A supernumerary marker chromosome originating from two different regions of chromosome 18

Benno Röthlisberger, Krystyna Chrzanowska, Damina Balmer, Mariluce Riegel, Albert Schinzel

Abstract
By random amplification of a microdissected chromosome using the degenerate oligonucleotide primed polymerase chain reaction (DOP-PCR) and forward painting (microFISH), we characterised an extra structurally abnormal chromosome (ESAC) or supernumerary marker chromosome in a mentally retarded girl with a pattern of dysmorphic features. It could be clearly shown that the small marker chromosome originates from two different regions of chromosome 18, 18p11.1→18q11.1 and 18q12.3→18q21.1 respectively. Maternal origin of the de novo ESAC and biparental origin of the normal homologues of chromosome 18 were shown by PCR of several highly polymorphic microsatellites. In this case, application of microFISH was a prerequisite for rapid and precise characterisation of an ESAC. A definite identification of this discontinuous supernumerary marker chromosome would not have been possible using FISH with centromere specific probes or multicolour FISH approaches.

Keywords: supernumerary marker chromosome; microdissection; FISH; chromosome 18

At least 168 cases with an ESAC (excluding those from chromosome 15) originating from all chromosomes have been documented. Birth prevalence is estimated at 0.14 to 0.72 per 1000. Despite the relatively high frequency of trisomy 18 detected pre- and postnatally, only four cases with a marker chromosome derived from chromosome 18 have been reported. Subjects with a supernumerary marker chromosome have a duplication and in some cases a triplication of the material comprising the ESAC. The risk of an abnormal phenotype is/are the chromosome(s) involved identified, but in addition the exact segment(s) of the latter can be clarified. The importance of investigating large ESACs by microFISH has been shown by Blennow et al, who described a marker originating from three different chromosomes. However, the origin of most supernumerary marker chromosomes is determined by FISH analysis using a centromere specific probe or a chromosome specific library or both. This is generally considered to be sufficient if the marker chromosome is very small, since the common opinion is that the marker contains only centromeric and pericentromeric sequences. However, as we show in a case with an ESAC originating from chromosome 18, this assumption is inadequate, since even smaller markers may be the product of more than one chromosome region.

Case report
This 13 year old girl is the second child of healthy, unrelated parents. At birth the father was 25 years old and the mother was 24 years old. The girl was born at 42 weeks of gestation after an uneventful pregnancy. Birth weight was 3600 g (50th centile), length was 56 cm (>90th centile), and occipitofrontal head circumference (OFC) was 35 cm (between the 25th and 50th centile). Clinical examination at the age of 11½ years showed height 147 cm, weight 39.8 kg, and OFC 52 cm, all within the normal range. Total hand length (18.2 cm) was on the 10th centile and middle finger length (7.0 cm) was on the 3rd centile. The girl displayed the following anomalies: hypertelorism with inner canthal distance (3.7 cm) and outer canthal distance (9.8 cm) above the 97th centile, bilateral epicanthic folds, upward slanting palpebral fissures, short and upturned nose, long philtrum, short neck, widely spaced nipples, short 5th fingers, cubitus valgus on both sides, and three hyperpigmented areas over the abdomen (fig 1). Pubertal development was at Tanner stage 3.

The patient’s psychomotor development was moderately retarded with a global IQ of 76. She started to walk at 10 months of age and spoke her first single words at 3 years; she is hyperactive and suffers from dyslalia.

Methods
Metaphase chromosome preparations were obtained from PHA stimulated lymphocyte cultures from the patient and both parents according to standard procedures. GTG banded chromosome preparations were examined.
Chromosome microdissection was performed according to a slightly modified protocol by Senger et al. Five ESACs were transferred to a 2 µl collection drop, then amplified by DOP-PCR, and labelled with biotin. FISH analysis was performed according to Lichter et al. using the specific chromosome library generated by microFISH. The hybridised metaphases were examined with a Zeiss Axioplan epifluorescence microscope. Subsequent analyses were performed by image acquisition using a CCD camera and digital image analysis (QUIPS™, VYSIS).

To determine the origin and formation of the supernumerary chromosome, genomic DNA from the patient and both parents was amplified by standard PCR with commercially available highly polymorphic microsatellites (Research Genetics®), loaded onto a 6% polyacrylamide/urea gel, and visualised by silver staining.

Results

Chromosome preparations from blood lymphocyte cultures of the patient showed an additional small marker chromosome in 70% of the metaphases investigated (fig 2). Chromosome analysis of the mother and the father showed normal karyotypes. In order to determine the origin of the ESAC, two completely independent microFISH experiments with reverse painting on normal metaphases were performed. It could be seen that the ESAC was formed by two regions of chromosome 18, 18p11.1→18q11.1 and 18q12.3→18q21.1 (fig 3). The presence of chromosome 18 specific alphoid centromeric sequences was confirmed by FISH analysis with a biotinylated centromere specific probe (D18Z1). Following the detailed system of the ISCN the aberration can be written as: 46,XX[30%]/47,XX,+mar[70%.rev ish der (18)(p11.1→q11.1::q12.3→q21.1:) de novo. It should be mentioned that the orientation of the segment 18(q12.3→q21.1) is not clear.

For determination of the parental origin of the de novo ESAC and confirmation of the breakpoints, microsatellite analysis was performed on genomic DNA. Owing to the patient’s relatively high percentage of cells with the marker chromosome, the microsatellite analysis resulted in clear differences in the intensities of the maternal and paternal bands. All informative markers mapping to the duplicated regions were consistent with a maternal origin of the intenser band, whereas markers lying outside the duplicated regions resulted in similar intensities of the maternal and paternal bands and thus were in agreement with biparental inheritance of the normal homologues of chromosome 18 (table 1).

To exclude a paracentric inversion or an intrachromosomal insertion of the long arm of chromosome 18 in one of the parents, a reverse painting experiment on metaphases of both parents using the library created by microdissection of the ESAC was performed. The signals detected were identical to the signals detected on the control slide. Thus, a cytogenetically undetectable rearrangement in the relevant region of the long arm of chromosome 18 in one of the parents, namely the mother, could be excluded.

Discussion

While classical cytogenetic procedures (G, Q, C banding, silver staining) can provide information regarding the structure of a marker, it is limited in its power to identify the origin of marker chromosomes. Molecular cytogenetic techniques, namely FISH, have provided a valuable method for characterising the majority of ESACs. Different types of probes are available for this use. Since it is generally assumed that ESACs usually derive from a single autosome, originating from the centromeric region and eventually from adjacent pericentric regions, most of the ESACs have been characterised by FISH studies using centromere specific probes. Nevertheless, at least 10 cases have been published in which the chromosomal origin could not be determined using alphoid repeats. In two reports, more complex ESACs
Many additional marker chromosomes are very small ring chromosomes. According to the classical model, these ring chromosomes are considered to arise from chromosome breaks occurring on either side of the centromere and subsequent rejoicing of the broken ends into a circular structure, while the segments distal to the breakpoints are lost. To generate small ring chromosomes, at least one of these two breaks is either at the centromere or adjacent to the centromere in the region containing alphoid repeats. The two cases described above and, in addition, our case with a discontinuous marker chromosome originating from two different regions of chromosome 18 clearly show that this assumption of formation of small rings is simplified. It is possible that many of the small ESACs identified only by FISH with centromere specific probes or chromosome specific libraries are inaccurately characterised. A correct description of ESACs is important since there may be recognisable clinical syndromes owing to the particular genetic content of the marker chromosomes. ESACs may contain genetic material located pericentromeric on the short or long arm or both but also, as shown in our case, may contain interstitial regions of the corresponding chromosome or even contain genetic material originating from other chromosomes. Therefore, different distinct syndromes may be associated with each group of morphologically similar markers. The precise characterisation of ESACs with this technology could provide the patient and the genetic counsellor with better predictive information on the phenotype of the carrier. MicroFISH would be a rapid method of investigating all cases in which the chromosomal origin could not be determined using alphoid repeats, and it could in addition provide a more accurate characterisation of all ESACs that have usually been studied only by FISH using centromere specific probe or chromosome specific libraries or both, including multicolour FISH approaches. MicroFISH provides a technique not only to identify the origin of marker chromosomes but also to detect the euchromatin involved and thus provide information on the composition of ESACs.

A more detailed description of small supernumerary marker chromosomes could also result in a better understanding of their formation. In our case we could show that the marker originated from two different regions of the normal maternally inherited chromosome 18. After exclusion of a paracentric inversion or an intrachromosomal insertion of the long arm of chromosome 18 in the mother, at least four breaks are necessary to explain the formation of this ESAC, probably by the breakdown of a larger ring chromosome 18 until the stable form was generated. The breakpoint at 18q21 gives further evidence that this is a hot spot for breakpoints in complex chromosomal rearrangements.  

As surprising as these microFISH results may appear, the clinical findings are equally unexpected. Published cases with duplication of a similar chromosomal region display a more severe phenotype characterised by microcephaly, short stature, and more pronounced

<table>
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<th>Locus</th>
<th>cM</th>
<th>Alleles</th>
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<tr>
<td>D18S71</td>
<td>18p11.23p11.22</td>
<td>47</td>
<td>bc,bd,ac</td>
</tr>
<tr>
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<td>18p11.1q11.2</td>
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<td>D18S769</td>
<td>18q11.1q11.2</td>
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<td>D18S1143</td>
<td>18q21.1q21.1</td>
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The alleles are given in the order patient, mother, father. The markers showing intense maternal alleles in the patient are given in italics and the intense alleles themselves in bold. Allele designations (a to d) are arbitrary. Genetic mapping according to the sex averaged linkage map by the Genetic Location Database (LDB).
developmental delay. In contrast, our patient had normal measurements at birth as well as later in life and a global IQ of 76. Minor dysmorphic signs such as epicanthic folds, upward slanting palpebral fissures, widely spaced nipples, and short 5th fingers are present in our case as well as in most of the formerly reported cases. The less severe phenotype in our case could be explained by mosaicism and by the smaller size of the duplicated region.

In conclusion, since the great variability of phenotypes in patients with ESACs of the same chromosomal origin is probably the result of variations in their genetic content (in addition to the degree of mosaicism and the possibility of uniparental disomy), microFISH is a prerequisite for an accurate characterisation of small supernumerary marker chromosomes, as shown in our case with a discontinuous ESAC originating from two different regions of chromosome 18. Such a detailed characterisation and precise description of the genetic content is not possible with other molecular cytogenetic methods, namely FISH with centromere specific probes or chromosome specific libraries, including multicolour FISH approaches. The more accurate characterisation of ESACs may result in the description of different distinct syndromes associated with each group of morphologically similar but genetically different markers.

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