Noninvasive Prenatal Testing: The Future Is Now

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Prenatal detection of chromosome abnormalities has been offered for more than 40 years, first by amniocentesis in the early 1970s and additionally by chorionic villus sampling (CVS) in the early 1980s. Given the well-recognized association between increasing maternal age and trisomy,1–3 the primary utilization of prenatal testing has been by older mothers. This has drastically reduced the incidence of aneuploid children born to older mothers.4 Although younger women have relatively low risks of conceiving a child with aneuploidy, the majority of pregnant women are in their late teens, 20s, and early 30s. As such, most viable aneuploid babies are born to these younger mothers.5 Invasive prenatal diagnosis (CVS and amniocentesis) is not a feasible option for all low-risk mothers, as these procedures carry a small but finite risk and would ultimately cause more miscarriages than they would detect aneuploidy. For this reason, a number of noninvasive tests have been developed—including first-trimester risk assessment at 11 to 14 weeks, maternal serum analyte (quad) screening at 15 to 20 weeks, and sonographic fetal structural survey at 18 to 22 weeks—all of which are designed to give a woman an adjusted (more accurate) estimate of having an aneuploid fetus using as baseline her a priori age-related risk. Ultrasound and maternal serum analysis are considered screening procedures and both require follow up by CVS or amniocentesis in screen-positive cases for a definitive diagnosis of a chromosome abnormality in the fetus. The ability to isolate fetal cells and fetal DNA from maternal blood during pregnancy has opened up exciting opportunities for improved noninvasive prenatal testing (NIPT). Direct analysis of fetal cells from maternal circulation has been challenging given the scarcity of fetal cells in maternal blood (1:10,000-1:1,000,000) and the focus has shifted to the analysis of cell-free fetal DNA, which is found at a concentration almost 25 times higher than that available from nucleated blood cells extracted from a similar volume of whole maternal blood. There have now been numerous reports on the use of cell-free DNA (cfDNA) for NIPT for chromosomal aneuploidies—especially trisomy

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extra copy of a chromosome—or monosomy (a missing chromosome)—and a number of commercial products are already being marketed for this indication. This article reviews the various techniques being used to analyze cell-free DNA in the maternal circulation for the prenatal detection of chromosome abnormalities and the evidence in support of each. A number of areas of ongoing controversy are addressed, including the timing of maternal blood sampling, the need for genetic counseling, and the use of confirmatory invasive testing. Future applications for this technology are also reviewed.

In 1959, Lejeune and colleagues demonstrated that Down syndrome is caused by an extra copy of chromosome 21 (trisomy 21 [T21]). This finding was of great clinical importance as Down syndrome is the single most common cause of mental retardation and has the highest incidence at birth of any chromosome abnormality. Children with Down syndrome have an increased risk for congenital defects and infectious morbidity, and all have some degree of mental retardation, with a mean intelligence quotient of 24. They also have a significantly shortened life span, with a 10- to 20-fold increased risk for leukemia and an increased risk for early-onset dementia. Although most fetuses with chromosomal aneuploidies are nonviable and lead to early miscarriage, a handful have the potential to survive to the newborn period. In some cases, as with trisomy 13 (T13) and trisomy 18 (T18), they are associated with significant clinical morbidity and a high rate of mortality shortly after birth. For all these reasons, considerable effort has been expended over the years to identify such fetuses early in pregnancy in order to provide couples with sufficient time to consider their reproductive options. Although patients may choose to decline prenatal testing, a discussion of the various options available for prenatal diagnosis is now considered a standard of care for all pregnant women in developed countries. Options currently recommended by both the American Congress of Obstetricians and Gynecologists (ACOG) and the American College of Medical Genetics (ACMG) include diagnostic testing, which requires that fetal cells be harvested using one of two invasive procedures (CVS or amniocentesis); or noninvasive screening, which uses a combination of first trimester risk assessment (FTRA) at 11 to 14 weeks and/or maternal serum analyte (quad) screening at 15 to 20 weeks and/or a sonographic fetal structural survey at 18 to 22 weeks to adjust a woman’s a priori risk of having an aneuploid fetus based on her age. The diagnostic accuracy of karyotyping cultured cells obtained by invasive testing has been found to be 97.5% to 99.8%. However, invasive testing also carries a risk for procedure-related miscarriage.

Aneuploidy screening by ultrasound and/or analysis of various maternal serum biochemical markers is primarily targeted at detecting Down syndrome and, to a lesser extent, T18; it has reported detection rates of 75% to 96% (depending on the screening approach utilized) with false-positive rates ranging from 5% to 10%. In addition, ultrasound and maternal serum analysis are considered screening procedures and both require follow-up by CVS or amniocentesis in screen-positive cases for a definitive diagnosis of a chromosome abnormality in the fetus.

The presence of fetal cells in maternal blood was initially reported in 1969, and the possibility that these cells could be isolated during pregnancy generated an exciting new noninvasive approach for identifying fetal genetic abnormalities. Several fetal cell types have been reported to exist in the maternal circulation, including fetal trophoblasts, lymphocytes, granulocytes, nucleated erythrocytes, and platelets. Fetal erythroid cells are the most commonly studied cell type as the existence of erythroid progenitors in adult blood is scarce in comparison...
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with its quantitative constitution in fetal blood. Nucleated erythrocytes are also fairly well differentiated and likely to have a limited life span in the maternal circulation. The detection of certain fetal aneuploidies and triploidy from a maternal blood sample has been accomplished using fluorescence in situ hybridization (FISH) for chromosome-specific DNA probes following various sorting and enrichment procedures. However, the inconsistent and low yield of fetal cells retrieved after sorting and enrichment has prevented the transition of this approach to routine clinical practice. Although a few commercial entities continue to persevere with aneuploidy testing of intact fetal cells, primarily utilizing newer molecular techniques, the focus of NIPT has shifted toward aneuploidy analysis of cfDNA.

In 1997, Lo and colleagues showed that cfDNA could be reliably detected in the maternal circulation during pregnancy. A year later, the same group reported that a surprisingly high mean concentration of fetal DNA (3.4%-6.2%) can be found in total maternal plasma DNA. This translates to approximately 20-25 times greater concentrations than that in the cellular fraction of maternal blood at the same gestational stage. Fetal DNA can also be detected as little as 10 μL of maternal plasma and serum, in amounts significantly higher than that available from nucleated blood cells extracted from a similar volume of whole blood. The approach of using cfDNA instead of fetal cells provided a far easier, less labor intensive, and less time consuming way to work with fetal DNA derived from the maternal circulation, and this opened up new opportunities for NIPT. Since that time, there have been numerous reports on the use of cfDNA for NIPT for fetal chromosome aneuploidies—primarily for T21, but also for T18 (Edwards syndrome), T13 (Patau syndrome), and sex chromosome anomalies—and a number of commercial products are already being marketed for this indication.

Cell/DNA Trafficking in Pregnancy

In his landmark 1953 publication designed to explain immunologic tolerance during pregnancy (and for which he was subsequently awarded the Nobel Prize in 1960), Dr. Peter Brian Medawar proposed that there may be a “true anatomic barrier between the mother and the fetus.” He proved to be wrong. Indeed, throughout pregnancy, fetal cells are constantly trafficking across the placenta into the maternal circulation and vice versa; some of these are undoubtedly fetal stem cells. As such, every woman who has ever been pregnant can be regarded as being a recipient of a stem cell “transplant,” and during pregnancy. It is present as early as 5 to 7 weeks of gestation, released continually by apoptotic cells throughout pregnancy, and is typically cleared from circulation within a matter of hours. It is now known that 3% to 10% of the cfDNA in the maternal circulation during pregnancy comes from the fetoplacental unit, most of which is shed from the placenta. Unfortunately, the absolute amount of fetal cfDNA is very small—typically less than 1 μg in 20 mL of whole blood—and reliably separating fetal cfDNA from maternal cfDNA is not technically feasible at this time. A number of companies have been spearheading the effort to develop a reliable and accurate commercial NIPT for fetal aneuploidy detection. All analyze the full cfDNA complement in the maternal blood without extracting or enriching the fetal fraction. Although the precise technology used by each

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20 fetal cells in 20 mL of maternal blood. Given their scarcity, efforts to date to isolate and purify these fetal cells for subsequent analysis have been largely unsuccessful, for this reason, attention has turned to cfDNA.

The presence of cfDNA in the circulation was documented in adult serum as early as 1947 (see Swaminathan and Butt for a review). It likely comes from lysis of cells within the circulation and from physiologic apoptosis occurring within various organ systems, and an extensive scavenging system exists to remove this DNA from the circulation. In 1997, Lo and colleagues reported the presence of cfDNA from the Y chromosome of male fetuses in the maternal plasma.
of these companies varies, they all rely on the same premise: that existing high-throughput approaches to DNA sequencing coupled with sophisticated data analysis will be able to detect abnormal amounts of chromosome-specific cfDNA in the maternal circulation in those pregnancies with fetal aneuploidy. The expectation is that one or all of these tests will prove to be superior to the currently available ultrasound/biochemical noninvasive tests and may eventually replace invasive prenatal testing altogether.

**Current Recommendations for Prenatal Diagnosis of Fetal Aneuploidy**

A discussion of the various options available for prenatal diagnosis is now considered a standard of care for all pregnant women in developed countries. Several options are discussed below.

**Diagnostic Testing**

This approach requires the direct harvesting of fetal cells during pregnancy for subsequent karyotype and/or genetic analysis. Two invasive procedures are commonly used.

CVS, which involves a biopsy of placental cells, can be performed either transcervically or transabdominally. The major benefit of CVS is that it can be performed early in the pregnancy, typically between 10 and 13 weeks of gestation. Preliminary results may be obtained in 1 to 2 days if a short-term direct culture of actively dividing villous cytotrophoblastic tissue is initiated, but in most cases a final karyotype result is issued within 7 to 10 days following long-term culture of mesenchymal cells from the villi. The disadvantage of CVS lies primarily in the source of cells being analyzed (the cells come from the trophoblast and not the fetus itself), which can lead, in a small number of instances (1%-2% of cases), to diagnostic ambiguity in the setting of confined placental mosaicism. In addition, the procedure-related pregnancy loss rate following CVS may be as high as 1%. Amniocentesis, on the other hand, involves placing a needle directly into the uterine cavity and aspirating some amniotic fluid containing fetal cells (amniocytes), which typically originate from fetal urine, pulmonary secretions, and skin. These amniocytes are cultured, harvested, and subjected to karyotype and/or genetic analysis. Compared with CVS, the cells primarily derive from the fetus itself, and the procedure-related pregnancy loss rate is generally regarded as being lower than for CVS (quoted as 1:200 to 1:600). Amniocentesis is usually only offered after 15 weeks (earlier amniocentesis is associated with higher pregnancy loss rates) and fewer viable cells are retrieved, precipitating long culture times and consequently a longer turnaround time (typically 8-14 days).

FISH, using chromosome-specific fluorescence-tagged probes, has provided a rapid way to detect the most common fetal aneuploidies found at the time of prenatal testing. The prenatal FISH panel typically targets chromosomes 13, 18, 21, X, and Y, and results from FISH analysis of interphase nuclei derived from chorionic villi or amniocytes may be available within 24 to 48 hours. The sensitivity and specificity of prenatal FISH for detection of all aneuploidies targeted by the typical FISH chromosome panel is > 99.6% and > 99.98%, respectively, and a large multicenter retrospective study demonstrated an extremely high concordance rate (99.8%) between FISH on interphase amniocytes and standard cytogenetic analysis for the specific chromosome abnormalities that the FISH panel is designed to detect.

Although prenatal FISH provides a swift way to diagnose common fetal aneuploidies, its diagnostic scope has reduced sensitivity compared with conventional cytogenetic analysis; that is, cases with cytogenetic abnormalities other than the most frequent ones (eg, translocations, inversions, markers) will not be identified by this technique. Furthermore, aneuploidy due to familial Robertsonian translocations cannot be identified by FISH because this requires visual inspection of the G-banded karyotype. For these reasons, FISH results should always be followed by routine chromosome analysis for a complete cytogenetic evaluation of the fetus.

Chromosomal microarray analysis (CMA) has provided a way to detect submicroscopic imbalances that remain undetected by conventional cytogenetic analysis. Unlike FISH, CMA permits high-resolution assessment of the entire genome. CMA can either be performed using array-comparative genomic hybridization (aCGH) or single nucleotide polymorphism (SNP) oligonucleotide microarray analysis (SOMA). Although both techniques offer comparable detection of copy number imbalances (also referred to as copy number variances), SOMA is able to utilize the SNP genotype information to detect triploidy, uniparental isodisomy, and consanguinity. SOMA can also assess zygosity in multiple gestations and, if a maternal sample is concurrently run, SOMA can detect maternal cell contamination. The recent Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD) Multicenter Prenatal Microarray Study indicated 100% concordance between CMA (aCGH and SOMA) and conventional cytogenetic analysis for the detection of nonmosaic common aneuploidies. In this cohort, CMA
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detected clinically relevant genomic imbalances in an additional 2.5% of cases that had a normal karyotype by G-band analysis.61

Noninvasive Testing

Noninvasive testing grew out of a desire to avoid direct contact with the growing fetus/placenta and concomitantly risking the health of the fetus. NIPT refers specifically to techniques that evaluate fetal cells or cfDNA in a blood sample drawn from the mother during pregnancy. However, for the purposes of this article, it applies to any form of NIPT for fetal aneuploidy. Importantly, it is a screening test, not a diagnostic test. This is a critical distinction and should be explained carefully to every patient during prenatal counseling. A screening test will determine if an individual patient is at high or low risk of having an aneuploid fetus; it does not confirm or refute the diagnosis. If a patient is assessed as high-risk on the basis of her NIPT results (typically regarded as a risk > 1:200) or if she is not satisfied with a simple risk estimate, then she may choose to proceed with invasive testing (CVS or amniocentesis). Such tests are diagnostic in that, with very rare exception (such as mixing up samples in the laboratory and low level mosaicism), they can definitively confirm or refute the presence of a chromosome abnormality.11-15

Prenatal screening has been defined as “the identification, among apparently normal pregnancies, of those at sufficient risk for a specific fetal disorder to justify subsequent invasive and/or costly prenatal diagnostic tests or procedures.”96 The objective is to develop a test with a high detection rate (ideally > 95%) and low false-positive rate (< 1%). Several NIPT options are currently approved and recommended by both ACOG9 and ACMG.10 Current NIPTs perform just as well in women over the age of 35 (advanced maternal age) as they do in younger women; as such, it is no longer necessary or appropriate to counsel women of advanced maternal age to proceed directly to amniocentesis.9,10,44

FTRA is typically done at 11 to 14 weeks of gestation or, more correctly, at a crown-rump length of 45 to 84 mm.9,10,20,22,24,26,28 It involves two elements, both of which are required to accurately interpret the test: the sonographic nuchal translucency, and maternal serum biochemical testing for two analytes (β-human chorionic gonadotropin [β-hCG] and pregnancy-associated plasma protein-A). Based on these results, as well as her age and a few other minor variables (such as body mass index and singleton or twin pregnancy), the patient can be given an adjusted risk that she may be carrying an aneuploid fetus.

Maternal serum analyte (quad) screening can be offered at 15 to 20 weeks of gestation.9,10,18,20,25,26 It involves measuring circulating levels of four biomarkers in the mother’s blood: maternal serum alpha-fetoprotein, β-hCG, unconjugated estriol, and inhibin-A. Again, based on these results, as well as her age, the patient can be given an adjusted risk that she may be carrying an aneuploid fetus. Such testing has been validated for twins (although the detection rate is lower20), but not for triplets or other higher-order multiple pregnancies.

FTRA can be combined with the quad screen, but this should only be done using an integrated algorithm that does not include both β-hCG measurements.10,23,27,29,56,63,64 A single result should then be given to the patient once the entire integrated screen is completed. Independent assessment of T21 risk followed by second trimester screening (serial testing) is generally discouraged because of the high false-positive rates associated with this approach.20,65

The sonographic fetal structural survey is not part of the second trimester quad screen for fetal aneuploidy. That said, it is commonly offered to women at 18 to 20 weeks of gestation, and allows the obstetric care provider to screen the fetus for two reasons: to identify the presence or absence of a major structural defect, such as a cardiac defect or neural tube defect, which may or may not be associated with a chromosomal abnormality; and to identify the presence or absence of so-called soft markers of fetal aneuploidy. These soft markers are not structural abnormalities per se, and include such findings as short femurs, thickened nuchal fold, renal pyelectasis, or echogenic bowel, but may suggest the presence of an underlying fetal aneuploidy. This is especially true when several of these sonographic markers are seen together. Absence of the fetal nasal bone has been shown to be both sensitive and specific in identifying fetuses with T21.66,67 However, overall, aneuploidy detection by ultrasound is not especially sensitive, and is dependent in large part on the skill and experience of the sonographer. In one landmark study of 15,000 women screened by routine ultrasound at two separate intervals during pregnancy, only 35% of fetal anatomic abnormalities were detected, and only 17% of these
were detected prior to 24 weeks of gestation. Moreover, because 40% to 50% of fetuses with T21 have no abnormalities on ultrasound, these sonographic markers have a low sensitivity and specificity for detecting T21. As such, ultrasound is not recommended as a primary screening tool for T21. Although rare (1:20,000 to 1:40,000 pregnancies), ultrasound is more sensitive when it comes to the diagnosis of T18 and T13, because the majority of these fetuses have major structural anomalies with or without fetal growth restriction.

### Weaknesses of Current NIPT Options

Obtaining a fetal genetic test result early in the pregnancy facilitates early reproductive decision making. In this regard, FTRA has a significant advantage over second trimester maternal serum quad screening or integrated screening. However, FTRA requires both a blood test and an ultrasound, which typically entails two prenatal visits. Although these noninvasive screening tests are safe for the pregnancy, they are primarily targeted at detecting T21 (and to a lesser extent T18) and they have poor accuracy with false-negative rates between 12% and 23% and false-positive rates between 1.9% and 5.2%. The performance of these tests for the detection of T21 is summarized in Table 1.

### Next-Generation NIPT Using cfDNA

Given these weaknesses, several companies have focused on the analysis of cfDNA in a sample of maternal blood collected in the first trimester to develop a more accurate and reliable NIPT. There are currently two primary next-generation sequencing approaches for gathering genetic data from cfDNA. The first, massively parallel shotgun sequencing (MPSS), sequences DNA fragments from the whole genome, whereas the second, targeted sequencing, selectively sequences specific genomic regions of interest.

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### MPSS and Counting

MPSS is a high-throughput technique that uses miniaturized platforms for sequencing large numbers of small DNA sequences called reads from the entire genome. This approach allows for tens of millions of short-sequence DNA tags or fragments (typically 25-36 bp in length) to be sequenced rapidly and simultaneously in a single run. After sequencing the cfDNA present in the maternal plasma, the chromosomal origin of each 25- to 36-bp DNA fragment is obtained by comparison of the sequence data from each DNA fragment with a euploid reference copy of the human genome. Fragments are categorized by chromosome (these include maternal and fetal DNA) and the number of reads mapping to the chromosomes of interest are compared with the number of reads mapping to one or more presumably normal reference chromosomes. This procedure is referred to as counting. If the amount of a chromosome-specific sequence exceeds the threshold that represents a normal (disomic) chromosome, the result is reported as positive for trisomy for that chromosome (Figure 1). A trisomic fetus has 50% more genetic material because of the extra chromosome (3 copies), resulting in an increase in the relative amount of cfDNA from the affected chromosome found in the maternal plasma. It is precisely this difference that the test attempts to detect. This difference is quantitative, not qualitative. In other words, no effort is made to distinguish maternal from fetal DNA. Because maternal DNA is the majority of cfDNA sample, the incremental difference due to fetal trisomy is very small when maternal and fetal DNA measurements are combined. This means that the ability to detect the increased chromosomal dosage resulting from fetal aneuploidy is directly related to the fraction of fetal cfDNA in the maternal circulation. For example, chromosome 21 represents approximately 1.5% of the total genome and an extra copy (T21) would increase the amount of DNA from chromosome 21 from the expected 1.5% to 2.25% (a 50% increase). If the fraction of fetal cfDNA in the maternal circulation is 10%, the relative change in the total cfDNA sample would only increase from 1.5% to 1.575% [1.5% (copy # = 2) × 0.9 (90% maternal)] + [2.25% (copy # = 3) × 0.1 (10% fetal)] = 1.575%. At lower fetal fractions, the increase becomes more marginal. Because fetal fraction tends to rise with increasing gestational age (although there is significant variation from individual to individual), this consideration is particularly important at early gestational ages. The ability to distinguish these minor differences with
### Performance Parameters of Noninvasive Screening Tests for Fetal Trisomy 21

<table>
<thead>
<tr>
<th>Risk Factor/ Biomarker&lt;sup&gt;a&lt;/sup&gt;</th>
<th>GA (wks)</th>
<th>Sensitivity (%)</th>
<th>Screen Positive (%)</th>
<th>Specificity</th>
<th>DR&lt;sup&gt;b&lt;/sup&gt; (for a 5% FPR) (%)</th>
<th>FPR&lt;sup&gt;c&lt;/sup&gt; (for an 85% DR) (%)</th>
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<td>First trimester risk assessment</td>
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<td>1 in 25</td>
<td>79-90</td>
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<tr>
<td>— Targeted sequencing + DANSR + FORTE</td>
<td>—</td>
<td>&gt; 10</td>
<td>100</td>
<td>—</td>
<td>99.7-100</td>
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<td>Sparks AB et al, Sparks AB et al, Ashoor G et al.</td>
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<tr>
<td>— Targeted sequencing + PS</td>
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<td>&gt; 9</td>
<td>100</td>
<td>—</td>
<td>99.8-100</td>
<td>—</td>
<td>Zimmermann B et al, Nicolaides KH et al.</td>
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<sup>a</sup>All serum biomarker results include maternal age.

<sup>b</sup>DR refers to the percentage of affected pregnancies (those with a Down syndrome fetus) called screen-positive by the test. In this table, DR data are reported for a fixed 5% FPR.

<sup>c</sup>FPR refers to the percentage of unaffected (normal) pregnancies called screen-positive by the test. In this table, FPR data are reported for a fixed 85% DR.

DANSR, Digital Analysis of Selected Regions; DR, detection rate; FORTE, Fetalfraction Optimized Risk of Trisomy Evaluation; FPR, false-positive rate; GA, gestational age; MPSS, massively parallel shotgun sequencing; PS, Parental Support™ (Natera; San Carlos, CA).
high degree of confidence requires a large number of reads from chromosome 21, typically > 95,000. Because MPSS obtains sequence information from all chromosomes and because chromosome 21 represents approximately 1.5% of the human genome, it is estimated that approximately 6.3 million uniquely mapped reads from the entire genome would be required to ensure sufficient chromosome 21 counts to make this distinction. The efficiency of MPSS is such that only approximately 25% of reads are uniquely mapped, which requires approximately 25 million raw sequencing reads per sample covering the entire genome in order to return sufficient data for analysis. This is especially important if assessment of additional clinically relevant chromosomes (13, 18, X, and Y) is considered. The requirement for a large number of reads necessitates the use of traditional large sequencing machines and represents enormous redundancy especially because the clinically significant chromosomes represent only approximately 14% of the human genome. In addition, throughput is limited, which could potentially affect turnaround times. An important limitation of counting appears to be the variability with which standard polymerase chain reaction conditions amplify certain chromosomes, and this variability seems to be linked to the GC base content of an individual chromosome. This limitation has ramifications for clinical testing because it means that the accuracy rate for detection of fetal aneuploidies using MPSS will be different for each chromosome assessed. Indeed, reported detection rates for T13, T18, and T21 all show accuracies to be highest for T21, followed by T18, and are lowest for T13. The issue of GC content has, to some extent, been dealt with using newer bioinformatics algorithms, which has led to improved sensitivity and specificity in the detection of T13 and T18.

**Targeted Sequencing**

Instead of sequencing random genomic fragments from all chromosomes, targeted sequencing selectively amplifies specific genomic regions of interest (such as nonpolymorphic loci on chromosomes 21 and 18, or specific polymorphic loci of interest) and then reads and counts only those specific sequences. This means that nearly all sequences are utilized for assigning a diagnosis. This strategy significantly reduces the total number of reads analyzed with a concomitant improvement in efficiency and a 10-fold reduction in overall costs. Also, the reduced read requirement enables the use of small bench-top sequencers that afford both financial and practical benefits to the laboratory. The ability to selectively sequence specific regions of the genome in cfDNA allows for a focused analysis of clinically important chromosomes such as 13, 18, 21, X, and Y.
targeting methods still appear to vary from chromosome to chromosome. Specifically, detection rates using quantitative read counting remain the highest for T21, followed by T18, and then T13.82

Analysis and Interpretation of Sequence Data

Just as important—and perhaps more important—than the technology used for genomic sequencing is the choice of bioinformatics platform for post-hoc data analysis. Several approaches have been taken.

Z-Score

The first approach to determine the difference in the total cfDNA attributed to fetal trisomy was based on a simple z-score. The z-score reflects the number of standard deviations the proportion of reads from a particular chromosome (in relation to the proportion of reads from all other chromosomes) is above the mean. If the amount of a chromosome-specific sequence exceeds a threshold that would be expected if the fetal karyotype was normal (euploid)—typically a z-score of > 2.5—the result is reported as positive for trisomy for that chromosome (Figure 1). The basic z-score method does not account for the variation in GC base content from chromosome to chromosome and, as such, is subject to different aneuploidy detection rates depending on the specific chromosome assessed.

Z-Score With GC Correction

The observation that the GC content of an individual chromosome determines how much of this chromosome is represented in the amplified sample and consequently its detection rate led to a bioinformatics approach to account for these differences. Instead of using read counts to directly calculate the proportion of a specific chromosome of interest, a GC-corrected read count is used with the purpose of eliminating the GC bias.79 As hypothesized, the detection rates for T13 and T18 are indeed improved using z-scores derived after quantitative correction for GC content,79 and additional improvements can be achieved using other modifications such as use of a non-repeat-masked reference genome instead of a repeat-masked reference genome.78

Z-Score With GC Correction Using an Internal Control

A more recent strategy utilizes a specific internal reference chromosome for each chromosome being assessed. The optimal internal reference chromosome is one that has a similar GC content to the chromosome of clinical interest.83 This approach appears to be markedly more adept at detecting aneuploidies other than T21, and a recent proof of concept study using this approach demonstrated 100% accuracy for detection of T13, T18, T21, 45,X, and 47,XXY in a small sample of 32 aneuploid cases.83

Normalized Chromosome Value

The normalized chromosome value (NCV) approach differs in the normalization process that compares the reads from the chromosome of interest with the number of counts from a reference set derived from an unaffected group of samples.77,84 The NCV algorithm helps to minimize the intra- and inter-run sequencing variation.84

Parental Support™

In contrast to the quantitative methods previously discussed in this review, Parental Support™ (PS; Natera, San Carlos, CA) focuses on measuring single nucleotide polymorphisms (SNPs). By measuring polymorphic loci, this approach is able to extract multiple pieces of information (including the number and identity of each allele) from each sequence read. PS then incorporates allelic information from the mother (and from the father, if available) to model a set of hypotheses (viz, monosomy, disomy, or trisomy), corresponding to different genetic inheritance patterns and crossover locations for every possible copy number count. Bayesian statistics then assign a probability to each hypothesis, and a maximum likelihood estimation analysis is performed to select the most likely hypothesis and calculate the probability of that hypothesis being correct.85

Commercial NIPT: Are We There Yet?

A number of companies have been spearheading the effort to develop the next generation of NIPT tests, including Sequenom Center for Molecular Medicine (San Diego, CA), Verinata Health (Redwood City, CA), Ariosa Diagnostics (San Jose, CA), and Natera. These companies all use a sequencing-based approach for gathering the genetic information contained within the cfDNA. In some cases, MPSS is the sequencing methodology of choice, whereas targeted sequencing is utilized by others. Each entity utilizes a unique and proprietary algorithm for interpretation of the genetic data. Although the exact technology may vary, the implications for clinical practice are the same; namely, these are all screening tests performed by analyzing cfDNA in a sample of maternal blood, and all positive test results should be confirmed by amniocentesis or CVS before acting upon the information. Detection rates reported by the commercial entities differ, as does the scope of chromosomal aneuploidies assessed. These, together

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with the specific analysis technique offered, are described below and summarized in Table 2.

The Sequenom Center for Molecular Medicine uses the MPSS approach with z-score GC correction. Data from its international collaborative study indicate sensitivity > 91% for T13 and > 99% for both T21 and T18.78 The study reported specificity for all three chromosomes to be > 99.6%.78

Verinata Health uses MPSS with the NCV algorithm. Published in 2012, the Maternal Blood Is Source to Accurately Diagnose Fetal Aneuploidy (MEDLISSA) study was conducted as a prospective, multicenter, observational study with blinded, nested, case-control analyses, and demonstrated a high sensitivity and specificity for the detection of T21, T18, and T13, as well as sex chromosome anomalies.77

Ariosa Diagnostics uses a targeted sequencing approach, which it calls Digital Analysis of Selected Regions (DANSR), coupled with its post-hoc bioinformatics algorithm, Fetal Fraction Optimized Risk of Trisomy Evaluation (FORTE), which is designed to account for an individual’s age-related risks and fetal fraction of the sample. This approach is closely related to MPSS in that it also uses counting, but includes an initial targeted amplification step in which approximately 400 loci of each chromosome of interest are selectively amplified prior to MPSS analysis. This targeted amplification results in an improvement in sequencing efficiency, with 100,000 to 300,000 reads achieved per target chromosome.75,80,86 DANSR has recently been shown to detect T21, T18, and T13 with good accuracy and greater efficiency than MPSS alone75,82 (Table 2).

### Table 2

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<tr>
<th>Company</th>
<th>Technical Approach</th>
<th>Accuracy of Detection</th>
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<td><strong>Sequenom Center for Molecular Medicine</strong> (San Diego, CA)</td>
<td>MPSS</td>
<td>T21: 98.6, T18: 100</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td></td>
<td>T13: 91.7</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td></td>
<td>45,X: —</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td></td>
<td>47,XXY: —</td>
<td>—</td>
</tr>
<tr>
<td><strong>Verinata Health</strong> (Redwood City, CA)</td>
<td>MPSS</td>
<td>T21: 100, T18: 97.2</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td></td>
<td>T13: 78.6</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td></td>
<td>45,X: 93.8</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td></td>
<td>47,XXY: —</td>
<td>—</td>
</tr>
<tr>
<td><strong>Ariosa Diagnostics</strong> (San Jose, CA)</td>
<td>Targeted sequencing + DANSR</td>
<td>T21: 100, T18: 98</td>
<td>Yes (SNPs)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>T13: —</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>45,X: —</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td></td>
<td>47,XXY: —</td>
<td>—</td>
</tr>
<tr>
<td><strong>Natera</strong> (San Carlos, CA)</td>
<td>Targeted sequencing + PS</td>
<td>T21: 100, T18: 100</td>
<td>Yes (SNPs)</td>
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<td></td>
<td></td>
<td>T13: 100</td>
<td>Yes</td>
</tr>
<tr>
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<td></td>
<td>45,X: 100</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td></td>
<td>47,XXY: 100</td>
<td>—</td>
</tr>
</tbody>
</table>

*Prospective study.

DANSR, Digital Analysis of Selected Regions; FORTE, Fetal Fraction Optimized Risk of Trisomy Evaluation; MPSS, massively parallel shotgun sequencing; PS, Parental Support™ (Natera); SNPs, single nucleotide polymorphisms; T13, trisomy 13; T18, trisomy 18; T21, trisomy 21.
permits aneuploidy detection using approximately 1 million raw reads per subject, enabling analysis of 96 subjects per sequencing lane. In contrast, MPSS evaluates the entire genome and requires approximately 25 million raw reads per subject, which limits sequencing throughput to 4 to 6 samples per lane. Thus, DANSR may enjoy an advantage over MPSS in terms of sequencing cost and throughput. As with all purely quantitative methods, however, the approach is dependent on chromosomes having low amplification variability (GC content influenced), which may limit its diagnostic accuracy for some chromosomes.

The ability to extend NIPT to the sex chromosomes is important, because these variants represent almost half of the at-birth chromosomal abnormalities and an even higher fraction in the mid-trimester when prenatal screens are typically offered.

Natera has been working in the field of genetic testing since 2009 with a focus on preimplantation genetic diagnosis, molecular karyotyping of products of conception, and noninvasive paternity testing using cfDNA. More recently, the company has become interested in NIPT and uses a novel approach called PS, which combines a targeted amplification step (that targets approximately 2000 polymorphic loci per chromosome under investigation chosen from cross-ethnic databases to avoid interethnic bias) with sophisticated statistical analysis. PS utilizes between 500,000 and 2 million sequence reads per chromosome, a number that falls between MPSS and DANSR. The PS approach has a number of distinct advantages. First, it is based on allelic distribution patterns rather than quantifying chromosome-specific read counts and, as such, does not suffer from chromosome amplification variability or require the use of a reference chromosome. PS is therefore able to determine fetal copy number at chromosomes 21, 18, 13, X, and Y with similar accuracy, and it has the same accuracy across all tested chromosomes regardless of fetal fraction. Owing to the lack of chromosome specificity, and because PS does not require comparison against a reference chromosome, the technology has the capacity to detect sex chromosome aneuploidies (45,X; 47,XXY, and 47,XYY), uniparental isodisomy, and triploidy. The ability to extend NIPT to the sex chromosomes is important, because these variants represent almost half of the at-birth chromosomal abnormalities and will enable testing to be carried out on a small desktop sequencing machine, thereby improving efficiency and significantly shortening turnaround time. Recent data have shown that the PS informatics-based method can noninvasively detect fetuses with T13, T18, T21, 45,X and 47,XXY with high sample-specific calculated accuracies for each individual chromosome and across all five chromosomes, and preliminary data from an improved version of the PS method, termed Next-Generation Aneuploidy Test Using SNPs (NATUS), show that it delivers similar performance but produces a no-call rate similar to other reported techniques. In a recent independent, prospective, blinded study of 242 singleton pregnancies in the United Kingdom, maternal blood was collected prior to clinically indicated CVS at 11 to 13 weeks of gestation. cfDNA was isolated from maternal plasma, subjected to targeted multiplex PCR amplification followed by sequencing of 19,488 SNPs covering five chromosomes (13, 18, 21, X, and Y), and analyzed using the NATUS algorithm. Results were reported in 94.6% (229/242) of cases and compared with the fetal karyotype obtained from CVS. In subjects for whom a result was reported, all fetuses with aneuploidy (including 25 with T21, 3 with T18, 1 with T13, 2 with Turner syndrome, and a case of triploidy) were accurately detected with no false-positive or false-negative results. Fetal gender was also correctly determined in all cases. The “no call” rate was 5.4% (13/242), which included two cases of T21. The authors conclude that “...cfDNA testing in maternal blood using targeted sequencing of SNPs at chromosomes 13, 18, 21, X, and Y and use of the NATUS algorithm holds promise as an accurate method for detecting fetal autosomal aneuploidies, sex chromosome...
aneuploidies, and triploidy in the first trimester of pregnancy."

In summary, these four approaches have more in common than they differ. The major differences appear to be in the post-hoc bioinformatics and how the data are analyzed once the DNA fragments are sequenced. They all give a result within 2 weeks from a single maternal blood sample drawn early in pregnancy, and all appear to have a better sensitivity (> 99%) for T21 detection (Table 1) than current recommended screening options. The major advantage of these tests is in their specificity. Using routine serum analyte screening, the likelihood that a positive screening test result actually represents a true positive (ie, a T21 karyotype on amniocentesis) is only approximately 2% to 4%. In contrast, using cfDNA technology, the vast majority (> 99%) of screen positives are likely to be true positives.

Future Applications for the Technology

In addition to prenatal diagnosis, analysis of cfDNA in maternal blood also has the potential to establish the Rh genotype of the fetuses and identify fetuses with congenital adrenal hyperplasia. Fetal sex determination is also possible from an early gestational age, although this raises a number of ethical concerns and, at least for now, is best reserved for clear medical indications. The advent of chromosomal microarray technology has uncovered multiple new genomic regions that lead to a clinical phenotype when deleted or duplicated. These types of aberrations are commonly referred to as microdeletions and microduplications, and their clinical consequences can be observed in pediatric populations. Examples include the 22q11.2 microdeletion syndrome, which causes DiGeorge syndrome and occurs with a frequency of 1 in 4000 births, and Prader-Willi syndrome, which can result from a microdeletion of the proximal 15q11.2 region. It seems obvious that future efforts in NIPT will be directed at detecting these smaller regions of genomic imbalance. However, the clinical consequences of microdeletions and microduplications are not always certain, they may be extremely variable, and in some cases may be unknown. Initial microdeletion/microduplication targets are sure to be restricted to regions with well characterized clinical outcomes. NIPT techniques using targeted amplification (DANSR and PS/NATUS) could be expanded to target detection of subchromosomal genomic imbalances (microdeletions/microduplications), as well as single gene disorders and/or carrier status for paternally inherited mutations.

Conclusions

Although safe for the pregnancy, currently available biochemical/ultrasound-based noninvasive screening tests for fetal aneuploidy have poor accuracy with false-negative rates between 12% and 23% and false-positive rates between 1.9% and 5.2%. Indeed, using current NIPT methodology, one out of every six cases of Down syndrome will go undetected. And, of all women deemed to be at high risk of having an aneuploid fetus, more than 95% are actually carrying a healthy baby, leading to unnecessary anxiety and miscarriage risk from amniocentesis or CVS. NIPT of cfDNA in a sample of the mother’s blood can significantly increase the detection rate of fetuses with T21 and decrease false-positive results. The accuracy of NIPT depends on the precise technology used and on the percentage of fetal DNA in a pregnant woman’s blood. NIPT has not been sufficiently validated in women at low risk for fetal aneuploidy. As such, it is not currently recommended for all pregnant women. Indications for the use of cfDNA for fetal aneuploidy are summarized in Table 3. If NIPT is ever to completely replace conventional cytogenetic analysis following CVS or amniocentesis, it will need to match the diagnostic accuracy as well as the scope of anomalies that can be detected. The diagnostic scope of traditional prenatal cytogenetic analysis has recently been extended to include detection of microdeletions and microduplications using new genomic technologies such as

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**TABLE 3**

Indications for Considering the Use of Cell-Free Fetal DNA Analysis for Fetal Aneuploidy

- Maternal age ≥ 35 years at delivery
- Fetal ultrasonographic findings indicating an increased risk for aneuploidy
- Personal or family history of a prior pregnancy with a trisomy
- Positive test result for aneuploidy, including first trimester, sequential, or integrated screen, or a quadruple screen
- Parental balanced Robertsonian translocation with increased risk for fetal trisomy 13 or trisomy 21

Data from Obstet Gynecol. 2012;120:1532-1534.
Noninvasive Prenatal Testing continued

microarray analysis. The recently completed NICHD Multicenter Prenatal Microarray Study indicated that 1.7% of routine low-risk cases (for such indications as advanced maternal age, abnormal FTRA, and parental concern) and 6% of cases referred for structural ultrasound abnormalities had a clinically significant copy number change that was not detected by routine cytogenetic analysis. As such, there is a much higher risk for these submicroscopic imbalances than there is for Down syndrome. NIPT still has a way to go before it can replace procedures such as CVS and amniocentesis.

Errol Norwitz, MD, PhD, is an unpaid member of the Advisory Board of Natera (San Carlos, CA); Brynn Levy, MSc (Med), PhD, is a consulting Laboratory Director of Natera.

References


MAIN POINTS

- Prenatal detection of chromosome abnormalities has been offered for more than 40 years; however, invasive testing also carries a risk for procedure-related miscarriage. For this reason, a number of noninvasive tests have been developed.

- Fetal DNA can be detected in as little as 10 µL of maternal plasma and serum, in amounts significantly higher than that available from nucleated blood cells extracted from a similar volume of whole blood. The approach of using cell-free fetal DNA instead of fetal cells provided a far easier, less labor intensive, and less time consuming way to work with fetal DNA derived from the maternal circulation, and this opened up new opportunities for noninvasive prenatal testing (NIPT).

- Noninvasive testing grew out of a desire to avoid direct contact with the growing fetus/placenta and concomitantly risk the health of the fetus. NIPT refers specifically to techniques that evaluate fetal cells or cfDNA in a blood sample drawn from the mother during pregnancy.

- The diagnostic scope of traditional prenatal cytogenetic analysis has recently been extended to include detection of microdeletions and microduplications using new genomic technologies such as microarray analysis.

- There are currently two primary next-generation sequencing approaches for gathering genetic data from cell-free DNA: massively parallel shotgun sequencing and targeted sequencing.

- A number of companies have been spearheading the effort to develop a reliable and accurate commercial NIPT for fetal aneuploidy detection. Although the precise technology used by each of these companies varies, they all rely on the same premise: that existing high-throughput approaches to DNA sequencing coupled with sophisticated data analysis will be able to detect abnormal amounts of chromosome-specific cfDNA in the maternal circulation in those pregnancies with fetal aneuploidy.

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