Quantitative analysis of DNA levels in maternal plasma in normal and Down syndrome pregnancies

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Abstract

Background: We investigated fetal and total DNA levels in maternal plasma in patients bearing fetuses affected with Down syndrome in comparison to controls carrying fetuses with normal karyotype.

Methods: DNA levels in maternal plasma were measured using real-time quantitative PCR using SRY and β-globin genes as markers. Twenty-one pregnant women with a singleton fetus at a gestational age ranging from 15 to 19 weeks recruited before amniocentesis (carried out for reasons including material serum screening and advanced maternal age), and 16 pregnant women bearing fetuses affected with Down syndrome between 17 to 22 weeks of gestation were involved in the study.

Results: The specificity of the system reaches 100% (no Y signal was detected in 14 women pregnant with female fetuses) and the sensitivity 91.7% (SRY amplification in 22 of 24 examined samples). The median fetal DNA levels in women carrying Down syndrome (n=11) and the controls (n=13) were 23.3 (range 0–58.5) genome-equivalents/ml and 24.5 (range 0–47.5) genome-equivalents/ml of maternal plasma, respectively (P = 0.62). The total median DNA levels in pregnancies with Down syndrome and the controls were 10165 (range 615–65000) genome-equivalents/ml and 7330 (range 1300–36750) genome-equivalents/ml, respectively (P = 0.32). The fetal DNA proportion in maternal plasma was 0%-6 % (mean 0.8%) in women carrying Down syndrome and 0%-2.6 % (mean 0.7 %) in the controls, respectively (P=0.86).

Conclusions: Our study revealed no difference in fetal DNA levels and fetal DNA: maternal DNA ratio between the patients carrying Down syndrome fetuses and the controls.
Background
Current experimental non-invasive methods for the prenatal diagnosis of fetal genetic characteristics use free extracellular fetal DNA and fetal cells isolated from maternal peripheral blood. Free extracellular fetal DNA which can be detected by polymerase chain reaction (PCR) in the serum or plasma of pregnant women [1] has been previously successfully used for the prenatal diagnosis of fetal sex and rhesus D status [2–5]. A novel approach is to monitor elevations in the fetal DNA levels in pregnancies with preeclampsia [7,8], aneuploid fetuses [9,10] and preterm labor [11] using the real-time quantitative PCR technology (RQ-PCR) [6]. The study of Zhong et al. also showed that in preeclampsia, the amounts of both circulatory fetal and maternal DNA were increased [8].

In this study, we investigated the fetal and total DNA levels in maternal plasma in patients bearing fetuses affected with Down syndrome in comparison to the controls carrying fetuses with normal karyotype. The detection of Y DNA was used in male fetuses because a generic fetal DNA marker is not yet available.

Methods
Twenty-one pregnant women with a singleton fetus at a gestational age ranging from 15 - 19 (Φ 17.) weeks with an increased risk for an aneuploid fetus who were undergoing an invasive prenatal diagnostic procedure (amniocentesis), and 16 pregnant women bearing fetuses affected with Down syndrome between 17 - 22 (Φ 20.) weeks of gestation were recruited for this study. Local Ethics Committee approval and informed consent was obtained for all patients in the study.

DNA extraction from plasma samples
Five ml of maternal peripheral blood from Down syndrome and euploid cases was collected into EDTA containing tubes and processed within a few hours (maximally 24 h). In details, blood samples were centrifuged firstly at 1200 g for 10 min., than plasma samples were recentrifuged again and the supernatants were collected and stored at -80°C until further processing. Down syndrome and euploid case samples were matched for gestational age ranging from 15 - 19 (Φ 17.) weeks with 5,1 (range 1–8) months before DNA extraction and RQ-PCR analyses.

DNA was extracted from 400 µl plasma using QIAamp DNA Blood Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. To minimize the risk of contamination, DNA was isolated in laminar airflow and aerosol resistant tips were used. DNA was eluted in 50 µl Buffer AE and 2,5 µl were used as a template for the RQ-PCR reaction.

Real-time quantitative PCR Analysis
The real time quantitative PCR analysis was performed using ABI PRISM 7700 Sequence Detection System (Applied Biosystem, Branchburg, New Jersey, USA). The β-globin TaqMan system consisted of two primers β-globin-354 (forward), 5'-GTG CAC CTA CCT GAG GAG GCA GA3'; β-globin-455 (reverse), 5'-CCT TGA TAC CAA CCT GCC CAG-3' and a dual-labelled fluorescent TaqMan probe β-globin-402T, 5'(FAM) AAG GTG AAC GTG GAT GAA GTT GGT GG (TAMRA)-3' [6]. The SRY TaqMan system consisted of SRY-109 (forward) primer, 5'-TGG CGA TTA AGT AGT ACC CTG ACA ATG TAT T-3' and a probe SRY-142T, 5'(FAM) AAG GTG AAC GTG GAT GAA GTT GGT GG (TAMRA)-3' [6]. Sequence data were obtained from the GeneBank Sequence database U01317 (β-globin gene) and L08063 (SRY gene) as was previously described by Lo et al. [6]. TaqMan amplification reactions were set up in a reaction volume of 25 µl using the TaqMan Universal PCR Master Mix (Applied Biosystems, Branchburg, New Jersey, USA). DNA amplifications were carried out in 8-well reaction optical tubes/stripes (Applied Biosystem, Branchburg, New Jersey, USA). The TaqMan PCR conditions were used as described in TaqMan guidelines using 40 cycles of 95°C for 15 s and 60°C for 1 min. with 2-min preincubation at 50°C required for optimal AmpErase UNG activity and 10-min preincubation at 95°C required for activation of AmpliTaq Gold DNA polymerase. Each sample was analysed in triplicate.

The calibration curve was run in parallel with each analysis. The concentration of the of SRY and β-globin were expressed in genome-equivalents per millilitre of maternal plasma.

Statistics
Two-tailed t-test was performed using the SPSS software package for Windows. A P value of < 0.05 was taken as statistically significant.

Results
Figure 1 and Figure 2 demonstrate standard curves for SRY and β-globin genes plotting the threshold cycle (Ct) against known concentrations of serially diluted mixed DNA. Target quantity of unknowns was interpolated from given the Ct of any unknown.

In our setting the sensitivity of the RQ-PCR system was 91.7 % as SRY specific PCR amplicons were detected in 22 out of 24 studied samples and the specificity 100 % since
no Y chromosome positive signals were detected in 14 women currently pregnant with a female fetus.

The median fetal DNA levels in maternal plasma obtained from women carrying Down syndrome (n = 11) and the controls carrying fetuses with normal karyotype (n = 13) were 23.3 (range 0–58.5) genome-equivalents/ml and 24.5 (range 0–47.5) genome-equivalents/ml. The difference between the two groups did not reach statistical significance (P= 0.62), (Figure 3).

The total median DNA levels in patients bearing fetuses affected with Down syndrome (n = 16) and the controls carrying fetuses with normal karyotype (n = 21) were 10165 (range 615 – 65000) genome-equivalents/ml and 7330 (range 1300 – 36750) genome-equivalents/ml of maternal plasma, respectively (P = 0.32), (Figure 4).

Discussion

Recently, Lo et al. [9] reported findings of very high concentrations of circulating cell-free fetal DNA in maternal plasma in a proportion of pregnancies involving trisomy 21 fetuses. The median cell-free fetal DNA in women carrying trisomy 21 fetuses was found to be 2.96-fold and 1.97-fold higher than that of women carrying euploid fetuses in prospective studies done on samples collected from two centres. Consequently, Zhong et al. [10] confirm these results by showing elevated levels of fetal DNA in pregnancies with trisomy 21 fetuses (mean 185.6 genome-equivalents/ml) when compared to pregnancies with normal male fetuses (mean 83.1 genome-equivalents/ml).
Our results showed that fetal DNA levels in maternal plasma were not significantly elevated in pregnancies with Down syndrome fetuses when comparing to the controls carrying fetuses with normal karyotype. The median fetal DNA levels in maternal plasma obtained from women carrying Down syndrome \( (n=11) \) and the controls carrying fetuses with normal karyotype \( (n=13) \) were 23.3 (range 0–58.5) genome-equivalents/ml and 24.5 (range 0–47.5) genome-equivalents/ml, respectively \( (P=0.62) \). Plasma samples from women carrying euploid fetuses were recruited before amniocentesis for indications including maternal serum screening and advanced maternal age and compared with samples from women carrying Down syndrome fetuses sampling immediately as soon as possible after karyotype determination. Our data might confirm recently published results of Ohashi et al. independent study who observed no significant differences in the levels of fetal DNA between pregnancies with fetuses of normal karyotype and those with Down and Edwards syndrome [12]. Several studies reported that plasma DNA concentration could be affected by different sample processing protocols [13,14]. The examination of only circulatory fetal DNA, any of initial processing steps concerning a single centrifugation of the blood sample or use of a discontinuous Percoll density gradient, as well as plasma fractions subsequently cleared further by high-speed centrifugation or filtration appear not to make significant difference in the concentration of fetal DNA [13,14]. However in case of maternal plasma, DNA concentrations may vary in various studies due to the level of clearing the plasma of any maternal cellular remnants. By using sample processing protocol concerning the double centrifugation of blood samples at 1200 g we found out comparable fetal DNA levels as those reported by Lo et al. who separated the plasma using double centrifugation at 3000 g [6]. Differences in total DNA levels and fetal DNA:maternal DNA ratio in maternal plasma between our results and those reported by Lo et al. [6] might be caused by the use of lower-speed and lower-time centrifugation protocol we used for plasma preparation in our study.

**Conclusions**

Our study revealed no difference in fetal DNA levels and fetal DNA:maternal DNA ratio between the patients carrying Down syndrome and the controls carrying fetuses with normal karyotype.

**Competing interests**

None declared.

**Authors' contributions**

IH designed the study and drafted the manuscript. BH, DH, SV carried out plasma sampling, DNA extraction and RQ-PCR studies. PC, KN, JK, DS, JD collected biological samples and reported clinical data. OC and JV participated in the design of the study.

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