Non invasive prenatal test for duchenne muscular dystrophy

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Objective
To investigate the accuracy and feasibility of a haplotype-based noninvasive prenatal test for Duchenne Muscular Dystrophy (DMD).

Methods
Seventeen families, each with a proband affected by DMD were recruited. The causative mutations in probands and their mothers were previously identified by multiplex ligation-dependent probe amplification (MLPA). Captured sequencing was performed on genomic DNA from parents and proband using customized hybridization probes targeted at highly heterozygous 2358 SNPs located within the 1M region flanking DMD gene and its coding region to acquire parental haplotypes and the linkage to pathogenic mutations. Maternal plasma DNA obtained at 12-25 weeks of gestation also underwent targeted sequencing to deduced fetal haplotypes assisted by parental haplotypes. The fetal genotypes in DMD gene were further validated by invasive procedures of prenatal diagnosis.

Results
The haplotype-based noninvasive prenatal test was successfully performed in all families. Four female and six male fetuses were identified to be normal. Four female fetuses was carriers and three male fetuses were DMD patients due to exons 49-52 deletion, exons 8-37 deletion and c.628G>T, respectively. All these results were consistent with those of invasive procedures.

Conclusion
Haplotype-based noninvasive prenatal Test for DMD using targeted sequencing is accurate and feasible and has potential applications in clinical practice.