A placental trisomy 2 detected by NIPT evolved in a fetal small Supernumerary Marker Chromosome (sSMC).

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Case Report

Keywords: array comparative genomic hybridization, fluorescence in situ hybridization, karyotyping, mosaicism, noninvasive prenatal test, small supernumerary marker chromosome
Abstract

**Background:** Non-invasive prenatal testing (NIPT) is a rapidly developing and widely used method in the prenatal screening. Recently, the widespread use of the NIPT caused a neglecting of the limitations of this technology.

**Case presentation:** The 38-year-old woman underwent amniocentesis because of a high risk of trisomy 2 revealed by the genome-wide Non-Invasive Prenatal Test (NIPT). The invasive prenatal diagnosis revealed the mosaicism for a small supernumerary marker chromosome sSMC derived from chromosome 2. Interphase fluorescence in situ hybridization (FISH) on uncultured amniocytes revealed three signals of centromere 2 in 30% of the cells. GTG-banded metaphases revealed abnormal karyotype (47,XX,+mar[21]/46,XX[19]) and was confirmed by array comparative genomic hybridization (aCGH). Cytogenetic analyses (FISH, aCGH, karyotype) on fetal skin biopsies were performed and confirmed the genomic gain of the centromeric region of chromosome 2. In the placenta, three cell lines were detected: a normal cell line, a cell line with trisomy 2 and a third one with only the sSMC.

**Conclusion:** Whole-genome Non-Invasive Prenatal Testing allows not only the identification of common fetal trisomies but also diagnosis of rare chromosomal abnormalities. Especially in such cases, it is extremely important to perform not only NIPT verification on a sample of material other than trophoblast, but also to apply appropriate research methods. Such conduct allows detailed analysis of the detected aberration, thus appropriate clinical validity.

**Background**

Non-invasive prenatal testing (NIPT) is a rapidly developing and widely used method in the world. The NIPT resolution depends on the technique applied. Recent research shows that different NIPT approaches can be used not only to detect the most common trisomies, but also for detection of rare aneuploidies and to assess submicroscopic deletions and duplications, as well as identify monogenic diseases [1,2,3,4,5,6]. The well-documented specificity and sensitivity of NIPT for the most common trisomies (13, 18, 21) are different for each chromosome. Mackie et al. estimated specificity of 99% to 100% for 13, 21, 18 trisomies and monosomy X. The sensitivity to detect trisomy 21 reaches 99% but for trisomy 13, 18 and monosomy X it is 90%, 98% and 93% respectively [7]. Similarly study Lee et. al showed 99-100% specificity and sensitivity for trisomies 21, 13 but sensitivity for 18 is high as 92% [8]. However, in literature there is no data of NIPT sensitivity and specificity for other chromosomes.

Moreover discordant results between the NIPT and the fetal karyotype have been reported. This can be due to the fact that the fetal cell-free DNA (cfDNA) in the maternal blood is derived from the cytotrophoblastic cells of the placenta [9] and confined placental mosaicism (CPM) can yield a discrepancy between NIPT and amniotic fluid analysis. CPM are observed in about 1 – 2% of chorionic villus samplings (CVS) [10]. Additionally, Twin pregnancies, vanishing twing, maternal copy number variations (CNV), maternal mosaicism, as well as maternal malignancies but also low-level imbalance-
mosaicism examined chromosomes are the most commonly reported reasons for discordance [11,12,13,14,15,16,17,18,19].

Therefore, in the case of fetal abnormalities identified during the ultrasound screening, invasive test instead of NIPT is recommended [20]. Here we report a discrepancy between the result of NIPT and the final genetic diagnosis. The indication for the prenatal examination in this case was a high risk of trisomy 2 revealed by NIPT test while in invasive diagnostic mosaicism of a small supernumerary marker chromosome (sSMC) derived from chromosome 2 has been detected.

Case Presentation And Methods

A 38-year old woman came to the clinic because of increased risk of trisomy 21 (T21=1/291) calculated based on the combined test. The ultrasound test did not reveal any abnormalities. The patient conducted NIPT at the 14\textsuperscript{th} week of gestation, which revealed a high risk of trisomy 2. Amniocentesis was performed at 17\textsuperscript{th} weeks of gestation. The amniotic fluid was aliquoted in order to perform FISH, aCGH and conventional karyotyping analysis. A fetal skin and placental were taken to perform FISH and aCGH on uncultured cells, and conventional karyotyping on cultured cells.

Methods

**Fluorescence in situ hybridization (FISH)**

Interphase FISH analysis was performed on uncultured amniotic fluid cells using commercially available DNA probe for region 2p11.1-q11.1 (Spectrum Red centromere of chromosome 2 probe, locus D2Z2) (Chromosome 2 Alpha Satellite Probe – Cytocell) and probe for region Xp11.1-q11.1 (Spectrum Green centromere of chromosome X probe, locus DXZ1) (Chromosome X Alpha Satellite Probe – Cytocell) as a control. Fetal skin and placenta cells were examined with the same probe for chromosome 2 (Chromosome 2 Alpha Satellite Probe – Cytocell) and locus specific probe RP11-676E9 from bacterial artificial chromosome (BAC) clone located at cytoband 2p21 (chr2:45,579,001-45,773,026; GRCh37). The position of the BAC clone was taken from the UCSC Genome Browser. DNA was labelled with Spectrum Green-dUTP (Vysis), using the Vysis nick-translation kit (Vysis). Labelling of probe was done as described previously by Merscher et. al [21].

**Array comparative genomic hybridization (aCGH)**

DNA was extracted directly from amniotic fluid cells and from fetal skin cells. Microarray was performed using CytoSure Constitutional v3 (8x60k), (Oxford Gene Technology, GRCh37/hg19) according to procedures described by Bartnik et al., [22]. Scanned images were quantified using Agilent Feature Extraction software (v10.0). The CytoSure (Oxford Gene Technology) software was used for chromosomal microarray analysis. All genomic coordinates are based on the March 2006 assembly of the reference genome (NCBI37/hg19).
Karyotype analysis

Analysis of GTG-banded metaphases at approximately 400-band resolution in amniocytes and skin fibroblasts after in situ culturing was performed according to the standard protocol.

Result

Interphase FISH analysis performed on amniotic uid showed three signals for the centromere of chromosome 2 in 45 cells and two signals in 105 cells, indicating 30% mosaicism for trisomy 2 in female fetus. However, based on the interphase FISH, structural chromosomal abnormalities cannot be identified. Analysis of GTG-banded metaphases revealed a 47,XX,+mar[21]/46,XX[19] karyotype indicating the presence of a marker chromosome in 52% of the analyzed metaphases. In order to determine the genetic content of the sSMC, aCGH was performed. Array CGH indicated a pericentromeric gain of 14.83 Mb in chromosomal region 2q11.1q13 (95420515_1102553160). Based on the morphology of the marker detected in GTG-banding (figure 2) and molecular results, we concluded that the marker was a small supernumerary ring chromosome (sSRC) derived from chromosome 2. Based on Liehr online database [23] (the panel B in figure 1), the region is known as pathogenic, causing dysmorphism, developmental delay, brain malformations, heart defect, hypotonia, mental retardation, finger or toe/foot malformations, growth retardation, kidney problems/ malformations and omphalocele or situs inversus. Following detailed genetic counselling, the parents decided to terminate the pregnancy. Ultrasound examination and postnatal autopsy at the 22nd weeks of gestation revealed a female fetus with growth restriction but without any further malformations. Fetal skin an placental tissue were sampled to perform further cytogenetic examinations. Array CGH of skin cells (red plot in figure 1) showed the same 14.83 Mb gain of chromosome 2q11.1q13 (95420515_1102553160). However, this time the plot showed higher level of mean log ratio (0.44 on average), compared to results from the amniotic fluid cells (0.19 on average), indicating higher level of mosaicism in skin cells. FISH analysis of fetal skin cells (figure 3B) showed three signals of centromere 2 in 84 of 132 analyzed nuclei, indicating 64% of mosaicism for the sSMC. The placenta (figure 3A) showed the presence of three cell lines: 39% of scored nuclei had a normal number of signals from chromosome 2, 40% presented trisomy of the whole chromosome 2 and 21% carried the sSMC. Finally, cytogenetic analysis by conventional karyotype of cultured skin and placental cells revealed 30% and 33% mosaicism of the sSMC respectively (table 1), however, no trisomy of whole chromosome 2 was detected.

Table 1. Summarized results from FISH and GTG-banding scoring.
| Biological material | Treatment | Method of analyses | Results | Mosaicism [%] |
|---------------------|-----------|--------------------|---------|---------------|
| Amniotic fluid      | uncultured | FISH               | nuc ish(D2Z2x3)[45/150] | 30 |
|                     | cultured  | GTG                | mos 47,XX,+mar[21]/46,XX[19] | 52 |
| Fetal skin          | uncultured | FISH               | nuc ish(D2Z2x3)[84/132] | 64 |
|                     | cultured  | GTG                | 47,XX,+mar[15]/46,XX[35] | 30 |
| Placenta            | uncultured | FISH               | nuc ish(D2Z2x3,RP11-676E9)x3[45/112]/(D2Z2x3,RP11-676E9x2)[44/112] | 40/39 |
|                     | cultured  | GTG                | 47,XX,+mar[6]/46,XX[12] | 33 |

Abbreviations: GTG - Giemsa trypsin G-banding, FISH – fluorescence in situ hybridization, Nuc ish – nuclear in situ hybridization, Mos – mosaic, Mar – marker chromosome

### Discussion And Conclusions

Herein, we report the prenatal diagnosis and molecular cytogenetic characterization of the mosaicism for sSMC derived from chromosome 2. The indication for the prenatal examination in this case was a high risk of trisomy 2 revealed by NIPT test. Our case underlines the screening nature of the NIPT test and also demonstrates that performing only the FISH analyses in uncultured amniocytes, as the most rapid way to verify the NIPT result, would be insufficient for the correct diagnosis. Therefore, bearing in mind that the fetal and placental karyotypes might be different, a good follow-up of abnormal NIPT results is necessary, but as the investigation showed, techniques as well as the tested tissue should be properly selected.

Chromosomal mosaicism can be associated with a wide spectrum of phenotypes extending from apparently normal to severe or lethal. Hsu and coauthors [24] summarized the outcome of 11 fetuses with whole chromosome trisomy 2 detected in mosaicism, which ranged from 4% to 33% and has been confirmed in other tissues. Within these cases, 1 with the lowest percentage of abnormal cells (4% in the amniocytes but without trisomy in blood and placenta) resulted in an apparently normal livebirth; 1 newborn presented IUGR, 1 had IUGR and multiple anomalies, 3 stillbirths or intrauterine deaths, and 4 in elective terminations (11-33% mosaic cells, all with abnormal findings).

Clinical consequences in cases of mosaicism of partial trisomy can be harder to predict. Marker chromosomes derivative from chromosome 2, described in the literature, have been associated with multiple clinical consequences. Children diagnosed after birth might present: developmental delay, brain malformations, seizures, heart defect, kidney malformations, hypotonia, mental retardation, growth
retardation, microcephaly [23]. However, little is known about prenatal ultrasound abnormalities in such cases.

Microarray analysis defined the gain of 14.83 Mb in the 2q11.1q13 region. Our case is highly similar to a case reported by Riegel and Schinzel [25]. Authors described a 4-year-old boy with multiple clinical features and a duplicated segment of 2q11.1-q13.2 presented in all analyzed cells. Low birth weight (<10th centile), a left cleft lip with cleft palate and natal left upper incisor were noted at birth. Consecutive examination revealed mental retardation, low set ears, irregular teeth, cryptorchidism and epilepsy. However, also in this case, three ultrasound routine examinations performed prenatally did not disclose abnormal findings. The difference in the clinical presentation between our case and the patient described by Riegel and Schinzel [25], might be due to the percentage of the level of normal cells, but also could be because of different impact of the simple region duplication in the genome versus marker chromosome.

Furthermore, cytogenetic discrepancy between results of uncultured fetal cells (skin 64%, amniocytes 30%) and cultured cells (skin 30%, amniocytes 52%) provides an additional challenge in genetic counselling. FISH analyses in uncultured skin cells presented higher percentage of mosaicism (64%) than was found later in cultured skin cells (30%). The discrepancy has also been observed between uncultured and cultured placenta cells. In uncultured placenta we observed three cell lines; with a normal number of signals from chromosome 2 (39%), trisomy of whole chromosome 2 (40%), and 21% of cells carrying the ssSMC. Whereas cultured placenta did not reveal the whole trisomy 2. The variations in cell numbers carrying an abnormality before and after cultivation are compatible with results of other authors and presumably stem from a selection against the trisomic cells after long-term culture [26]. Different values of mosaicism indicated in uncultured (30%) and cultured (52%) amniocytes can be caused by the contamination of amniotic fluid with maternal cells. Moreover, the number of available for analysis metaphases was relatively low (n=40) in comparison to uncultured nuclei analyzed with FISH (n=150). The fact that the abnormal cells may divide slower and undergo apoptosis more easily causes an increase in the proportion of normal cells, should be taken into consideration during genetic consultation [27].

In the presented case, based on FISH and GTG-banding results, the ssSMC was in fact a small ring chromosome. There is a high probability that this ring chromosome formation is caused by low repetitive elements present in the pericentric region of chromosome 2 [28,29].

We have observed a discrepancy between the indication for invasive testing, and the final genetic diagnosis results. However, the presence of the trisomy of whole chromosome 2 was confirmed in the placenta. It is probable that the marker chromosome has arisen from trisomic embryo cells, which existed at an initial stage of development. The presence of the marker chromosome mosaicism in subsequent studies points to the existence of a functional trisomy rescue mechanism in this case.

A genetic counselling of fetuses with mosaicism is especially problematic because of the relatively poor phenotypic data and time-limitation. Considering the fate of the pregnancy in the case of detecting abnormalities, NIPT can be assessed as a screening test, and should be accompanied by an ultrasound
examination. Hence, invasive diagnosis is necessary to confirm the non-invasive results. In the presented case, it should be emphasized that there is a discrepancy between the result of the NIPT study and the results of the genetic diagnostic tests performed in the fetal tissues. Therefore it is crucial to choose the most suitable investigation strategy in order to perform the most rapid genetic diagnosis.

**Abbreviations**

NIPT - non-invasive prenatal testing

FISH - fluorescence in situ hybridization

aCGH - array comparative genomic hybridization

sSMC - small supernumerary marker chromosome

cfDNA - cell-free DNA

IUGR - intrauterine growth restriction

sSRC - small supernumerary ring chromosome

GTG - Giemsa trypsin G-banding

Nuc ish – nuclear in situ hybridization

Mos – mosaic

Mar – marker chromosome

Mb - mega base pairs

CVS - chorionic villus samplings

CPM - confined placental mosaicism

CNV - copy number variations

**Declarations**

**Ethics approval and consent to participate**

This work was approved by Committee on Ethics in Institute of Mother and Child, Warsaw, Poland.

**Consent for publication**

Consent to participate in this study was signed by the gravid.
Availability of data and materials

All data generated or analysed during this study are included in this published article.

Competing interests

The authors declare that they have no competing interests.

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Authors’ contributions

JD and MD, NB, JRV contributed in writing the manuscript, NB, JRV, KvdB performed NIPT, AL, KJ-D, MK, JD and MD performed karyotypes and FISH studies, MB performed array CGH, AK-CH, TR performed ultrasound scan. EO conducted genetic consultations. BN was a major contributor in writing the manuscript. All authors read and approved the final manuscript.

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