Is There a Yet Unreported Unbalanced Chromosomal Abnormality without Phenotypic Consequences in Proximal 4p?

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The finding of unbalanced chromosomal abnormalities (UBCA) was recently reviewed and summarized from a total of 200 families [Barber, 2005; http://www.ngrl.org.uk/Wessex/collection/ubca_chart.htm]. UBCA usually involve several megabases of DNA, and carriers of such UBCA are ascertained either through an abnormal phenotype, adverse reproductive effects or by chance [Liehr et al., 2009].

Here, we report a possibly new chromosomal region involved in cytogenetic UBCA in a case detected through prenatal diagnosis. The indication for amniocentesis was nuchal translucency of 3.9 mm in week 14 + 3, with a nasal bone of 1.5 mm in week 13. Cytogenetic analysis revealed a karyotype mos 47,XX,+mar\textsuperscript{[29]}/46,XX\textsuperscript{[12]} in 3 independent amniotic fluid cell cultures. The origin of the small supernumerary marker chromosome (sSMC; for sSMC see also http://www.med.uni-jena.de/fish/sSMC/00START.htm) was determined to be from chromosome 4 by multiplex fluorescence in situ hybridization (M-FISH) [Speicher et al., 1996; see fig. 1 A]. By subcentromere-specific M-FISH (subcenM-FISH) [Starke et al., 2003], it was shown that only centromere-near short-arm material was present on the sSMC (fig. 1 B), and array-proven multicolor banding (aMCB) [Weise et al., 2008] suggested that the sSMC was derived from 4p14\textasciitilde;p13 to 4q11.1 (fig. 1C). Array comparative genomic hybridization...
tion (array-CGH) done according to the manufacturer’s instructions using whole genomic DNA confirmed and refined these results (fig. 1D). The sSMC lead according to Agilent platform using a 44K chip to gain of copy number of 39,799,760 to 48,924,359 bp. Thus, the final karyotype was mos 47,XX,+min(4)(:p14-1q11.1:)[29]/46,XX[12]; the karyotype is given according to Liehr [2009], describing a centric minute shaped sSMC. Marker analysis ruled out uniparental disomy 4. After genetic counseling, the parents opted for the pregnancy.

At birth, placental cells and fibroblast cultures from the umbilical cord were analyzed. Interphase FISH using a centromere-specific probe for chromosome 4 revealed 3 specific signals in 60% of the placental nuclei and chromosome analysis confirmed mosaicism in fibroblasts as well. Parents refused cytogenetic analysis of peripheral blood. The parental chromosomes were normal.

A healthy girl was born at week 39 with a weight of 3,740 g. The baby was phenotypically normal and developed normally during the time of monitoring, i.e. until the age of 1 year.

Here we report the first case with an sSMC derived from 4p14 to 4q11.1 present in the mosaic of 60–72% of the studied cells, leading to only minor clinical symp-
toms with respect to the size of the observed imbalance. At the age of 3 years 6 months, the patient has no external malformations and size, weight and head circumference are all at the 50th percentile. She showed slight psychomotor retardation, i.e. sitting at 8 months, walking at 1 year of age, delayed language development, and now speaking 4–5-word sentences. The size of the duplicated region is 8.9 Mb, which is a common size for UBCA (http://www.med.uni-jena.de/fish/sSMC/00START.htm). The region is known to be comprised of many copy number variations and other repetitive elements [Jackson et al., 1999; Iafrate et al., 2004], but also contains more than 40 genes of which 22 have ‘OMIM numbers’ and at least 2 of which are associated with late-onset neurological conditions. Maybe it has also to be considered that the mosaicism present in the reported case could also be responsible for the lack of symptoms, as it has recently been shown that different body tissues can vary significantly in mosaic-expression of sSMC carriers [Fickelscher et al., 2007].

Overall, apart from the case reported here, there is no comparable case in the literature with such a small imbalance in 4p; there is only a larger one which includes 4p16 to 4p12 reported by Kakinuma et al. [2008]. Thus, we have to wait for further reports on duplications to clarify the question if proximal 4q may harbor a UBCA.

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