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<td><strong>Author(s)</strong></td>
<td>Chiu, RWK; Akolekar, R; Zheng, YWL; Leung, TY; Sun, H; Chan, KCA; Lun, FMF; Go, ATJI; Lau, ET; To, WWK; Leung, WC; Tang, RYK; Au-Yeung, SKC; Lam, H; Kung, YY; Zhang, X; Van Vugt, JMG; Minekawa, R; Tang, MHY; Wang, J; Oudejans, CBM; Lau, TK; Nicolaides, KH; Lo, YMD</td>
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<tr>
<td><strong>Citation</strong></td>
<td>BMJ (Clinical Research Ed.), 2011, v. 342, article no. c7401</td>
</tr>
<tr>
<td><strong>Issued Date</strong></td>
<td>2011</td>
</tr>
<tr>
<td><strong>URL</strong></td>
<td><a href="http://hdl.handle.net/10722/180701">http://hdl.handle.net/10722/180701</a></td>
</tr>
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Non-invasive prenatal assessment of trisomy 21 by multiplexed maternal plasma DNA sequencing: large scale validity study

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ABSTRACT

Objectives To validate the clinical efficacy and practical feasibility of massively parallel maternal plasma DNA sequencing to screen for fetal trisomy 21 among high risk pregnancies clinically indicated for amniocentesis or chorionic villus sampling.

Design Diagnostic accuracy validated against full karyotyping, using prospectively collected or archived maternal plasma samples.

Setting Prenatal diagnostic units in Hong Kong, United Kingdom, and the Netherlands.

Participants 753 pregnant women at high risk for fetal trisomy 21 who underwent definitive diagnosis by full karyotyping, of whom 86 had a fetus with trisomy 21.

Intervention Multiplexed massively parallel sequencing of DNA molecules in maternal plasma according to two protocols with different levels of sample throughput: 2-plex and 8-plex sequencing.

Main outcome measures Proportion of DNA molecules that originated from chromosome 21. A trisomy 21 fetus was diagnosed when the z score for the proportion of chromosome 21 DNA molecules was >3. Diagnostic sensitivity, specificity, positive predictive value, and negative predictive value were calculated for trisomy 21 detection.

Results Results were available from 753 pregnancies with the 8-plex sequencing protocol and from 314 pregnancies with the 2-plex protocol. The performance of the 2-plex protocol was superior to that of the 8-plex protocol. With the 2-plex protocol, trisomy 21 fetuses were detected at 100% sensitivity and 97.9% specificity, which resulted in a positive predictive value of 96.6% and negative predictive value of 96.6%.

Conclusion Multiplexed maternal plasma DNA sequencing analysis could be used to rule out fetal trisomy 21 among high risk pregnancies. If referrals for amniocentesis or chorionic villus sampling were based on the sequencing test results, about 98% of the invasive diagnostic procedures could be avoided.

INTRODUCTION

Trisomy 21, Down’s syndrome, occurs in 1 in 800 live births.1 Prenatal diagnosis of trisomy 21 requires invasive sampling of fetal genetic material through amniocentesis or chorionic villus sampling. However, these tests carry a risk of miscarriage of about 1%,2 and they are therefore reserved for pregnancies considered to be at high risk of fetal trisomy 21. The traditional method of identifying the high risk group has been increased maternal age, but screening by this method would require invasive testing in about 5% of pregnant women and identify only 30% of affected fetuses.1 In the past 20 years maternal age has been combined with ultrasonographic examination of the fetus and biochemical measurement of various proteins or hormones in the maternal circulation to improve identification of high risk pregnancies. This combined approach of screening can now identify more than 90% of affected fetuses, but there is still a need for invasive testing in 3–5% of the population.3–9

Cell-free DNA from the fetus has been found in the plasma of pregnant women, and this has been used successfully for non-invasive determination of the fetal sex and fetal RhD genotype in RhD negative women.10–13 The basis of these tests is the detection of fetal-specific DNA sequences in maternal plasma.14 The same approach of searching for fetal-specific nucleic acids, such as DNA methylation and mRNA
markers in maternal plasma, has been proposed for non-invasive detection of fetal aneuploidies.

The advantages and disadvantages of these approaches have been reviewed in detail. An alternative approach for non-invasive prenatal diagnosis of fetal trisomy 21 is to show the presence of an elevated amount of chromosome 21 sequences in maternal blood, because there are three rather than two copies of fetal chromosome 21. However, fetal DNA molecules amount to just 10–20% of the total DNA circulating in the maternal plasma, so any increment in the total amount (fetal and maternal) of chromosome 21 DNA molecules in the plasma of a trisomy 21 pregnancy would be substantially diluted by contributions from the mother.

The difficulty in measuring such a small increment in chromosome 21 DNA concentration has recently been overcome with the use of massively parallel genomic sequencing. This technique can identify and quantify millions of DNA fragments in biological samples in a span of days. Three cohort studies have shown the feasibility of using the technique to identify fetuses with trisomy 21 by analysis of maternal plasma DNA. The sample numbers studied were small because, typically, only a few samples could be analysed in each sequencing run.

The objective of this study is to validate the diagnostic performance and practical feasibility of massively parallel genomic sequencing for the non-invasive prenatal assessment of trisomy 21 in a large number of pregnancies that have undergone conventional screening and were clinically indicated for definitive testing by amniocentesis or chorionic villus sampling.

METHODS

The primary research question of the study was to evaluate if maternal plasma DNA sequencing could accurately confirm or exclude fetal trisomy 21 compared with full karyotyping among pregnancies clinically indicated for amniocentesis or chorionic villus sampling. A secondary goal of the study was to develop laboratory protocols to improve the throughput of massively parallel genomic sequencing for handling large sample numbers.

Recruitment criteria

We prospectively recruited pregnant women from eight obstetric units in Hong Kong, one unit from the Netherlands, and one prenatal diagnostic centre in the United Kingdom between October 2008 and May 2009. We also retrieved archival maternal plasma samples collected between October 2003 and September 2008 from trisomy 21 and non-trisomy 21 pregnancies matched for gestational ages in a ratio of approximately 1:5 from the participating sites in the Netherlands and UK. The inclusion criteria were singleton pregnancies with clinical indications for chorionic villus sampling or amniocentesis as per the existing obstetric practice of each recruitment unit. Women were excluded if full karyotyping results were not available.

We recorded the maternal ages at the expected time of delivery, gestational ages at blood sampling, and indications for invasive testing.

Sample processing

Peripheral venous blood samples (5–10 mL) were collected into tubes containing EDTA. A plasma sample would be accepted for analysis if it was collected before invasive obstetric procedures, was harvested within six hours of venepuncture, was at least 2 mL in volume, and was not haemolysed. The prospectively collected and retrieved archival plasma samples from the UK and the Netherlands were sent to Hong Kong in batches by overnight courier while kept frozen on dry ice. We performed all subsequent analyses prospectively at The Chinese University of Hong Kong.

Maternal plasma DNA sequencing

We extracted and sequenced DNA molecules from 2.0–4.8 mL of maternal plasma using a protocol similar to that reported previously but with the introduction of multiplexing. DNA molecules exist in maternal plasma as short fragments. The genomic identities of millions of DNA molecules per maternal plasma sample can be decoded in a sequencing run. The analytical goal of our test was to determine the proportion of sequenced plasma DNA molecules originating from chromosome 21 (percentage chromosome 21). This proportion is expected to be elevated in maternal plasma during a pregnancy with a trisomy 21 fetus. Multiplexing allows more than one plasma sample to be mixed and sequenced jointly in each discrete segment of a sequencing glass slide and thereby increases the number of samples that can be analysed in each sequencing run. We studied two levels of multiplexing, 2-plex and 8-plex, whereby DNA from two or eight maternal plasma samples were co-sequenced in each slide segment.

Briefly, a unique synthetic DNA “barcode” of six base pairs, referred to as an index, was introduced on to one end of each plasma DNA molecule. The index served as a signature for a sample, with one index used per maternal plasma sample. For example, for 8-plex sequencing, eight different indices were needed for each of the eight test or control samples that would be co-sequenced. We compiled the multiplexed sample mixtures by pooling eight or two maternal plasma DNA preparations for the 8-plex or 2-plex sequencing protocols.

We performed sequencing on the Genome Analyzer II (Illumina) for the 8-plex sample mixtures and on the Genome Analyzer IIx (Illumina) for the 2-plex sample mixtures. After the sequencing run, the actual DNA molecules that belonged to a specific sample could then be distinguished from those belonging to other samples by sorting the index sequences attached to the DNA molecules. We considered a sequencing result as valid and reportable only if the analysis of
the sample passed a set of quality control measures (listed in appendix 1).

Sequencing data analysis
We processed the sequencing data with the use of a bioinformatics algorithm based on our previously reported strategy (appendix 2 on bmj.com).24 In essence, the chromosomal origin of each sequenced “read” (the short fragments of DNA that are decoded) was identified by comparing with the reference human genome available from the Ensembl website (ftp://ftp.ensembl.org/pub/release-48/fasta/homo_sapiens/dna/). The percentage chromosome 21 within each maternal plasma sample was then calculated. The proportion of reads from chromosome Y (percentage chromosome Y) was similarly calculated and was used to estimate the fetal DNA concentrations (that is, the proportion of DNA molecules in the maternal plasma sample that originated from the fetus) in pregnancies with male fetuses. We derived a z score for chromosome 21 in a test sample by subtracting the mean percentage chromosome 21 of a reference set of euploid pregnancies (controls) from the percentage chromosome 21 of the test case and divided by the standard deviation of the value for percentage chromosome 21 among the reference sample set according to the equation:

\[
Z\text{ score for percentage chromosome 21 in test case} = \left( \frac{\text{percentage chromosome 21 in test case} - \text{mean percentage chromosome 21 in reference controls}}{\text{standard deviation of percentage chromosome 21 in reference controls}} \right)
\]

We used a z score of >3 (representing a percentage chromosome 21 value greater than that of the 99.9th centile of the reference sample set for a one tailed distribution) as the cut-off value to determine if the percentage chromosome 21 was increased and hence fetal trisomy 21 was present.

All members of the sequencing and bioinformatics teams were blinded to the karyotype information. Disease classification based on the sequencing results and z scores was generated automatically by the bioinformatics algorithm. We then determined the diagnostic performance of the sequencing test by comparing its results with those of full karyotyping of the amniotic fluid or chorionic villus sample.

Statistical analysis
We reported the observed diagnostic sensitivity and specificity values of the sequencing protocols based on the predefined cut-off point—namely, chromosome 21 z score of 3. We plotted the receiver operating characteristic (ROC) curves for the measurements of percentage chromosome 21 and determined the areas under the curves (AUC). ROC curve analysis was also performed to derive cut-off values for percentage chromosome Y to distinguish male and female fetuses. We measured the analytical imprecision of the sequencing protocols by calculating the coefficient of variation (SD/mean × 100%) of the percentage chromosome 21 values among the control samples.

Fig 1 | Recruitment of participants. Numbers in parentheses are the number of cases with sequencing results from the 2-plex protocol
Full karyotyping showed that there were 86 pregnancies with trisomy 21 fetuses (40 cases from the archival sample set) and 597 pregnancies with euploid fetuses (fig 1). The remainder comprised 40 fetuses with trisomy 18, 20 with trisomy 13, eight with Turner’s syndrome, and two sex chromosome mosaics, all of which were included as non-trisomy 21 cases in this study.

Among the sequenced samples, we randomly chose 96 pregnancies with euploid male fetuses as the reference control set for z score calculation. All 86 trisomy 21 cases, 82 reference controls, and 146 non-trisomy 21 cases were sequenced with the 2-plex protocol (fig 1).

Review of the clinical profile of the cases revealed there were three broad groups of clinical indications for referral for full karyotyping: (a) pregnancies with a risk higher than 1 in 300 for fetal trisomy 21 as estimated by conventional prenatal screening; (b) pregnancies with an intermediate risk (risk between 1 in 300 and 1 in 1000) for fetal trisomy 21; and (c) pregnancies with other risk indications, including a previous trisomy 21 pregnancy, ultrasound abnormalities, or risks for monogenic diseases. The numbers of pregnancies in the high, intermediate, and other risk groups were 582, 39, and 132, respectively (table 1). Among the high and intermediate risk groups, 570 (91.8%) pregnant women underwent combined screening in the first trimester while the remaining 51 (8.2%) women received other forms of prenatal screening, including the integrated test and second trimester maternal serum screening.

### Diagnostic performance of maternal plasma DNA sequencing

A mean of 0.3 million (SD 88 000) reads per sample from the 8-plex sequencing analysis and 2.3 million (SD 474 000) from 2-plex sequencing matched our bioinformatics criteria for calculation of percentage chromosome 21. As each sequencing glass slide can sequence only a finite number of DNA molecules, higher levels of multiplexing would lead to fewer DNA molecules to be sequenced per sample. Figure 2 shows the distribution of the chromosome 21 z scores among the controls, non-trisomy 21 group, and trisomy 21 group. Table 2 summarises the diagnostic performance of the sequencing test for detecting fetal trisomy 21.

With a chromosome 21 z score of 3 as the diagnostic cut-off point, the 2-plex protocol detected 100% of trisomy 21 fetuses at a 2.1% false positive rate—namely, 100% sensitivity and 97.9% specificity. This gave a positive predictive value of 96.6% and a negative predictive value of 100%.

The 8-plex protocol, however, detected 79.1% of the trisomy 21 fetuses at a false positive rate of 1.1%—that is, 79.1% sensitivity and 98.9% specificity. The positive predictive value was 91.9% and the negative predictive value

### Table 1 Characteristics of 753 pregnancies with maternal plasma DNA sequencing. Values are numbers (percentages) of subjects unless stated otherwise

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Reference controls (n=96)*</th>
<th>Non-trisomy 21 fetuses (n=571)</th>
<th>Trisomy 21 fetuses (n=86)</th>
<th>All (n=753)</th>
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<tbody>
<tr>
<td>Fetal sex:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>96 (100)</td>
<td>251 (44)</td>
<td>39 (45)</td>
<td>386 (51)</td>
</tr>
<tr>
<td>Female</td>
<td>0</td>
<td>318 (56)</td>
<td>47 (55)</td>
<td>365 (48)</td>
</tr>
<tr>
<td>Sex chromosome mosaic</td>
<td>0</td>
<td>2 (0.4)</td>
<td>0</td>
<td>2 (0.3)</td>
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<td>Maternal age at expected date of delivery (years):</td>
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<td></td>
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<td></td>
</tr>
<tr>
<td>&lt;35</td>
<td>42 (44)</td>
<td>255 (45)</td>
<td>22 (26)</td>
<td>319 (42)</td>
</tr>
<tr>
<td>≥35</td>
<td>54 (56)</td>
<td>316 (55)</td>
<td>64 (74)</td>
<td>434 (58)</td>
</tr>
<tr>
<td>Median age</td>
<td>35.7</td>
<td>35.0</td>
<td>38.0</td>
<td>35.4</td>
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<td>Gestational age at maternal blood sampling (weeks+days):</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤13+6</td>
<td>80 (83)</td>
<td>410 (72)</td>
<td>67 (78)</td>
<td>557 (74)</td>
</tr>
<tr>
<td>&gt;14+0 to 14+6</td>
<td>7 (7)</td>
<td>80 (14)</td>
<td>15 (17)</td>
<td>102 (14)</td>
</tr>
<tr>
<td>&gt;15+0</td>
<td>9 (9)</td>
<td>81 (14)</td>
<td>4 (5)</td>
<td>94 (12)</td>
</tr>
<tr>
<td>Median gestational age</td>
<td>12+5</td>
<td>13+1</td>
<td>13+0</td>
<td>13+1</td>
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<td>Means of invasive testing:</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chorionic villus sampling</td>
<td>78 (81)</td>
<td>462 (81)</td>
<td>81 (94)</td>
<td>621 (82)</td>
</tr>
<tr>
<td>Amniocentesis</td>
<td>18 (19)</td>
<td>109 (19)</td>
<td>5 (6)</td>
<td>132 (18)</td>
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<td>Risk grouping:</td>
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<tr>
<td>High</td>
<td>64 (67)</td>
<td>437 (77)</td>
<td>81 (94)</td>
<td>582 (77)</td>
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<tr>
<td>Intermediate</td>
<td>6 (6)</td>
<td>33 (6)</td>
<td>0</td>
<td>39 (5)</td>
</tr>
<tr>
<td>Other</td>
<td>26 (27)</td>
<td>101 (18)</td>
<td>5 (6)</td>
<td>132 (18)</td>
</tr>
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<td>Median fetal risk for trisomy 21:</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>In high risk group</td>
<td>1 in 50</td>
<td>1 in 51</td>
<td>1 in 2</td>
<td>1 in 63</td>
</tr>
<tr>
<td>In intermediate risk group</td>
<td>1 in 486</td>
<td>1 in 513</td>
<td>NA</td>
<td>1 in 502</td>
</tr>
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</table>

*Euploid male fetuses. NA=not applicable.
value was 96.9%. Eleven of the 18 false negative cases were from the archived samples. The distribution of the false negative and true positive cases among the prospectively collected and archived samples was not statistically significant (χ² test, \(P = 0.258\)). The median gestational ages for the true positive and false negative trisomy 21 cases from the 8-plex protocol were 13.0 weeks (interquartile range 12.3–13.5) and 12.9 weeks (12.3–13.5), respectively. The median risk level for trisomy 21 estimated by conventional screening for the true positive and false negative cases were 1 in 4 (interquartile range 1 in 5 to 1 in 2) and 1 in 5 (1 in 7 to 1 in 2), respectively. The true positive and false negative cases were not significantly different in terms of gestational ages (Mann-Whitney, \(P = 0.538\)) or risk levels (Mann-Whitney, \(P = 0.466\)). There were six false positive results in the 8-plex results and three in the 2-plex results. Because of the small number of false positive cases, no systematic features were identified. Profiles of these cases are shown in appendix 3 on bmj.com.

ROC curve analysis of the percentage chromosome 21 measurements for all test cases was done to estimate the diagnostic efficacy of the 2-plex and 8-plex sequencing protocols (fig 3). Area under the curve values for the 2-plex and 8-plex sequencing protocols were 1.00 (95% confidence interval 0.98 to 1.00) and 0.98 (0.97 to 0.99) respectively.

Utility of maternal plasma DNA sequencing for the prenatal assessment of trisomy 21

In this study we focused on pregnancies clinically indicated for invasive prenatal diagnostic testing because we required the availability of full karyotyping for comparison of the diagnostic performance of the sequencing test. However, this study design essentially limits the applicability of our data to pregnancies at increased risk for trisomy 21. To investigate if the sequencing test might be useful as a screening test for all pregnancies in general, we determined the post-test probabilities for women of different ages and hence with different prevalences of trisomy 21.\(^2\) The data are shown in table 3. For a 20 year old woman at 12 weeks of gestation, her pretest probability for trisomy 21 by maternal age alone was 1 in 1068. If maternal plasma DNA sequencing with the 2-plex protocol was done, the post-test probability for a positive test result was 1 in 23 while the post-test probability for a negative test result was 1 in infinity. With the 8-plex protocol, the post-test probabilities for a positive result and a negative result becomes 1 in 16 and 1 in 5082 respectively.

These data suggest that the sequencing test result, whether positive or negative, can substantially alter the probabilities for having a trisomy 21 fetus. The 2-
plex sequencing particularly highlights the key value of the sequencing test result for ruling out trisomy 21.

Factors affecting diagnostic performance

The diagnostic performance of the 2-plex protocol was superior to that of the 8-plex sequencing, with significantly higher z score values for the trisomy 21 cases (Mann-Whitney, P<0.001). The high z scores were due to a small standard deviation and, hence, better precision for estimating percentage chromosome 21. The coefficients of variation of percentage chromosome 21 values for the controls on each flow cell were calculated as a measure of precision (see appendix 4 on bmj.com). The mean coefficient of variation for measuring percentage chromosome 21 with the 8-plex protocol was 1.59% (SD 0.36%) and was higher than that for the 2-plex protocol, which was 0.66% (SD 0.25%) (t test, P<0.001).

The extent of increase in percentage chromosome 21 in maternal plasma for the trisomy 21 cases is governed by the fetal DNA concentration. We calculated the fetal DNA concentrations based on the percentage chromosome Y values among the 39 male fetuses with trisomy 21 and plotted them against the chromosome 21 z scores (shown in appendix 5 on bmj.com). The two variables are significantly correlated in both the 8-plex sequencing data (Pearson correlation coefficient 0.703, P<0.0001) and 2-plex data (Pearson correlation coefficient 0.462, P=0.003). The five pregnancies with male fetuses with trisomy 21 that we failed to detect using the 8-plex protocol (that is, with chromosome 21 z scores <3) had fetal DNA concentrations less than 10%.

Theoretically, at 10% fetal DNA concentration, percentage chromosome 21 is just 1.05 times higher in a plasma sample obtained from a trisomy 21 pregnancy than a euploid pregnancy.18 To detect this increase in percentage chromosome 21, the coefficient of variation of the test would need to be ≤0.83% (see appendix 6 on bmj.com). The mean coefficient of variation of the 8-plex protocol was greater than this value, whereas that of the 2-plex protocol was <0.83%. Hence, the 2-plex protocol performed better at detecting trisomy 21 among samples containing low fetal DNA concentrations.

Fetal sex determination by maternal plasma DNA sequencing

The presence of chromosome Y sequences among sequenced DNA molecules from maternal plasma should theoretically signify that the pregnancy involved a male fetus. However, we previously reported that the bioinformatics algorithm would incorrectly assign a small fraction of sequenced DNA reads to chromosome Y in pregnancies with female fetuses.24 The percentage chromosome Y values of all studied pregnancies are shown in appendix 7 on bmj.com. Optimal cut-off values for percentage chromosome Y to distinguish male and female fetuses were identified by ROC curve analysis to be 0.0114% and 0.0095% for the 8-plex and 2-plex protocols, respectively. Table 2 shows the estimated diagnostic performance for fetal sex determination. We then determined the fetal DNA concentrations of 314 pregnancies with euploid male fetuses based on the percentage chromosome Y values obtained from the 8-plex protocol. The median fetal DNA concentration was 15.2% (interquartile range 10.6%–19.1%), and the distribution is shown in fig 4. Among the pregnancies with euploid male fetuses, the median fetal DNA concentration for the archival and prospectively collected samples were 14.7% and 15.4%, respectively (Mann-Whitney, P=0.334).

<table>
<thead>
<tr>
<th>Maternal age (years)</th>
<th>Pretest probability*</th>
<th>8-plex sequencing</th>
<th>Post-test probability†</th>
<th>2-plex sequencing</th>
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<tr>
<td></td>
<td></td>
<td>Positive test result†</td>
<td>Negative test result</td>
<td>Positive test result†</td>
</tr>
<tr>
<td>20</td>
<td>1 in 1068</td>
<td>1 in 16</td>
<td>1 in 5082</td>
<td>1 in 23</td>
</tr>
<tr>
<td>25</td>
<td>1 in 946</td>
<td>1 in 14</td>
<td>1 in 4501</td>
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</tr>
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<td>30</td>
<td>1 in 626</td>
<td>1 in 10</td>
<td>1 in 2977</td>
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<tr>
<td>31</td>
<td>1 in 543</td>
<td>1 in 9</td>
<td>1 in 2582</td>
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<tr>
<td>32</td>
<td>1 in 461</td>
<td>1 in 7</td>
<td>1 in 2191</td>
<td>1 in 11</td>
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<td>33</td>
<td>1 in 383</td>
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<td>1 in 1820</td>
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<td>34</td>
<td>1 in 312</td>
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<td>35</td>
<td>1 in 293</td>
<td>1 in 4</td>
<td>1 in 1182</td>
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<td>36</td>
<td>1 in 196</td>
<td>1 in 4</td>
<td>1 in 930</td>
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<td>37</td>
<td>1 in 152</td>
<td>1 in 3</td>
<td>1 in 720</td>
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</tr>
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<td>38</td>
<td>1 in 117</td>
<td>1 in 3</td>
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<td>1 in 89</td>
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<td>1 in 38</td>
<td>1 in 2</td>
<td>1 in 177</td>
<td>1 in 2</td>
</tr>
</tbody>
</table>

*Pretest probabilities are based on prevalence of fetal trisomy 21 at 12th week of gestation.28
†A positive test result is a sample with a z score for percentage chromosome 21 >3.

Table 3 | Probabilities for a trisomy 21 fetus in women by age alone and according to result of maternal plasma DNA sequencing test
DISCUSSION

We show that fetal trisomy 21 could be detected in high risk pregnancies with high sensitivity and specificity by means of multiplexed sequencing of maternal plasma DNA. A key strength of the study is that the diagnostic performance was compared against full karyotyping, which is the gold standard for aneuploidy diagnosis. Another strength of this study is that the test performance was assessed based on a large number of trisomy 21 and unaffected pregnancies. In fact, this study reports the largest scale use of massively parallel genomic sequencing (so called next generation sequencing) for medical diagnostics to date and shows that it is practically feasible on a large scale.

Another strength of the study lies in the comparison of the diagnostic performance of sequencing protocols with two levels of sample throughput. The comparison allows us to deduce factors that are fundamental to the sensitive detection of fetal trisomy 21 and provide insight into ways for further improvements. Measurement of the percentage of plasma DNA that was from chromosome 21 was more precise with the use of the 2-plex protocol because seven times more plasma DNA molecules were analysed with this method than with the 8-plex protocol. All trisomy 21 fetuses, including those missed by the 8-plex protocol, were detected with the 2-plex protocol. In short, the diagnostic sensitivity of the sequencing approach improves with an increase in the number of plasma DNA molecules analysed per sample.18

There are some weaknesses in the study design. Our study was based on high risk pregnancies—those clinically indicated for amniocentesis or chorionic villus sampling—which ensured that full karyotyping results were available from each pregnancy. To investigate if the sequencing test might be useful for prenatal assessment of fetal trisomy 21 for all pregnancies in general, we determined the post-test probabilities for having a trisomy 21 fetus in women of different ages, and hence with different pretest risks. The data suggest that the sequencing test result can substantially alter the probabilities for having a trisomy 21 fetus even in women with low pretest risk.

Another potential weakness of the study was that an archived sample set was included. However, both the prospectively collected and archived maternal plasma samples were processed using the same protocols and analysed prospectively. There were also no significant differences in the fetal DNA concentrations and test performances between the two sample groups. In this study 5.6% of the collected maternal plasma samples were of compromised quality. Reportable sequencing results were not achieved for 11 samples. All these samples were identified by the predefined quality control steps. In routine practice, laboratory reports would not be issued for such samples, and another blood specimen would be requested.

We found other aneuploidies besides trisomy 21 among the recruited pregnant women. For this study, we focused on the diagnostic performance of the sequencing approach for fetal trisomy 21 because our previous data showed that the measurements of the proportion of DNA molecules from chromosomes 18 and 13 were much less precise.24,25 More research is required to develop protocols to improve the precision for measuring amount of DNA molecules from chromosomes 18 and 13.26

Our data reveal that the main value of the maternal plasma DNA sequencing test is to rule out fetal trisomy 21. Hence, with the current diagnostic performance, it is more suitable as a screening test to stratify pregnancies whose risk for trisomy 21 warrants the consideration of amniocentesis or chorionic villus sampling. In this study, we performed the sequencing test after a pregnancy was deemed to be at increased risk for trisomy 21 according to the current prenatal screening programmes. The false positive rates of the current screening programmes are about 5%, and all of these pregnant women are offered the option of invasive testing. However, if we took into consideration the results of the sequencing test, trisomy 21 could be ruled out in 98% of those pregnancies. This would leave just 0.1% (that is, 5%×(100%−98%)×100%) of all pregnant women needing referrals for amniocentesis or chorionic villus sampling. Most of our studied pregnancies were in the first trimester, which suggests that it is possible to implement the test even in early pregnancy as a second tier screening test after the first trimester combined test, which is already in use in many parts of the world.

On the other hand, our post hoc analysis shows that the sequencing test result may alter clinical decisions even in women at low risk for trisomy 21. Hence, there is the potential to apply the sequencing test as a first tier screening test. However, massively parallel genomic sequencing is currently expensive. To determine its suitability as a first tier screening test would require formal investigation of both its diagnostic performance and cost benefit. We used the sequencing test in this study for detection of fetal trisomy 21, but, with some improvements in bioinformatics, it could be used to detect other aneuploidies, such as trisomy 18 and trisomy 13.29 Furthermore, our group has recently shown that when more sophisticated sequencing protocols and bioinformatics algorithms are applied to the analysis of maternal plasma DNA, it may be possible to
WHAT IS ALREADY KNOWN ON THIS TOPIC
Non-invasive prenatal detection of fetal trisomy 21 is achievable by massively parallel sequencing of maternal plasma DNA. Its diagnostic performance and practical feasibility in the clinical setting has not been tested on a large scale.

WHAT THIS STUDY ADDS
Among high risk pregnancies clinically indicated for invasive prenatal diagnosis, non-invasive detection of fetal trisomy 21 can be achieved with the use of multiplexed massively parallel sequencing of maternal plasma DNA at 100% sensitivity and 97.9% specificity, giving a 96.6% positive predictive value and 100% negative predictive value.

The sequencing test could be used to rule out trisomy 21 among high risk pregnancies before proceeding to invasive diagnostic testing to reduce the number of cases requiring amniocentesis or chorionic villus sampling.

perform a genetic and mutational scan across the whole genome of the fetus in a non-invasive manner.30 Thus, it is likely that maternal plasma DNA sequencing will play an increasingly important role in the future developments of prenatal screening and diagnosis.

We thank members of the Lo laboratory for technical assistance in performing the sequencing runs.

Contributors: RWKC and YMDL designed and initiated the study, coordinated and managed the research process, interpreted the results and wrote the first draft of the report. RA, ATJIG, ETL, JMGvV, and RM contributed to the implementation of the clinical recruitment protocols and management of clinical data. TYL, MHYT, CBMO, TKL, and KHN contributed to the design of the clinical recruitment protocols, coordination of the recruitment of patients and management of clinical data. YWZ, KCAC, and FMFL contributed to the development of the laboratory protocols and monitoring of the laboratory analyses. HS and KCAC contributed to the development of the bioinformatics analysis protocols. WWK, WKT, WLYK, SKCA-Y, HL, and YYK contributed to the recruitment, collection and management of clinical data. XZ and JW contributed to the laboratory analysis. All authors provided critical comments on the drafts of the manuscript and approved the final version.

Competing interests: All authors have completed the Unified Competing Interest form available on request from the corresponding author and declare that (1) RWKC, HS, ETL, MHYT, TKL, and YMDL had support from the University Grants Committee of the Government of the Hong Kong Special Administrative Region, China, under the Areas of Excellence Scheme (AoE/M-04/06) for the submitted work; RWKC and YMDL had support from Sequenom for the submitted work; YMDL has support from an endowed chair from the Li Ka Shing Foundation. (2) YMDL is a consultant to, and holds equities in, Sequenom. RWKC and YMDL have received travel grants from Life Technologies and Illumina. RWKC, WLYZ, HS, KCAC, FMFL, and YMDL have received travel grants from University Grants Committee of the Government of the Hong Kong Special Administrative Region, China, under the Areas of Excellence Scheme (AoE/M-04/06). (3) The authors’ spouses, partners, and children have no financial relationships that may be relevant to the submitted work. (4) RWKC, WLYZ, KCAC, FMFL, and YMDL have filed patent applications on the detection of fetal nucleic acids in maternal plasma for non-invasive prenatal diagnosis. Part of this patent portfolio has been licensed to Sequenom and the Institut Jacques Boyd. The funders and sponsors have no role in the study design and the collection, analysis, and interpretation of data and the writing of the article and the decision to submit it for publication. The researchers are independent of the funders and sponsors.

Ethical approval: Approvals were obtained from the institutional review boards of each recruitment site: Joint Chinese University of Hong Kong-New Territories East Cluster Clinical Research Ethics Committee, Joint Institutional Review Board of the University of Hong Kong-Hospital Authority Hong Kong West Cluster, Clinical and Research Ethics Committees of the Hospital Authority in the Kowloon Central/Kowloon East, Kowloon West, Hong Kong East, New Territories West Clusters, King’s College Hospital Ethics Committee, and Ethics Committee of the VU University Medical Center. All participants gave informed written consent.

Data sharing: No additional data available.

29 Fan HC, Quake SR. Sensitivity of noninvasive prenatal detection of fetal aneuploidy from maternal plasma using shotgun sequencing is limited only by counting statistics. PLoS One 2010;5:e10439.

Accepted: 14 December 2010