Supernumerary marker chromosome and global developmental delay: Role of microarray - case report and review of literature

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Received - 14 August 2019 Initial Review - 17 September 2019 Accepted - 24 September 2019

ABSTRACT

Small supernumerary marker chromosomes (sSMCs) are defined as structurally abnormal chromosomes that cannot be identified or characterized by conventional karyotype analysis and are generally equal in size or smaller than chromosome 20. Here, we present the molecular characterization of an sSMCs derived from chromosome 15 in prenatal diagnosis in a 38-years-old female.

Keywords: Array CGH, Global Developmental delay, Karyotype, Marker chromosome.

Small supernumerary marker chromosomes (sSMCs) are defined as structurally abnormal chromosomes that cannot be identified or characterized by conventional karyotype analysis and are generally equal in size or smaller than chromosome 20 [1]. sSMCs are present in nearly 0.044% of live births and 0.075% of prenatal cases [2,3]. Approximately 77% of sSMCs arise de novo and 23% are inherited [2,3]. Nearly 70% of sSMCs are derived from the short arms and pericentromeric regions of acrocentric chromosomes [4].

Most (70%) of de novo sSMCs have no phenotypic effects [5]. However, in at least 30-50% of prenatally detected sSMCs cases, the pregnancy is terminated [2,3] which means unnecessary abortions were induced in a certain percentage of potentially healthy children with sSMCs. Therefore the precise characterization of marker chromosomes is crucial for prenatal diagnosis and proper genetic counseling. Here, we present the molecular characterization of an sSMCs derived from chromosome 15 in prenatal diagnosis.

CASE REPORT

A 38-years-old female, G3P1A1L1 presented at 17 weeks of gestation with a history of her girl child aged 6 years having Global developmental delay. Her period of gestation was corresponding to the Ultrasonography. She had undergone combined first-trimester screening by which was reported to be low risk. There was no gross congenital malformation in the fetus of present pregnancy.

She had delivered a girl child six years back at term by an Emergency Cesarean section for non-progress of labor. The child had delayed milestones and mild facial dysmorphism for which she was evaluated and diagnosed to have Global developmental delay. The genetic evaluation was done including Comparative Genomic Hybridization (CGH) array and Fluorescent in situ hybridization (FISH) which diagnosed the presence of a marker chromosome in the child, 47, XX + mar. On parental evaluation, the paternal karyotype was normal 46 XY and mother was detected to have the same marker chromosome. However, the mother was phenotypically and functionally normal and part of the medical profession. A 2.05 Mb duplication was detected on the long arm of chromosome 15 (cytogenetic location 15q11.1q11.2).

Since both the mother and the child had the identical marker chromosome and mother being normal, the Global development delay could not be completely attributed to the marker chromosome. The microarray report of both proband and the mother was awaited. Meanwhile, the parents opted for prenatal testing by amniocentesis and full Karyotype and FISH studies in the fetus of present pregnancy. She underwent an uneventful procedure and was discharged the same day. FISH was performed on the cells from the amniotic fluid sample using Vlysis DNA probe LSI SNRPN/LSIPML/CEP (tricolor). Signal enumeration

Figure 1: Amniotic fluid Karyotype of fetus.
was done in 50 cells. The final report of the amniotic fluid karyotype was the same as the mother and the index case (Fig. 1).

Post-counselling, the couple refused further analysis and opted for termination. The abortus underwent an autopsy wherein no structural abnormality was detected. The final microarray reports of the proband fetus and the mother were reported as follows: Fetus: Genomic duplication of 2.6 Mb on cytoband 15q11.1q11.2 starting from 20,055,137bp to 22,698,581bp on the long arm of Chromosome #15. This has been reported benign on ISCA Database (Fig. 2). Mother: Genomic duplication of 2.6Mb on cytoband 15q11.1q11.2 starting from 20,095,481 to 22,698,581bp on long arm q of chromosome#15. This has been reported benign on ISCA Database (Fig. 3). Loss of heterozygosity (LOH) of 19.2Mb was seen on Chromosome #16. It may be due to parental consanguinity which was absent in this case and can lead to uniparental disomy. Proband: Genomic duplication of 2.6Mb on cytoband 15q11.1q11.2 starting from 20,055,137 bp to 22,698,581 bp on the long arm of Chromosome#15. This has been reported benign by ISCA Database. Genomic duplication of 177 Kb on cytoband 9p 24.3 starting from 204,193 bp to 381,489 was present on the short arm p of chromosome # 9. Reported uncertain likely benign by ISCA (Fig. 4).

DISCUSSION

It is estimated that 50% of all sSMCs are derived from chromosome 15 [6]. Children with an sSMC are individually very different from each other, even when the sSMCs causes a known syndrome. Wong et al reported a de novo marker chromosome part of 15 detected on amniocentesis in a 21-year-old woman with high risk for Trisomy 21 on combined screen. Parental Karyotype was normal and the fetus did not have any anomaly or dysmorphism on sonography. Pregnancy outcome was successful with the baby having karyotype 47XY+Marker. At one year of age, the baby was progressing normally [7]. Jon Soo Kim et al reported a 10-month-old female infant referred to their pediatric neurology clinic for uncontrolled seizures and global developmental delay. Array CGH showed amplification from 15q11.1 to 15q13.1 spanning 8.47 Mb [6]. The reported clinical features of patients with 15q13.3 microdeletion and duplication are rather heterogeneous, ranging from mental retardation and psychiatric illness to seizures. In the present case, the proband and the mother had the same marker chromosome on routine karyotype and FISH; however, the mother was phenotypically and functionally normal.

Approximately 70% of SMCs are de novo and 30% are inherited. The most common SMCs are derived from acrocentric chromosomes and have a satellited or bisatellited structure. Chromosome 15 accounts for the highest percentage (~50%) of this group. Melo et al [8] reported a case of a 39-year-old woman who underwent amniocentesis at 16 weeks of gestation for advanced maternal age. An abnormal fetal karyotype – 47, XX,+mar – with one sSMCs was detected in all metaphases. Cytogenetic and molecular analyses revealed a fetal karyotype 47, XX,+mar patishidic (15)(q12)(D15Z1++,SNRPN–), in which the sSMCs (15) was a paternally inherited inverted duplicated chromosome and did not contain the critical region of Prader–Willi/Angelman syndromes. Uniparental disomy was excluded and genetic sonogram was normal, pregnancy outcome was a normal baby.

This case supports the literature in two aspects: sSMCs (15) that do not contain PWACR generally have a normal phenotype, and sSMCs transmitted by normal carriers to their offspring are not commonly correlated with clinical problems [8]. Some sSMCs lead to specific syndromes. Entire and partial gene deletions/ duplications can produce a completely different phenotypic effect. The low copy repeats (LCRs) in chromosome 15q11q13 have been recognized as Breakpoints (BP) for not only intra chromosomal deletions and duplications but also small supernumery marker chromosomes 15, sSMCs (15) [9].

Most sSMCs (15) take the form of a dicentric inv dup and can be classified into two groups: small sSMCs (15) and large sSMCs (15). The small sSMCs (15) are metacentric chromosomes without euchromatic material and do not contain the Prader-Willi/Angelman Critic Region (PWACR), which usually clinically irrelevant. In contrast, the large sSMCs (15) are acrocentric chromosomes containing copies of PWACR and are frequently associated with abnormal phenotype (mELO). For the other sSMCs, it is still not possible to predict outcomes, although

Figure 2: Amniotic fluid Microarray of Fetus.

Figure 3: Microarray of mother.
new cases can now be compared with the cases collected on the sSMCs website [9,10].

If a child has problems with growth or development, it is likely that the sSMCs has affected them but the cause could also be something different, not the sSMCs. The correct characterization of gene deletions and duplications is a crucial point in order to identify the genotype phenotype correlation. Some conclusions can be drawn by comparing other cases from the medical literature. Conventional karyotype analysis can detect numerical and structural chromosomal abnormalities but cannot determine the origin and genetic content of sSMCs. Array CGH has the ability to detect DNA dosage imbalance including deletions and duplications in the euchromatic regions and is useful for the characterization of the origin and hereditary effects in the sSMCs [11].

Sun et al [12] reported the analysis of twenty cases with sSMCs initially were detected by G-binding karyotype. Routine cytogenetic analysis showed mosaic marker chromosome in six out of the 20 cases. All the 20 cases were subjected to a CGH assay, and seventeen of them were successfully identified for the chromosome origin. Array CGH has many advantages that make it extremely useful for characterizing sSMCs. It clearly and easily determines the components of sSMCs in a single assay. This advantage was especially useful for complex sSMCs and multiple sSMCs. Complex sSMCs is a subgroup of sSMCs which consist of chromosomal materials derived from more than one chromosome [12]. Vetro et al [13] reported 4 cases of sSMCs that after array CGH was interpreted rather differently reporting two types of complex markers which DNA content was overlooked by conventional approaches. All are likely derived from partial trisomy rescue events thus changing the present ideas on composition of conventional approaches. All are likely derived from partial trisomy types of complex markers which DNA content was overlooked by more than one chromosome [12].

Glessner et al in the first meta-analysis of copy number variations (CNVs) studied five cohorts including Autism spectrum disorders (ASD) and Attention deficit hyperactivity disorders (ADHD). They stated that cumulative evidence indicates a shared genetic etiology of neurodevelopmental and neuropsychiatric diseases. They identified the DOCK8/KANK1 locus as containing exonic CNVs with genome-wide significant meta P values and consistent direction of effects across all five cohorts. There was a significant association of genomic duplication in Chromosome #9 short arm with ADS, ADHD and Neuropsychiatric illnesses [14]. In the present case, though the parents opted for termination based on the FISH report, the array CGH would have given the clarity towards whether the fetus would have been actually affected as its microarray did not show the genomic duplication in chromosome #9 present in the proband. The marker chromosome was only coincidental to the developmental delay.

Presence of uniparental disomy as a cause for sSMCs is being investigated. In a review article Kotzot stated that the incidence of UPD in cases with sSMCs is increased by coincidence, ascertainment bias and because for most chromosomes only then is the fetus viable [15]. In this case study, we validate that array CGH analysis provides an alternative to telomere FISH and disease-specific FISH in the cytogenetic diagnostic laboratory.

**CONCLUSION**

We conclude that array CGH is a modern and precise diagnostic tool that will complement and enhance current methods of detecting chromosomal imbalances prenatally. In a combination of a detailed ultrasound examination and karyotype analysis, it can provide more precise and rapid prenatal diagnosis of sSMCs. It is suggested that UPD is always tested for when an sSMCs is detected. In order to establish a stronger base for clinical service in the future and avoid miss characterization, more sSMCs cases need to be detailed characterized. This will help to clarify the variable clinical characteristics of sSMCs and provide additional information to aid clinical service and future research.

**REFERENCES**