasal voice</td>
<td>39/47</td>
</tr>
<tr>
<td>High arched palate</td>
<td>25/47</td>
</tr>
<tr>
<td>Cleft soft palate</td>
<td>1/47</td>
</tr>
<tr>
<td>Malformed auricles</td>
<td>35/50</td>
</tr>
<tr>
<td>Prominent ears</td>
<td>19/50</td>
</tr>
<tr>
<td>Incomplete scapha helix development</td>
<td>14/50</td>
</tr>
<tr>
<td>Low set ears</td>
<td>6/50</td>
</tr>
<tr>
<td>Moderate conductive deafness</td>
<td>2/50</td>
</tr>
<tr>
<td>Mild mental retardation</td>
<td>18/50</td>
</tr>
</tbody>
</table>

The belief that DiGeorge, Shprintzen, and CTAF syndromes are distinct may ultimately be seen to be the result of each group's special area of expertise in, respectively, immunology, craniofacial malformation, and cardiology. It seems increasingly likely that these eponymous syndromes will be phenotypic variations on a theme.

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