Exome Sequencing in Fetuses with Structural Malformations

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Abstract: Prenatal diagnostic testing is a rapidly advancing field. An accurate diagnosis of structural anomalies and additional abnormalities in fetuses with structural anomalies is important to allow “triage” and designation of prognosis. This will allow parents to make an informed decision relating to the pregnancy. This review outlines the current tests used in prenatal diagnosis, focusing particularly on “new technologies” such as exome sequencing. We demonstrate the utility of exome sequencing above that of conventional karyotyping and Chromosomal Microarray (CMA) alone by outlining a recent proof of concept study investigating 30 parent-fetus trios where the fetus is known to have a structural anomaly. This may allow the identification of pathological gene anomalies and consequently improved prognostic profiling, as well as excluding anomalies and distinguishing between de novo and inherited mutations, in order to estimate the recurrence risk in future pregnancies. The potential ethical dilemmas surrounding exome sequencing are also considered, and the future of prenatal genetic diagnosis is discussed.

Keywords: exome sequencing; prenatal; fetus; prenatal diagnosis
1. Introduction

The incidence of congenital abnormalities in the UK is approximately 2.2% [1]. These are frequently first identified by ultrasound scan during pregnancy. There is a wide range of potential outcomes for fetuses with abnormalities. Some abnormalities, such as isolated cleft lip, can be corrected in early childhood with a surgical procedure, and often has minimal long-term impact on the child [2]. Other abnormalities, such as bilateral renal dysplasia or cerebral malformations, are associated with high morbidity and mortality [3,4]. There is little doubt that a chromosomal anomaly associated with a structural malformation significantly worsens prognosis.

Numerous genetic mutations have been associated with fetal structural abnormalities. These include aneuploidies, copy number variants (CNVs), loss-of-function single nucleotide variants (SNVs) and mis-sense SNVs [5–9]. Knowing the cause of a fetal structural abnormality may help clinicians to make an accurate prognosis regarding the pregnancy, and estimate recurrence risk for any future pregnancies. This helps the families to make informed decisions, including whether to terminate the pregnancy. Despite the importance of a diagnosis, currently only a minority of children affected by a developmental disease receive a genetic diagnosis, to the frustration of families, clinicians and researchers alike [10].

In this review, we will outline the current approaches used to investigate the genetic cause of prenatal structural abnormalities. We will then describe several studies that have recently used next-generation sequencing (NGS) for this purpose, and we will discuss advantages and disadvantages of this approach. Finally, we will outline some ethical issues that this technology raises, and discuss likely future directions of the prenatal genetic diagnostics field.

2. Current Techniques for Prenatal Genetic Diagnosis

2.1. Sampling Methods

Fetal DNA for genetic testing may be obtained invasively or non-invasively. Invasive methods include amniocentesis, fetal blood sampling and chorionic villus sampling (CVS). They yield cells from which fetal genomic DNA can be extracted. The major disadvantage of invasive sampling is that the risk of miscarriage increases by around 1% following a procedure [11–14]. Alternatively, fragmented cell-free fetal DNA (cffDNA) can be obtained non-invasively from maternal plasma [15], but there are presently limitations to its use in prenatal diagnostics.

2.2. Karyotyping

One invaluable tool for the detection of chromosomal aberrations associated with fetal and congenital abnormalities is karyotyping, where whole chromosomes are stained and examined using a light microscope. In classical cytogenetics, the stains (such as Giemsa (G-) stain) reveal patterns of light and dark bands that are unique to each chromosome. The technique was developed in the late 1960s and early 1970s, and it allowed researchers to distinguish between chromosomes of similar sizes for the first time [16,17]. As conventional full karyotyping provides information on the number and gross appearance of chromosomes, it can be used to detect potentially pathogenic chromosomal
aberrations including aneuploidy, deletions, duplications, and unbalanced translocations. Giemsa (G) banding has a highest resolution of 3–10 Mb [18].

A useful supplement to classical karyotyping is fluorescence in situ hybridisation (FISH). Fluorescent-tagged oligonucleotide probes complementary to a DNA sequence of interest are used to visualise chromosomal regions of interest. Locus-specific or gene-specific probes can be used to identify known aberrations that cause fetal or congenital abnormalities; for example, 22q11 deletions in DiGeorge syndrome, 7q11.23 deletions in Williams syndrome, and dystrophin mutations in Duchenne muscular dystrophy [19–21]. In another nice example of the clinical use of FISH, specific subtelomeric probes were used to identify an unbalanced subtelomeric translocation in a child with multiple congenital abnormalities, where classical cytogenetic analysis had indicated a normal karyotype [22]. Because FISH has to be “targeted” to a specific chromosomal anomaly, it requires a clinician to have a high degree of suspicion of that specific anomaly from the phenotype of the patient. Generally, fetal chromosome karyotyping, along with FISH where appropriate, is offered to families when a significant fetal anomaly is identified on ultrasound, or when there is a high risk of such an anomaly. In this population, karyotyping identifies a chromosomal anomaly in around 9% of cases [23].

2.3. Chromosomal Microarray (CMA)

Chromosomal Microarray, also known as array comparative genomic hybridisation (aCGH), detects CNVs that may be pathogenic, benign, or variants of unknown significance (VOUS). CMA has a higher resolution than G-band karyotyping, and can detect deletions or duplications as small as 1 kb, depending on the platform used [24]. However, these very high resolution arrays have disadvantages, including higher cost and increased detection of VOUS. Therefore, for clinical diagnostic purposes, microarrays with a resolution in the range of 10–400 kb are generally preferred [25,26]. The choice and resolution of the microarray “platform” used in prenatal diagnosis is the major discussion point occurring prior to CMA universal adoption for this role.

One limitation of CMA is that it is only able to detect unbalanced chromosomal rearrangements. Balanced rearrangements, such as reciprocal translocations, may lead to disease by disrupting genes, without detectable gains or losses at breakpoints [23,27]. However, in practice, many apparently balanced rearrangements detected by G-banding are not truly balanced at the DNA level, and CMA testing can be used to detect small regions of DNA loss or gain, and so clarify the exact nature of the rearrangement [28]. Many CMA platforms may not detect triploidy or low-level mosaicism [23,29]. However SNP based CMA platforms will. In the UK, quantitative fluorescence polymerase chain reaction (QF-PCR) is used prior to CMA as a rapid and cost-effective screen for common aneuploidy and triploidy [29].

Despite the limitations, CMA has been the diagnostic test of choice for several years in children and adults with developmental delay [30]. For fetuses with structural abnormalities, CMA has a diagnostic yield of ranging from 6% to 10% higher than chromosomal karyotyping [6,23,31]. Therefore, the UK is considering the introduction of prenatal CMA for all fetuses with structural anomalies only by the end of 2014 [32]. This is slightly different to the guidance from the American College of Obstetrics and Gynecology who recommend that CMA should be performed as the diagnostic procedure of
choice on fetuses with a structural abnormality, and may also be performed in structurally normal fetuses undergoing invasive prenatal testing [33].

2.4. Non-Invasive Prenatal Testing

Up to 50% of cell-free DNA in the plasma of a pregnant woman is fetal-derived [34–36]. It consists of DNA fragments with a size range of 30–510 bps, and a median of 162 bps [37]. The cffDNA can be obtained non-invasively; therefore in recent years there has been huge interest in using it for prenatal genetic diagnosis. Non-invasive prenatal testing (NIPT) refers to sequencing cffDNA to identify genetic mutations in the fetus. NIPT is able to detect autosomal trisomies and sex chromosome aneuploidies [36,38–41], CNVs [42,43], identify the sex of the fetus [44], Rhesus status [45], and single gene disorders such as achondroplasia and cystic fibrosis [46–48].

Regarding clinical practice, in the United States and China, use of NIPT to detect aneuploidies and fetal sex is already widespread [49,50]. Implementation for single-gene disorders is much slower because of lower demand and higher technical challenges [48]. In the UK, NIPT is currently only being provided by the National Health Service for sex determination, genotyping of fetuses at risk of Rhesus disease, and some single-gene disorders. However, the “Reliable accurate prenatal non-invasive diagnosis” (RAPID) and “Non-invasive prenatal diagnosis for single gene disorders” (NIPSIGEN) studies are investigating how to expand the implementation, and UK health professionals generally view NIPT positively, therefore it is likely that provision will be expanded to other genomic disorders in the near future [51].

Two proof-of-concept studies published in 2012 showed that it is possible to sequence the whole genome of a fetus non-invasively using cffDNA, to a sufficient depth to be able to call SNVs, using parental haplotypes to distinguish fetal from maternal variants [52,53]. However, the sensitivity and specificity of the SNV calling are as yet insufficient to consider using this approach in clinical practice. The sensitivity and specificity of identification of de novo mutations were particularly low [53], and yet for diagnostics this is very important, as de novo mutations are especially likely to be pathogenic [9,54–58]. Therefore, to identify potentially pathogenic SNVs in fetuses with structural abnormalities, NGS on fetal DNA obtained through invasive methods remains, for now, the superior choice.

3. Prenatal Exome Sequencing

3.1. Exome Sequencing as both a Research Tool and a Diagnostic Test

NGS can be used to identify SNVs throughout the genome, thus it has a much higher resolution than cytogenetic approaches. Exome sequencing is often favoured over whole genome sequencing, as it targets only coding regions, which represent 1%–2% of the entire genome, but contain around 85% of the mutations that cause known genetic disorders [59–61].

The first report of exome sequencing as a method to discover the genetic cause of a Mendelian disease was made in 2010, with the identification of mutations in DHODH as the cause of Miller syndrome [62]. In the few short years since then, exome sequencing has proved to be a remarkably fruitful research tool. At least one hundred genes that harbour mutations causing Mendelian disease
have been identified, and this rate of progress shows no signs of abating as yet [63]. Additionally, exome sequencing is increasingly being used in the clinical setting as a diagnostic test for patients with rare diseases. It has a diagnostic yield of around 25% [55, 64].

3.2. Prenatal Exome Sequencing: Proof-of-Concept

Despite the success of NGS, only a handful of studies have used it for prenatal gene discovery or diagnosis. The first two such studies, both published in 2012, used NGS to identify aneuploidy and chromosomal rearrangements. Dan et al., used very low-coverage whole-genome sequencing to detect aneuploidies and unbalanced chromosome variants in 13/62 fetuses [65], and Talkowski et al., used whole genome “jumping library” sequencing of amniocytes to identify an apparently balanced de novo translocation that disrupts CHD7, causing CHARGE syndrome in a single fetus [66].

The next two studies used exome sequencing at a depth sufficient to identify SNVs, in a very small number of fetuses. Yang et al., performed exome sequencing on 250 patients with Mendelian disorders, four of which were fetuses from terminated pregnancies [55]. In one of the fetuses, which had Cornelia de Lange syndrome, they found the cause of disease, which was a de novo splicing variant in the known gene NIPBL. Finally, Filges et al., used exome sequencing to identify the cause of a recessive, lethal ciliopathy phenotype in one family [67]. They sequenced the parents, their unaffected daughter, and post-mortem samples from two fetuses that were affected by the disease, and found compound heterozygous mutations in KIF14 in both affected fetuses.

3.3. Prenatal Exome Sequencing: Diagnostic Yield

In our recent study, we demonstrated that for fetuses with structural abnormalities, exome sequencing improves diagnostic yield over karyotyping and CMA alone [9]. The study was collaboration between the University of Birmingham (Birmingham, UK) and the Wellcome Trust Sanger Institute (Cambridge, UK), and it is the largest published cohort of fetuses with structural abnormalities to have been exome sequenced to date.

We gathered a cohort of 30 euploid fetuses with a diverse range of structural abnormalities detected by ultrasound. We exome sequenced them using the Illumina HiSeq platform in a trio design. That is, maternal and paternal samples were sequenced in addition to the affected fetus. We considered de novo rare coding mutations, and inherited recessive and X-linked rare coding mutations. It is important to distinguish between de novo and inherited mutations, in order to estimate the recurrence risk for future pregnancies. We considered SNVs and CNVs (which were identified directly from the exome sequencing reads using the CoNVex software [68]). These mutations were classified through a systematic manual process as highly likely to be causal, possibly causal, or unknown.

A list of 77 potential candidate de novo coding or splicing variants (mean = 2.6 per fetus, range 0–5) was found. Thirty-four (mean = 1.13 per fetus, range 0–4) variants were then confirmed as truly de novo as a result of subsequent capillary sequencing, in-keeping with the known germline mutation rate [69, 70]. For 3/30 fetuses we identified mutations that are highly likely to be causal [9]. That is, similar mutations in the same gene have previously been shown to cause a very similar phenotype in humans. One of these mutations was in the FGFR3 gene which is a negative regulator of bone growth; the features of the fetus were that of a lethal skeletal dysplasia thus supporting that the genetic
abnormality detected on exome sequencing was pathogenic. Another mutation was found in a different fetus in \textit{COL2A1}, mutations in this gene are known to cause type II collagenopathies including heart and limb defects, although the particular mutation that we detected had not previously been reported. This second fetus had increased nuchal translucency, tricuspid regurgitation, abnormal hands and feet and bilateral talipes, again supporting the results of exome sequencing. A third fetus which had ventriculomegaly and agenesis of the corpus collosum demonstrated a \textit{de novo} Xp22.2 deletion which resulted in the removal of most of the \textit{OFD1} gene which is known to cause orofaciodigital syndrome, and is associated with absence of the corpus callosum is approximately 40\% [71].

The diagnostic yield of this study was 10\%. Only one of these mutations was a CNV detected by CMA, highlighting the utility of exome sequencing, and increasing the detection rate of prenatal genetic abnormalities over that achieved by karyotyping and CMA alone [6,23,31]. Nevertheless, our diagnostic rate is lower than that found in exome sequencing studies of rare postnatal diseases [55,64]. There are several possible reasons for this. Our estimate of 10\%, being based on a relatively small sample size, has a broad confidence interval. Furthermore, it is likely that in some cases, mutations in the same gene will have different phenotypic manifestations between prenatal and postnatal stages of development [72]. Also, in some cases we only had phenotypic data based on ultrasound scans. There are many phenotypes that cannot be identified from an ultrasound scan.

Interpreting the clinical significance of the many unknown variants identified is generally the most challenging aspect of any exome sequencing study of rare disease [73,74]. For the reasons described, doing so in the context of prenatal phenotypic data is particularly difficult. However, as more exome sequencing is performed, databases of pathogenic mutations such as the DECIPHER database will be expanded [75], the diagnostic yield of exome sequencing, including in the prenatal context, is likely to increase. The ability to match SNVs to organ-specific anomalies, for example, the heart, would be particularly useful and may allow a “targeted” approach.

4. The Ethics of Prenatal Exome Sequencing as a Screening Tool

There is no technical reason why exome sequencing of fetal DNA obtained during pregnancy could not be used on structurally normal fetuses, as a screening tool rather than a diagnostic test. While this is not a proposal that is likely to be implemented in the near future, it is a responsibility of the prenatal genetic diagnostics community to begin to consider the relevant ethical issues [76].

The many thorny ethical issues surrounding exome sequencing in the clinical postnatal context have been extensively debated, chief among which are whether to report incidental findings, and how to report VOUS. Last year, the American College of Medical Genetics and Genomics (ACMG) recommended that mutations in a set of 57 known disease genes should always be reported where sequencing is used as a clinical test [77]. Fetuses were excluded from the ACMG recommendations. They have since revised these recommendations, saying patients have a right to opt out of receiving information on incidental findings [78]. The European Society of Human Genetics have adopted rather more conservative guidelines preferring a targeted approach to avoid discovering incidental findings [79], and the Deciphering Developmental Disorders project which is investigating the causes of rare genetic diseases in children in the UK is not returning incidental findings to participants under any circumstances [80]. Some researchers and ethicists object to the ACMG recommendations on the
grounds that the line between “actionable” and “non-actionable” conditions is far from clear, and the probabilistic nature of the link between genotype and phenotype will result in unnecessary distress for a proportion of families who receive information about mutations which will in fact never cause them to develop disease [81]. Some have argued that routinely returning incidental findings regarding a list of genes amounts to genomic screening [82].

When considering prenatal sequencing as a screening tool, the ethical dilemmas are similar but amplified, partly due to the possibility of termination of the pregnancy. If prenatal sequencing were to be used as a screening tool, in some cases, it would identify a variant that causes a severe, distressing, and lethal phenotype and is highly penetrant, at an earlier stage than an ultrasound scan could have found structural abnormalities. An example of such a variant might be missense changes in FGFR3 that cause thanatophoric dysplasia [8,9]. In this scenario, earlier detection is undoubtedly better for families. It avoids potentially devastating news later in pregnancy, in the neonatal period, or even later in childhood when developmental delay is clinically apparent. If the families elect to terminate the pregnancy, distress is generally less severe at an early stage of pregnancy [83]. For families who choose to continue with the pregnancy, early diagnosis may offer a more accurate prognosis, more time to prepare, and in some cases the option to start treatments earlier. Therefore, these families would definitely benefit from prenatal sequencing.

In other, less clear-cut cases, the disadvantages of prenatal sequencing as a screening tool may outweigh the advantages. Parents are potentially at risk of “truth dumping”, whereby a large quantity of information is given to parents and they are expected to interpret the information which professionals are unable to interpret with certainty, and then make a decision. Identification of “normal” genomic variation and VOUS is virtually inevitable during prenatal sequencing. For example, a predicted pathogenic variant may be identified in a known developmental disorder gene, but if it has never been reported before it may be very difficult to accurately predict the phenotype. The ethical issues of returning VOUS to families have been considered in the context of CNVs discovered by CMA. Some research suggests that receiving information on VOUS during pregnancy can be very stressful and distressing [84]. Therefore, some researchers and clinicians think that they should not be reported to families, and that their detection should be limited in the first place by using targeted tests [26,85]. Others think that it is paternalistic to withhold this information [86]. If VOUS were to be returned, it is imperative that families receive extensive genetic counselling pre- and post-test [33]. These issues are still under debate, but it would be important for clinicians and researchers to come to a consensus on the issue of reporting VOUS, prior to any use of prenatal exome sequencing as a screening technique, because interpreting mutations identified by exome sequencing is generally more difficult than those identified by CMA, and there will be a higher number of VOUS identified.

Another question is whether to return to families information on mutations that are likely to cause late-onset disease, and/or have incomplete penetrance, such as a BRCA1 variant that confers an 80% risk of developing cancer later in life [87]. Some argue that families have a right to this information to do with what they will, even if it will result in increased termination rates, and termination of some healthy fetuses [76]. An alternative is to do more targeted sequencing based on the indication for the test, so as to avoid incidental diagnoses.

Additional problems include issues surrounding paternity, both in terms of needing correct information to aid the diagnosis of de novo mutations, but also decision making when couples
disagree. It also reignites the perpetual ethical debate regarding the rights of the fetus itself. The potential to use exome sequencing to test for non-medical genetic markers and consequently create a “designer baby” is also a fear.

There are currently more questions than answers regarding the ethics of prenatal exome sequencing as a screening tool. Nevertheless, many pertinent issues have already been thoroughly discussed in the context of postnatal clinical sequencing, or interpretation of prenatal CMA results. While prenatal exome sequencing clearly poses additional specific ethical challenges, it is likely that with continued open debate amongst clinicians and researchers, along with sensitive and thorough genetic counselling to families, these can be overcome.

5. Next-Generation Sequencing: The Future of Prenatal Genetic Diagnostics

From a scientific perspective, it seems likely that next-generation sequencing is the future of prenatal genetic diagnostics. Nevertheless, many questions remain to be answered before prenatal next-generation sequencing could become widespread. These include issues of cost effectiveness, clinical utility, ethics, and interpretation of mutations. To address some of these, the Wellcome Trust and the Department of Health in the UK have awarded a Health Innovation Challenge Fund grant to the collaborative Prenatal Assessment of Genomes and Exomes (PAGE) project. This will involve the Wellcome Trust Sanger Institute, the University of Cambridge, the University of Birmingham, Birmingham Women’s Foundation Trust, University College London and Great Ormond Street Hospital (London, UK). One thousand fetuses with structural abnormalities, along with maternal and paternal samples, will undergo exome sequencing or whole genome sequencing. Qualitative work and an ethical review will also be performed to address the many issues this rapidly developing area is raising. The results of this study are expected to yield insights into the genetic causes of fetal abnormalities, and pave the way scientifically, clinically and socially for large-scale implementation of NGS in the UK’s prenatal arena.

In this review we have focused on exome sequencing, as it is currently more cost-efficient than whole-genome sequencing for clinical diagnostic purposes. However, for several reasons, we predict an eventual move towards whole-genome sequencing rather than exome sequencing for clinical diagnostic purposes, including in prenatal samples. First, there are many examples of non-coding mutations that can cause congenital abnormalities including pancreatic agenesis and malformations of the digits [88–91]. These variants would usually not be detected by exome sequencing. Additionally, while the costs of NGS are falling rapidly [92], if the costs of the exome capture step do not fall in line with this, at some point whole-genome sequencing may become more cost-effective than exome sequencing [93]. Finally, a major reason why whole-genome sequencing is currently often avoided is that interpretation of non-coding variants is very difficult. However, with large-scale whole-genome projects being planned, this is also likely to start becoming easier [94].

6. Conclusions

Next Generation Sequencing and examination of exomes is a powerful tool. In fetuses with congenital malformations it is helpful when delineating prenatal prognosis to be able to exclude chromosomal anomalies. However, the use of these new genetic techniques potentially allows
identification of causative single gene anomalies and equally may allow the exclusion of such risk and the reassurance to parents that the underlying aetiology of fetal structural anomalies is “de-novo”.

The PAGE study, undertaken in the UK from 2014 to 2016 will allow us to collect information on the use of such technology in prenatal diagnosis and to evaluate the potential for its use in clinical practice.

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Author Contributions

Fiona Mackie performed the literature search and first draft of the manuscript. Keren Carss, Sarah Hillman, Matthew Hillman and Mark Kilby conducted the proof-of-concept study described in Section 3.2 and all contributed to amendments of the manuscript to produce the final version of the paper.

Conflicts of Interest

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