Cryptic terminal rearrangement of chromosome 22q13.32 detected by FISH in two unrelated patients

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Abstract

Two unrelated patients with cryptic subtelomeric deletions of 22q13.3 were identified using FISH with the commercially available Oncor probe, D22S39. Proband 1 was found to have a derivative chromosome 22 resulting from the unbalanced segregation of a t(1;22)(q44;q13.32) in her mother. Additional FISH analysis of proband 1 and her mother placed the breakpoint on chromosome 22 in this family proximal to D22S55 and D22S39 and distal to D22S45. We have mapped D22S39 to within 170 kb of D22S21 using pulsed field gel electrophoresis. D22S21 is genetically mapped between D22S55 and D22S45. These data indicate that the deletion in proband 1 is smaller than in eight of nine reported del(22)(q13.3) patients. Proband 1 and 2 share features of hypotonia, developmental delay, and expressive language delay, also seen in previously reported del(22)(q13.3) patients, although proband 1 appears to be more mildly affected. Proband 1 is also trisomic for the region 1q44–qter. This very small duplication has been previously reported only once and the patient had idiopathic mental retardation. This is the first report where 22q13.3 terminal deletion patients have been identified through the use of FISH, and the first report of a deletion of this region occurring because of missegregation of a parental balanced cryptic translocation. We feel that investigation of the frequency of del(22)(q13.3) in the idiopathic mentally retarded population is warranted and may be aided by the ability to use a commercially available probe (D22S39), which is already currently in use in a large number of cytogenetic laboratories.

Keywords: del(22)(q13.3); cryptic subtelomeric rearrangement; idiopathic mental retardation; D22S39

It has been estimated that at least 6% of idiopathic mental retardation is caused by cryptic subtelomeric chromosome rearrangements.1 The recent description of several patients with deletions of the subtelomeric region of the long arm of chromosome 22 suggests that rearrangements of this region may account for an important subset of this group of patients. To date, we are aware of at least 15 patients reported to have de novo subtelomeric deletions involving 22q13.3. Thirteen patients were identified through high resolution cytogenetics,2,3 while two were identified with Southern blot screening.4

We now describe the use of a widely available probe for the detection of this group of patients by FISH analysis. Using the commercially available Oncor probe (D22S39), included as a control with the probe (D22S75) for DiGeorge syndrome, we have recently identified two additional unrelated patients with deletions of 22q13.3.

This is the first report where 22q13.3 terminal deletion patients have been identified through the use of FISH, and the first report of a deletion of this region occurring because of missegregation of a parental balanced cryptic translocation.

Subjects and methods

CLINICAL REPORTS

Case 1 (proband 1) presented at 22 months of age for evaluation of developmental delay. She was born a product of a 37 week gestation to a 35 year old G2, P0 woman. Concern was first raised at 4 months when she was not rolling over, but she has made developmental progress. She walked at 16 months of age. At 22 months, she had three words, but had appropriate receptive language. At 2 years of age, her weight was 12.6 kg (50th centile), length 84 cm (25th centile), and head circumference was 48 cm (50th centile). She had bilateral epicanthic folds, a haemangioma on her forehead, and full cheeks (fig 1). She had a small umbilical hernia. She was moderately hypotonic, walked with a waddling gait, and was clumsy, but no ataxia was noted. Reflexes were present and toes were flexor. An MRI performed at 30 months of age was normal. The family history was non-contributory except for a seven year period of infertility in her parents.

Case 2 (proband 2) is a 13 month old girl who presented with developmental delay. She was the 2665 g (10th centile) product of a term gestation born by vaginal delivery to a 21 year old G2, P1 woman. Fetal movements were not noted until 7 months and were reported to be less vigorous than the previous pregnancy. In the newborn period, she was noted to have a small ventricular septal defect which closed spontaneously, feeding difficulties, and slow
Subtelomeric rearrangement of 22q13.32

weight gain. She has vescicoureteral reflux without anatomical abnormality. Developmentally, she sat at 11 months, pulled to stand at 12 months, and babbled but did not have any words. On physical examination, her length was 79.5 cm (95th centile), weight was 9.8 kg (50th centile), and her head circumference was 45.8 cm (50th centile). There were no unusual facial features. She had a 6 x 3 cm irregular macule on the medial surface of her leg. She had decreased tone, but was able to sit unsupported and stand holding on. Reflexes were present and normal except for clonus at the ankles. An MRI showed mild delay in myelination with some thinning of the corpus callosum. The family history was unremarkable.

CYTOGENETIC AND MOLECULAR STUDIES
Peripheral blood lymphocytes were initially examined by G banded chromosome analysis. Fibroblast lines from patient FB and his parents, previously reported by Phelan et al.,11 were obtained from Dr Harold Taylor at the Greenwood Genetics Center, Greenwood, South Carolina. Metaphase preparations were G banded with the trypsin method and stained with Wright stain.2 FISH studies were done using D22S75, DiGeorge chromosome region (DGCR) probe with D22S39 chromosome 22 control probe, obtained from Oncor, Gaithersburg, MD (Cat No PS140-DIG). Hybridisation was carried out according to the manufacturer’s recommendations using the rapid wash conditions and no signal amplification.

FISH studies carried out to define the extent of missing chromosome 22 material were done using yeast strains containing YACs M412H3 and M120E4 obtained from Dr Callum Bell at The Children’s Hospital of Philadelphia.1 In addition, a 14/22 alpha satellite probe (Oncor, Inc) was used to identify the chromosomes 22. Total yeast genomic DNA was labelled with biotin-14-dATP using a BioNick labelling system (BRL) and FISH was performed according to the method described by Pinkel et al7 with some modifications: 1 μg of labelled yeast genomic DNA and 5 μg Cot-1 DNA were used per hybridisation, slides were denatured at 70°C in 70% formamide, 2 x SSC, pH 7.0, and hybridised for 48 hours at 37°C; posthybridisation wash was 50% formamide, 2 x SSC, pH 7.0, at 40°C.

Pulsed field gel electrophoresis (PFGE) was performed using a CHEF-DRII apparatus (BioRad). Cells from three subjects normal for the 22q13.3 region were embedded in agarose and processed as previously described.4 DNA was digested with NotI, NruI, Ascl, or BstWI and separated by PFGE at 150 V using a 120 second switch time for 22 hours followed by a 180 second switch time for 22 hours. The resulting blot was probed with the clone for D22S21. The blot was then stripped of signal and reprobed with the clone for D22S39.

Results

CASE 1
Cytogenetic analysis was performed on proband 1 as well as FISH analysis to rule out Prader-Willi/Angelman syndrome because of her hypotonia. No deletion of D15S10 or SNRPN was found. A subtelomeric abnormality in the terminal band of chromosome 22q was suspected after routine cytogenetic analysis (band level 650) but the alteration in the G banding pattern was sufficiently subtle that without additional confirmation the exact nature of the alteration could not be established (fig 2A). To define the nature of the suspected abnormality more precisely, we carried out FISH analysis using the Oncor DiGeorge probe containing the control probe (D22S39) localised to the subtelomeric region (q13.3) of chromosome 22. Hybridisation of this cosmids mixture to metaphase spreads from proband 1 showed a deletion of cosmid D22S39. Two signals (proximal and distal) were seen on one chromosome 22 and only one signal (proximal) on the other chromosome 22 in 20/20 metaphase spreads examined (fig 3A). This result together with the G banding interpretation resulted in the karyotype of proband 1 being originally reported as 46,XX,del(22)(q13.33).ish del(22)(q13.3q13.3) (D22S39-) or 46,XX,add(22)(q13.32).ish del(22)(q13.3q13.3)(D22S39-). FISH analysis of metaphase spreads from the mother of proband 1 with the DGCR probe mixture showed two signals on one chromosome 22, one signal (proximal) on the other chromosome 22, as well as one signal at either the p or q terminus of chromosome 1 (fig 3B). Thus, the mother of proband 1 was carrying a balanced cryptic translocation involving the D22S39 cosmids region. G banding analysis showed that the translocation involved an
exchange of material from the termini of the long arms of one chromosome 1 and one chromosome 22, 46,XX,t(1;22)(q44;q13.32).ish t(1;22)(D22S39+;D22S39-) (fig 2B). Her daughter, proband 1, inherited only her derivative chromosome 22 resulting in monosomy for the region 22q13.32 to 22qter and trisomy for 1q44. The karyotype of proband 1 was revised in light of this finding to 46,XX,der(22)t(1;22) (q44;q13.32)mat.ish der(22)t(1;22)(D22S39-; D22S39-). The father of proband 1 and one maternal aunt were found to have normal karyotypes.

Additional FISH analysis with chromosome 22q terminal YAC probes further localised the breakpoint for this rearrangement. Hybridisation with YAC M412H3 (which contains the DNA marker D22S55) resulted in a fluorescent signal on only one chromosome 22qter in metaphase spreads from proband 1. When metaphase spreads from her mother were assayed, a signal was seen on one chromosome 22qter as well as at the terminal end of one chromosome 1 (fig 3C). This result localises the marker D22S55 distal to the breakpoint in this family (fig 2C). Hybridisation with YAC M120E4 (which contains marker D22S45) resulted in fluorescent signals at the terminus of both chromosomes 22 in metaphase spreads from proband 1 and her mother (fig 3D). This result localises the marker D22S45 proximal to the breakpoint in this family (fig 2C).

CASE 2
Routine cytogenetic analysis was performed on proband 2. A subtelomeric deletion in the ter-
Table 1  Features of del(22)(q13.3) patients

<table>
<thead>
<tr>
<th>Features</th>
<th>Patients described in refs 1, 2, and 3*</th>
<th>Proband 1 (this study)</th>
<th>Proband 2 (this study)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Height and weight appropriate for gestational age (not IUGR)</td>
<td>12/12</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Hypotonia</td>
<td>10/13</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Developmental delay</td>
<td>15/15</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Delay of gross motor milestones</td>
<td>13/15</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Age at 1st step: 20.5–39 months</td>
<td>9</td>
<td>16 months</td>
<td>Not walking at 13 months</td>
</tr>
<tr>
<td>Expressive language status</td>
<td>Delay (4)</td>
<td>&lt;10 words at 2½ y</td>
<td>Babbles at 13 months</td>
</tr>
<tr>
<td></td>
<td>No words/babbles (7)</td>
<td>&lt;10 words (2)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Too young to ascertain</td>
<td>(2)</td>
<td></td>
</tr>
<tr>
<td>Mild dilatation of cerebral ventricles</td>
<td>5/12</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Seizures</td>
<td>3/12</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Dolichocephaly</td>
<td>7/12</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Ptosis</td>
<td>4/12</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Epicanthic folds</td>
<td>8/12</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Dysplastic ears</td>
<td>10/13</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Tall stature</td>
<td>3/14</td>
<td>No</td>
<td>95%</td>
</tr>
<tr>
<td>Vesicoureteral reflux</td>
<td>1†</td>
<td>No</td>
<td>Yes</td>
</tr>
</tbody>
</table>

*Number in denominator reflects whether the feature was mentioned in the report, please note that patient A in Zwaigenbaum et al 1 is the same as QM in Nesslinger et al 2.
†Herman et al. 3

Discussion

The clinical features in these two probands with monosomy for 22q13.3–22qter are similar to those described in previously reported cases of 22q13.3 deletions 1–3 (table 1). Proband 1 appears to be more mildly affected in that she walked earlier (16 months), speaks a few words, and does not have any dysmorphic features other than epicantic folds (table 1). This would seem to correlate with the fact that her breakpoint is located between D22S45 and D22S39, distal to the breakpoints defined for the patients described in Nesslinger et al 2 (fig 2C). Enzyme studies measuring arylsulphatase A (ARSA) activities in proband 1 and her parents were consistent with deletion of this locus in proband 1 (data not shown). A pulse field gel electrophoresis map of 22q13.3 (H McDermid, unpublished data) gives a rough estimate (a gap exists between H91 and ARSA which cannot be measured) of a distance of over 1800 kb from D22S39/D22S21 to the telomere. The only reported 22qter deletion patient (NT) with a similar deletion than proband 1 is reported to be non-dysmorphic with normal
motor milestones, mild mental retardation, and first words at the age of 4 years. His deletion has been reported to be approximately 120 kb in size, with the breakpoint located distal to ARSA. Proband 1, in addition to monosomy for 22q13.32→22qter is trisomic for the region 1q44→1qter. A review of published reports identified only one previous report of dup(1)(q44) in which the proband had idiopathic mental retardation with no associated malformations. A review of patients carrying a duplication of the distal third of chromosome 1 (breakpoints from q23-q44) showed no features in common with proband 1 other than developmental delay.

Proband 2 is still too young to ascertain the degree of severity of her language and motor delay. The general clinical impression of her progress indicates that she is more severely affected than proband 1, as would be expected from the result that her chromosome 22 breakpoint is located proximal to D22S45. One unusual feature seen in proband 2, vesicoureteral reflux, has also been seen in one previously described patient with del(22)(q13.3). The evidence for deletion of 22q13.3 being the cause, and not an unrelated, relatively common polymorphism, includes controls published by Flint et al. They studied by Southern analysis 186 nuclear families and 80 meioses from CEPH pedigrees for deletions, as well as 3000 unrelated subjects, for duplications of this region and identified no person aneusomic for the terminal region of chromosome 22q. Our internal laboratory controls include 97 patients for which we have used the DiGeorge chromosome region Oncor probe with FISH analysis to rule out microdeletions of the 22q11.2 region. No-one in this patient population (t/o DiGeorge syndrome, velocardiofacial syndrome, and isolated conotruncal defects) has been deleted for the distal marker cosmid D22S39, as was seen in our two probands.

We feel that investigation of the frequency of del(22)(q13.3) in the idiopathic mentally retarded population is warranted given the fact that two of the three subjects identified in the study by Flint et al had 22qter deletions and the recent identification of the two probands described here. The ability to use a commercially available probe (D22S39), which is already currently in use in a large number of cytogenetic laboratories, as an assay for patients with subtelomeric deletions of chromosome 22 will hopefully facilitate this investigation. A unique 22q telomere specific FISH probe has also been recently isolated and described by Ning et al.

In light of our experience with these two cases, when a very subtle deletion is suspected after G banding analysis, we suggest that currently available FISH probes could be useful in defining the abnormality. Similar to the cases described here, we have also recently identified a "cryptic" de novo 20p terminal deletion (suspected after G banding analysis) with a subtelomeric FISH probe in an idiopathic mentally retarded subject. A complete set of subtelomeric chromosome specific probes has recently been described. When such a set is available in a multicolour FISH format, screening of the idiopathic mental retardation population will hopefully aid in the rapid identification of a cause for developmental delay in a subset of these subjects. As shown by the family of proband 1, there is then potential for the identification of a cryptic familial terminal rearrangement in such cases. The recurrence risk for another affected child in such families is obviously greatly increased.

We would like to thank all the members of the Kennedy Krieger Institute Cytogenetics Laboratory for their technical expertise in the preparation of these cases, especially Shirley Perman and Susan Morse for the identification of the subtelar aberrant G banding patterns in probands 1 and 2. We would like to thank Callum Bell for generously providing us with YACs and Forrest Spencer for the use of reagents and equipment. In addition, we would like to thank the family of proband 1 for allowing the publication of clinical photographs. RFD and GHT are partially supported by NICHD Mental Retardation Research Center Core Grant HD32041. A permanent cell line of proband 1, L95-1, is available from the Kennedy Krieger Institute, Baltimore, MD, through Dr G H Thomas.