Rapid prenatal diagnosis of Facioscapulohumeral Muscular Dystrophy 1 by combined Bionano optical mapping and karyomapping

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Abstract
Purpose: To explore the feasibility of performing rapid prenatal diagnoses of FSHD1 using a combination of Bionano optical mapping and linkage-based karyomapping.
Methods: DNA specimens from a family that had been previously diagnosed with FSHD1 using Southern Blot analysis were used for this study. Genetic diagnosis of the proband, fetus chorionic amniotic fluid, and aborted fetal tissue was performed using Bionano optical mapping (BOM) together with linkage-based karyomapping.
Results: BOM analysis showed that the proband's 4q35.2 region contained four D4Z4 repeats and the 4qA permissible allele, consistent with the previous FSHD1 diagnosis obtained by Southern Blotting. BOM analysis of the fetus' 4q35.2 region was consistent with that of the proband. Karyomap analysis revealed that the fetus inherited the affected chromosome segment from the proband. After genetic counseling, the couple choose termination of pregnancy, and we performed gene diagnosis of the abortus tissue by BOM.
Conclusions: Bionano optical mapping can determine the number of D4Z4 repeats and exclude interference of the 10q26.3 homologous region, and in combination with karyomapping, can be used for rapid and accurate prenatal diagnosis of FSHD1.

1 INTRODUCTION
Facioscapulohumeral Muscular Dystrophy (FSHD; MIM 158900; MIM 158901) is a common form of inherited human myopathy, with an estimated prevalence of 1:20,0001, presenting typically before age 20, with progressive weakness of the muscles of the face, shoulders and upper arms 2. FSHD is inherited as an autosomal dominant disease, due to a genetic deletion in chromosomal region 4q35. It can be divided into two types: FSHD1 and FSHD2 according to the underlying genetics. The current consensus is that the causative genetic defect in FSHD is the loss of transcriptional repression of the Double Homeobox Protein 4 (DUX4) gene present in each repeat of the macrosatellite array (D4Z4) at chromosome 4q35 3.

FSHD1 is more frequent (~ 95% of cases). The majority of FSHD1 patients (70-90%) inherit the deletion from one parent (50% risk of inheriting), a significant number (10-30%) of cases are due to de novo mutations. FSHD1 is characterized by: 1) variable length deletions of 3.3 Kb (D4Z4) repeated elements at chromosomal region 4q35.
Normally people have 11-100 repeats; but most FSHD patients have 1-10 repeats; 2) disease-linked 4qA allele in 4q35 subtelomere; 3) hypomethylation of D4Z4 repeats. FSHD2 (about 5% of cases) is usually caused by mutations at the SMCHD1 (Structural Maintenance of Chromosomes Hinge Domain 1 gene, on chromosome 18p11.32) or DNMT3B gene which also leads to the hypomethylation of D4Z4 repeats. Both of FSHD1 and FSHD2 cause misexpression of the double homeobox 4 (DUX4) and subsequently the toxic protein (Figure 1).

FSHD is not fatal disease, but there is no effective treatment. And it presents with clinical heterogeneity and continual aggravation, so that patients and their families bear a huge psychological burden. Most families wish to perform prenatal diagnosis. The traditional genetic method of FSHD diagnosis is by pulse field gel electrophoresis (PFGE) and Southern Blot which is process-complicated, labor- and time-intensive, needs a large quantity of high quality DNA, and can't accurately recognize critical length. These deficiencies are tolerable in gene diagnosis but intolerable in prenatal diagnosis, so a simpler and more accurate method would be advantageous. In a recent approach shown by Dai Y et al, the commercial Bionano Saphyr single DNA molecule optical mapping system is well suited for the rapid diagnosis of FSHD1. This method produces ordered genome-wide restriction maps by optical inspection and sizing of restriction fragments from single linearized DNA molecules.

In this study, we explored the new method of Bionano optical mapping combined with Karyomapping as new strategy for prenatal diagnosis.

2 MATERIALS AND METHODS

The affected family (Figure 2A) consisted of the proband, a 28 years old male with typical clinical symptoms including: typical facial weakness, scapular winging and proximal weakness (Figure 2B). He sleeps with eyes slightly open, and could not puff out his cheeks or whistle. His mother and sister all had similar muscle symptoms. All had obtained previous genetic diagnoses of FSHD1 by Southern Blot, which identified a 21 kb 4qA-type EcoRI fragment on chromosome 4 as being pathogenic. We obtained samples for prenatal diagnosis from this family: blood samples from the proband, his wife and his mother, fetal amniotic fluid at 17 weeks of pregnancy, amniotic fluid cells after culture and abortus tissue after the abortion. Informed consent was obtained from all family members. This study was approved by the Ethics Committee for Medical Research of the First Affiliated Hospital of Zhengzhou University.

2.1 Bionano Optical Mapping

2.1.1 Sample DNA preparation

Peripheral blood and culture cells DNA were isolated and extracted according to the Bionano Prep Blood DNA Isolation Protocol (Bionano Genomics #30033). Clinical centrifugation for 3 min at 2000 g was used to separate the white blood cells (WBC) from 3 ml of fresh blood, and these were then resuspended in Cell Suspension Buffer (CSB). High molecular weight DNA of aborted fetal tissue was extracted using the Bionano Prep Animal Tissue DNA Isolation Fibrous Tissue Protocol, (Bionano Genomics #30071). 50 mg tissue was ground up with liquid nitrogen in 2.5 ml of Animal Tissue Homogenize Buffer (HB) and 2.5 ml of anhydrous ethanol (Bionano Genomics). The mixture was left on ice for 1 h, centrifuged, and resuspended in 2 mL HB. The suspension was embedded using a 2% What's already known about this topic?

FSHD is inherited as an autosomal dominant disease. And the causative genetic defect in FSHD is the loss of transcriptional repression of the Double Homeobox Protein 4 (DUX4) gene present in each repeat of the macrosatellite array (D4Z4) at chromosome 4q35. FSHD is not fatal disease, but there is no effective treatment. The traditional genetic method of FSHD diagnosis is by pulse field gel electrophoresis and Southern Blot which is a lengthy procedure.

What does this study add?

In this study, we explored the approach of combining Bionano optical mapping with Karyomapping as new strategy for prenatal diagnosis for rapid and accurate prenatal diagnosis of family FSHD1.
CHEF Genomic DNA Plug Kit (Bio-Rad) to avoid fracture of DNA molecules. The DNA plugs were incubated in 2.5 ml Lysis Buffer (Bionano Genomics, USA) containing 167 μl of proteinase K (Qiagen) at 50°C overnight, then washed with wash buffer and TE buffer (Bionano Genomics, USA). 2 μl of 0.5 U/μl Agarase (Thermo Fisher) enzyme was added to the plugs into a 1.5 ml microcentrifuge tube and the samples left to stand for 45 min at 43°C. The DNA mixture was dropped into the center of 0.1 μm Dialysis Membrane (EMD Millipore, USA) for 45 min to dry at room temperature.

2.1.2 | Bionano optical mapping detection

DNA was quantified using a Qubit 3.0 Fluorometer (Thermo Fisher Scientific, USA). The integrity of high-molecular weight DNA was determined by pulse-field gel electrophoresis (Pippin Pulse, Sage Science). Samples with ≥ 150kb molecular weight at the concentration of 45-200 ng/μl were used for labeling experiments. DNA labeling was carried out according to Bionano PrepTM Labeling - NLRS Protocol (Bionano Genomics #U30024), and consisted of four consecutive steps (Nick, Label, Repair and Stain). First, 300 ng of high molecular weight DNA was nicked using endonuclease Nb.BssSI in 10× Buffer 3.1 (Bionano Genomics, USA) for 2 h at 37°C. Second, 1.5 μl of 10× Labeling Mix (Bionano Genomics) and 1.0 μl Taq DNA Polymerase (Bionano Genomics) was added and the sample left to stand for 1 h at 72°C. Third, labeled DNA was repaired using 5 μl Repair Master Mix for 30 min at 37°C. Forth, DNA was stained overnight using 41.8 μl of Staining Master Mix (Bionano Genomics, USA). The stained DNA was quantified as above. DNA was used at 3-10 ng/μl, loaded into a Saphyr chip (Bionano Genomics, USA), and the fluorescent DNA molecules were imaged using the Saphyr instrument (Bionano Genomics, USA).

2.1.3 | The pretreatment and comparison of data

Data from the Bionano Saphyr bioinformatics mapping platform was used to exclude molecules with length < 150 kb or of low quality. We performed in silico digestion of the human reference genome (GRCh38) to generate the Nb.BssSI enzyme (CACGAG) reference map (CMAP file). Then the sample data and reference CMAP file were compared using Bionano Solve 3.2.1 software. A custom script was built by Liu Q et al. to extract and align the required D4Z4 target regions from whole-genome maps, to speed up subsequent data analysis and visualization.

2.1.4 | Determining the number of D4Z4 repeats, 4qA/4qB haplotype and 4q/10q chromosome

We extracted data from the D4Z4 target region from the whole genome maps of family FSHD1 members. There is only one Nb. BssSI restriction enzyme site in each D4Z4 repeat, and the distance between each restriction site is about 3.3kb, thus the number of D4Z4 repeats can be simply determined by calculating the number of restriction sites in the target region. The FSHD1 pathogenic haplotype 4qA has an additional Nb.BssSI site, 1.7kb further into the region distal to the last D4Z4 repeat unit, while the non-pathogenic 4qB haplotype does not. Therefore, we can also determine the 4qA/4qB haplotype of the sample DNA by identifying and counting this extra site. And the identification of 10qA/B is same as 4qA/B. The average length of DNA reads is about 150kb, which can be well covered all D4Z4 repeats in FSHD1. So it’s easy to recognize 4q35 and 10q26 homologous sequences through the analysis of fluorescence spectra of upstream and downstream near D4Z4 repeats.

2.2 | Karyomapping

2.2.1 | DNA sample preparation

Genomic DNA was extracted from the peripheral blood samples and fetal chorionic amniotic fluid samples using a DNA extraction kit (Omega blood/tissue DNA kit, Georgia, United States) according to the manufacturer’s instructions.

2.2.2 | Karyomapping and haplotype analysis

The target region haplotypes identified using the Humankaryomap-12v1.0 BeadChip array (Illumina, San Diego, USA) for whole-genome SNP detection, and iScan for SNP scanning. BlueFuse Multi 4.4 (Illumina, San Diego, USA) was used for karyomap
FIGURE 3  Bionano optical maps and genetic diagnosis of the proband and fetus of family FSDH1. The green bar shows the reference normal human DNA in the target chromosome 4, region 4q35, while the blue bars each show one optical map reading in the target region. Each red vertical line in the red boxes indicates schematically the presence of a D4Z4 repeat (3.3kb). The vertical line immediately to the right of the red box indicates the presence of the pathogenic 4qA allele, while its absence indicates the non-pathogenic 4qB allele. The blue box shows the different region between 4q35.2 and 10q26.3. (A) the proband’s pathogenic chromosome 4, showing that it has only 4 D4Z4 repeats, and the 4qA allele; (B) the proband’s normal chromosome 4, showing 19 D4Z4 repeats, and the 4qA allele; (C) the fetal pathogenic chromosome, showing 4 D4Z4 repeats and the 4qA allele. (D) the fetal normal chromosome showing 30 D4Z4 repeats and the 4qA allele. (E) our normal chromosome 4 control showing 26 D4Z4 repeats, and the 4qB allele; (F) our normal chromosome 10 control showing 12 D4Z4 repeats, and the 10qA allele.
The DUX4 gene is located in the region 4q35.2. The start and stop area (191005267-191007077) of this gene in analysis is selected for parameter setting, and the analysis area is about 1.7 Kb. The upper and lower 2 Mb regions were taken as the auxiliary analysis areas. A total of 138 SNP sites were identified in this region. All of them were located at the 5' end of the D4Z4 repeats.3

3 | RESULTS

3.1 | Bionano optical mapping

was used to perform genetic diagnose of the affected proband of family FSHD1. The sequencing depth of the target area was 45x. The results were visualized using Bionano Solve (version 3.2.1) software (Figure 3). The analysis revealed that proband had pathogenic sequences in the target chromosome 4 region (see Figure 3A): only 4 D4Z4 repeats, and also the pathogenic 4qA allele. The proband’s normal chromosome 4 is shown in Figure 3B, and had 19 D4Z4 repeats, and the 4qA allele. Compared with the results of the proband’s Southern Blot, the Bionano test more accurately identified: 1) the number of D4Z4 repeats; 2) the chromosomal allocation of the D4Z4 repeats; 3) the 4qA/B alleles; and, 4) the 4q35.2 and 10q26.3 regions, by their upstream and downstream sequences.

3.2 | Karyomap analysis

The FSHD1 family haplotypes were constructed using the Karyomap SNP-array method. The results of this analysis are shown in Figure 4.

The haplotype of the DUX4 allele of the affected proband (the fetus’ father) was named P1 and P2, while the haplotype of the fetus’ unaffected mother was named M1 and M2. FSHD1 is an autosomal dominant genetic disease and the family members that inherited the FSHD1 pathogenic sequences in the chromosome 4q35.2 region are shown as blue bars. The results indicated that there is no recombination in 4q35.2 region. The fetus clearly inherited the pathogenic chromosome fragment P1 and thus would most likely be affected by FSHD1 disease.

3.3 | Bionano optical mapping of fetal amniotic fluid cells

The results are shown in Figure 3C and 3D, and indicated that the fetus had one pathogenic chromosome 4 with 4 D4Z4 repeats and the 4qA allele, while its normal chromosome contained 30 D4Z4 repeats, and also the 4qA allele. The fetus’ pathogenic sequences were similar to those of its affected father (the proband), indicating that the fetus would likely be positive for FSHD1 disease. The results of Bionano optical mapping using fetal amniotic fluid cells were consistent with the results obtained by Karyomapping linkage-analysis using fetal amniotic fluid cells.

4 | DISCUSSION

The genetic diagnosis of FSHD1 is very difficult, because of the special DNA structural features involved in the genetics of this disease:
from culture cells. The main problem in amniotic fluid cell culture is the amount of DNA required is small and can be tested directly using the Karyomap technique requires two patients in the family, the nation in these or other distal regions of chromosome 4. Although 10q26 homology region, and also detect the presence of recombinant regions. This information may exclude the interference of the pathomechanism (PGD) 15. However, in the present study, we utilized Bionano optical mapping for prenatal diagnosis in a FSHD1 family with positive history. We used white blood cells from the proband (the father), proband's mother and fetal amniotic fluid cells. The pathogenic 4q35 region of the 3 family members (fetus, father, grandmother) all had only 4 D4Z4 repeats and the 4qA allele, and the D4Z4 repeats were clearly distinguished from the 10q26 repeats. Southern Blot is still the "gold standard" 6, but it is not so accurate because it detects the length of all D4Z4 repeats, and not their exact number. Research by Yi et al 8 and Qian et al 13 showed that Bionano optical mapping provided an accurate molecular diagnosis of FSHD, deriving haplotypes consistent with gold standard Southern blotting.

Karyomapping is mostly used in pre-implantation genetic diagnosis (PGD) 15. However, in the present study, we employed this technique for prenatal diagnosis along with Bionano optical mapping. Karyomapping can be used to perform linkage analysis for prenatal diagnosis of FSHD1 as an indirect diagnosis, through the detection of SNP loci, to construct family members' haplotypes, and to determine whether a fetus has inherited the family’s pathogenic chromosome 10. These haplotypes provide information on the entire 4q35 region, including the pathogenic D4Z4 and flanking 4qA regions. This information may exclude the interference of the 10q26 homology region, and also detect the presence of recombination in these or other distal regions of chromosome 4. Although the Karyomapping requires two patients in the family, the amount of DNA required is small and can be tested directly using amniotic fluid or villus DNA to make the test results quick and responsive to the original sample.

In this study, the DNA samples used for Bionano detection were from culture cells. The main problem in amniotic fluid cell culture is cell chimeric or cultural artifact 16. But we used the Karyomap chip to detect the DNA of the original fetal amniotic fluid sample, eliminating the possibility of fetal chromosome mosaic, and Bionano optical mapping was able to detect the microchimerism problem caused by the cell culture. The two results can be mutually confirmed, excluding the error caused by cell culture, reflecting the original situation of the fetus.

In summary, we utilized the approach of combining Bionano optical mapping with Karyomapping for rapid and accurate prenatal diagnosis in a FSHD1 family with positive history in this study. Bionano optical mapping was used for prenatal diagnosis of FSHD1, and effectively exclude interference by the homologous 10q26 repeat elements. The absence of Southern Blot analysis of fetal tissue is a limitation of the current study but some samples have demonstrated the application of Bionano optical mapping in the FSHD1 gene diagnosis. Besides, Bionano optical mapping can detect chromosome deletion, repeat, inversion 17,18. Qian Zhang et al 13 used this method to detect a repetition of a 4q35 region successfully. Although the price of Bionano optical mapping is a little higher now, with the commercial development and promotion of Bionano, the price will gradually decrease. Bionano optical mapping and karyomapping are performed respectively for detection of fetal amniotic fluid and culture cells samples, improving the accuracy and confidence of the prenatal diagnosis of FSHD1. We present that Bionano optical mapping combined karyomapping may provide a new approach of rapid and accurate prenatal diagnosis for FSHD1 families with positive history.

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CONFLICT OF INTEREST STATEMENT

The authors declare no competing interests.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available due to privacy or ethical restrictions.

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