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Application of an individualized nomogram in first-trimester screening for trisomy 21

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KEYWORDS: facial profile; first-trimester screening; machine learning; nomogram; trisomy 21; ultrasonographic markers

CONTRIBUTION

What are the novel findings of this work?

We present an individualized nomogram, incorporating fetal nuchal translucency thickness (NT) and a series of ultrasonographic facial profile markers, which can potentially be utilized as a convenient and effective tool for screening for trisomy 21 in the first trimester of pregnancy.

What are the clinical implications of this work?

At present, the most widely used ultrasonographic marker for trisomy 21 is fetal NT at 11 to 13 + 6 weeks' gestation. The nomogram developed in this study can improve the ultrasonographic detection rate of trisomy 21 and can be used for screening during the routine prenatal ultrasound examination in early pregnancy.

ABSTRACT

Objectives To develop and validate a nomogram based on fetal nuchal translucency thickness (NT) and ultrasonographic facial markers for screening for trisomy 21 in the first trimester of pregnancy.

Methods This was a retrospective case–control study using stored two-dimensional midsagittal fetal profile images captured at 11 + 0 to 13 + 6 weeks' gestation in

singleton pregnancies. We included images from 302 trisomy-21 pregnancies and 322 euploid pregnancies. Cases were divided into a training set (200 euploid + 200 with trisomy 21) and a validation set (122 euploid + 102 with trisomy 21) at a ratio of approximately 2:1. For each, the maternal age, gestational age, fetal NT and karyotype were noted, and 12 ultrasonographic fetal facial markers were measured. The least absolute shrinkage and selection operator (LASSO) method and multivariable analysis were used to select automatically the discriminative markers. Logistic regression was used to develop a LASSO model, based on the selected markers, to screen for trisomy 21 in the first trimester of pregnancy. Furthermore, 60 of the 624 images were selected randomly as a retest set to evaluate the model's robustness. The predictive performance of screening for trisomy 21 of a model based on fetal NT and maternal age and of the LASSO model was assessed using the area under the receiver-operating-characteristics curve (AUC). A nomogram was developed as an individualized tool to predict patient-specific probability for trisomy 21, which is a more visual presentation of the LASSO model. The performance of the nomogram was assessed using the C-index and calibration curve.

Results Into the LASSO model were incorporated eight markers, including fetal NT, prenatal-thickness-to-nasal-bone-length ratio, facial profile line, frontomaxillary facial

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angle, frontonasal facial angle, mandibulomaxillary facial angle, maxilla-nasion-mandible angle and d2 (distance between the anterior edge of the prefrontal skin and the mandibulomaxillary line) (all $P < 0.05$). The AUCs of the LASSO model for screening for trisomy 21 were 0.983 (95% CI, 0.971–0.994) in the training set and 0.979 (95% CI, 0.966–0.993) in the validation set, and these were higher than the AUCs of all eight individual ultrasonographic markers included in the model. The AUC of the LASSO model in the retest set was 0.997 (95% CI, 0.990–1.000), indicating good robustness of the LASSO model. The AUC of the LASSO model was significantly higher than that of the model based on fetal NT and maternal age in both training and validation sets ($P < 0.001$ for both). The nomogram of the LASSO model showed good discrimination of trisomy 21, with C-indices of 0.983 in the training set and 0.981 in the validation set.

Conclusions We present an individualized nomogram which incorporates fetal NT and a series of ultrasonographic facial profile markers selected by the LASSO method and multivariable analysis. This nomogram can potentially be utilized as a convenient and effective tool in screening for trisomy 21 in the first trimester of pregnancy. © 2020 The Authors. *Ultrasound in Obstetrics & Gynecology* published by John Wiley & Sons Ltd on behalf of International Society of Ultrasound in Obstetrics and Gynecology.

INTRODUCTION

Trisomy 21 is the most common chromosomal abnormality. Growth delay and intellectual disability are common in children with Down syndrome¹. Screening for trisomy 21 in the first trimester of pregnancy is of critical importance, in order to provide women with affected fetuses reproductive choices at the earliest opportunity.

The 2013 ISUOG Practice Guidelines² state that fetal nuchal translucency thickness (NT) and absence of the nasal bone can be used to screen for chromosomal abnormalities in the first trimester of pregnancy. Furthermore, it has been observed that some trisomy-21 fetuses have abnormal facial features, such as a flat face and nasal-bridge collapse. However, the evaluation of these observations is subjective. Considerable effort has been devoted to transforming such abnormal facial features into objective ultrasonographic facial markers. There are essentially three main categories of quantitative fetal facial marker: (1) length and ratio, including prenasal thickness, nasal bone length, prenasal-thickness-to-nasal-bone-length ratio and prefrontal space ratio^{3,4}; (2) angles, including frontomaxillary facial angle, maxilla-nasion-mandible angle, frontonasal facial angle and mandibulomaxillary facial angle^{5–8}; and (3) facial profile line⁴. Some of these markers, such as prenasal-thickness-to-nasal-bone-length ratio, frontomaxillary facial angle, prefrontal space ratio and facial profile line have been assessed individually for the

prediction of trisomy 21 in the first trimester^{3–5}, whilst the others have only been utilized in the second and third trimesters. No study has reported on the performance of screening for trisomy 21 based on a combination of these quantitative markers in the first trimester of pregnancy.

Machine-learning methodology is a branch of research on artificial intelligence^{9,10}. Machine-learning algorithms build a mathematical model based on sample data (known as ‘training data’) to make predictions without being programmed explicitly to perform the task¹¹. In recent years, the machine-learning method has been utilized widely for the identification of tumors from ultrasonographic images¹². This method has yet to be applied in prenatal screening¹³. The aim, therefore, of this study was to develop and validate a nomogram that incorporates fetal NT and ultrasonographic fetal facial markers for screening for trisomy 21 in the first trimester of pregnancy, in cases in which the fetal face could be visualized adequately.

METHODS

The process of development of the nomogram for screening for trisomy 21 in the first trimester of pregnancy is illustrated in Figure 1.

Patients

This was a retrospective case–control study using stored fetal ultrasound images obtained at the Department of Ultrasound at the Beijing Obstetrics & Gynecology Hospital, Maternal & Child Health Centre, Capital Medical University, Beijing, China, between January 2009 and February 2019. We selected from our database two-dimensional images in the midsagittal plane of the fetal face from singleton pregnancies at 11 to 13 + 6 weeks’ gestation. The criteria for selection were as follows: (a) fetal karyotype confirmed by chorionic villus sampling or amniocentesis; (b) fetal nasal bone, frontal bone, skin over the nasal bone and the anterior edges of maxilla and mandible clearly visible on the image.

During the study period, 408 cases of trisomy 21 were confirmed by chorionic villus sampling or amniocentesis. Of these, 83 (20%) cases were excluded because of poor image quality, and a further 23 (6%) were excluded due to absence of the nasal bone, as many of the ultrasonographic markers rely on its presence. Thus, images from 302 trisomy-21 cases were retrieved for inclusion in the study. These were matched with images from 322 euploid fetuses that were selected randomly using the same criteria as for the trisomy-21 cases. Images were divided into a training set (200 euploid + 200 with trisomy 21) and a validation set (122 euploid + 102 with trisomy 21) at a ratio of approximately 2:1 (Figure 2).

Markers

For each of the 624 fetuses, the following data were collected: maternal age, gestational age, fetal karyotype

and fetal NT, and the following 12 facial ultrasonographic markers were measured: prenasal thickness, nasal bone length, prenasal-thickness-to-nasal-bone-length ratio, d1 (distance between the anterior edge of the frontal bone and the anterior edge of the prefrontal skin), d2 (distance between the anterior edge of the prefrontal skin (at the same point as the d1 measurement) and the mandibulomaxillary line), prefrontal space ratio (calculated as $d2/d1$), frontomaxillary facial angle, mandibulomaxillary facial angle, maxilla-nasion-mandible angle, frontonasal facial angle, facial profile line and the F distance (the largest perpendicular distance between the facial profile line and the outermost part of the frontal bone). These facial markers were measured manually, using retrieved midsagittal images which had been acquired for the measurement of fetal NT, by a single sonographer (Y.S.), using ITK-snap software (Version 3.6.0, <http://www.itksnap.org>). The sonographer was blinded to the karyotypic results.

Ultrasound images demonstrating the facial ultrasonographic marker measurements in euploid and trisomy-21 fetuses are given in Appendix S1. Prenasal thickness was the shortest distance between the anterior edge of the

lowest part of the frontal bone and the anterior line of skin. Nasal bone length was measured between the two ends of the ossification line (Figure S1 in Appendix S1)³. For d1 and d2, the mandibulomaxillary line was drawn between the anterior edge of the mandible and the anterior edge of the maxilla, and extended in the direction of the frontal bone. Both d1 and d2 were measured approximately parallel to the inferior edge of the maxilla. d1 was the distance between the anterior edge of the frontal bone and the anterior edge of the prefrontal skin and d2 was the distance between the anterior edge of the prefrontal skin at the same point as the d1 measurement and the mandibulomaxillary line. When the mandibulomaxillary line crossed behind the anterior edge of the prefrontal skin, d1 was measured between the frontal bone and the prefrontal skin, while d2 was measured between the mandibulomaxillary line and the prefrontal skin and multiplied by -1 . The prefrontal space ratio was calculated as $d2/d1$ (Figure S2 in Appendix S1)⁴. The frontomaxillary facial angle was measured between a line along the upper surface of the palate and a line from the upper-anterior corner of the maxilla to the external surface of the frontal bone or the external surface of an echogenic line under the

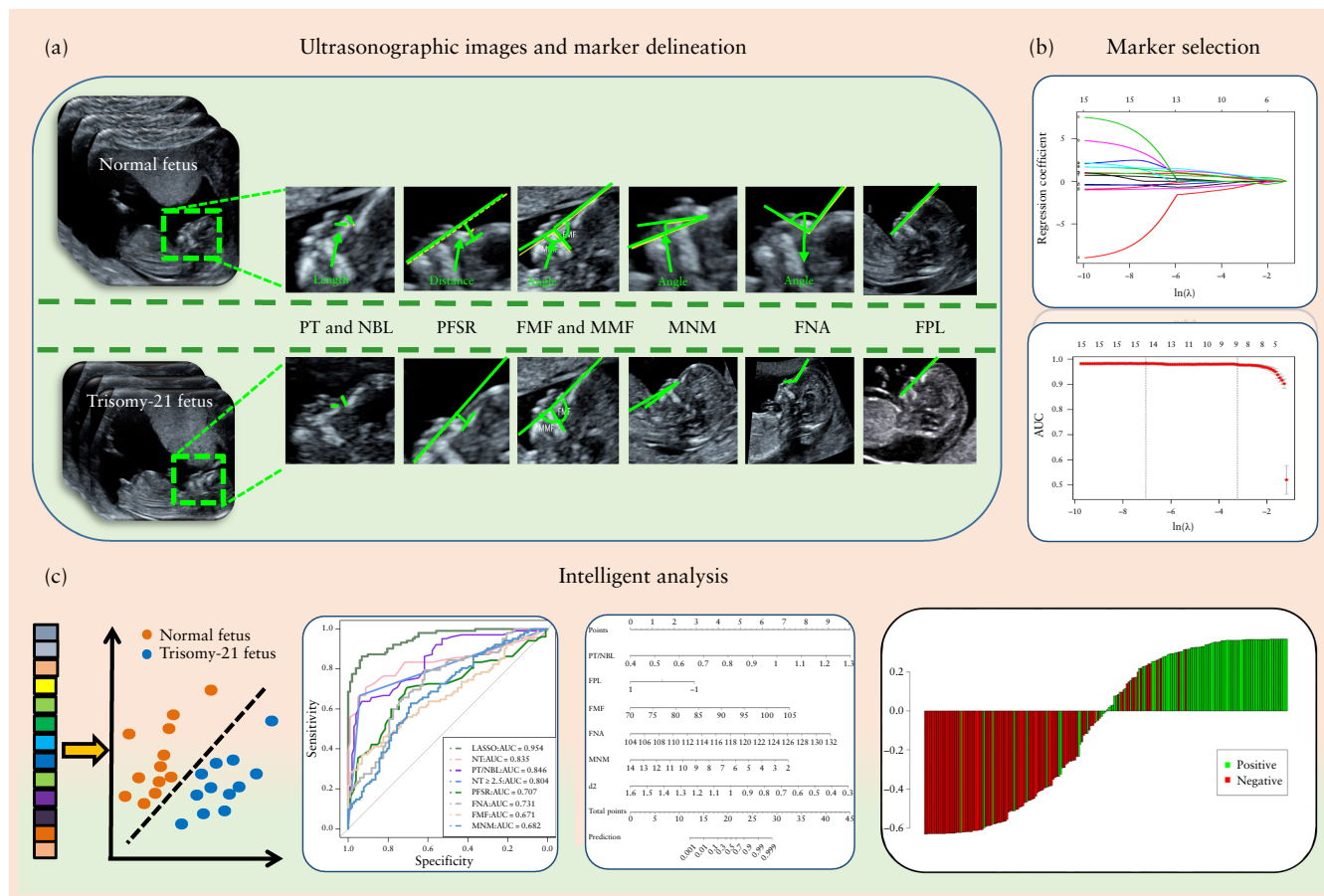


Figure 1 Flowchart summarizing development of nomogram for screening for trisomy 21 in first trimester of pregnancy. (a) Marker delineation of euploid fetuses and fetuses with trisomy 21. (b) Marker selection using least absolute shrinkage and selection operator (LASSO) method. (c) Development of LASSO model and nomogram. AUC, area under the receiver-operating-characteristics curve; d2, distance between anterior edge of prefrontal skin and mandibulomaxillary line; FMF, frontomaxillary facial angle; FNA, frontonasal facial angle; FPL, facial profile line; MMF, mandibulomaxillary facial angle; MNM, maxilla-nasion-mandible angle; NBL, nasal bone length; NT, nuchal translucency thickness; PFSR, prefrontal space ratio; PT, prenasal thickness.

skin, representing the metopic suture that remained open (Figure S3 in Appendix S1)⁵. The mandibulomaxillary facial angle was measured between a line along the upper surface of the palate (as for the frontomaxillary facial angle) and a line from the upper-anterior corner of the maxilla to the upper anterior corner of the mandible (Figure S3 in Appendix S1)⁸. The nasion was the apex of the maxilla-nasion-mandible angle; this angle was measured between a line from the midpoint of the anterior edge of the maxilla to the midpoint of the nasion and a line from the nasion to the midpoint of the anterior edge of the mandible. The nasion is the intersection of the frontal and nasal bones. When there was no intersection between nasal and frontal bones, the nasion was defined as the intersection between the tangential line of the nasal bone and the tangential line of the lowest part of the frontal bone (Figure S4 in Appendix S1)¹⁴. The frontonasal facial angle was measured between a line

along the outer profile of the nasal bone and a line traversing the upper corner of the external profile of the nasal bone, extending to the external surface of the forehead (Figure S5 in Appendix S1)¹⁵. The facial profile line was drawn from the nasion to the midpoint of the anterior edge of the mandible, and was classified as 'negative', 'zero' or 'positive', which indicated that the facial profile line passed the frontal bone anteriorly, longitudinally or posteriorly, respectively. The F distance (the greatest perpendicular distance between the facial profile line and the outermost part of the frontal bone) was measured when the facial profile line was positive (Figure S6 in Appendix S1)⁴.

In total, we evaluated 15 markers, including maternal age, fetal NT, fetal NT ≥ 2.5 mm and the 12 facial ultrasonographic markers. We selected the most useful predictive markers for trisomy 21 from these markers using the least absolute shrinkage and selection operator

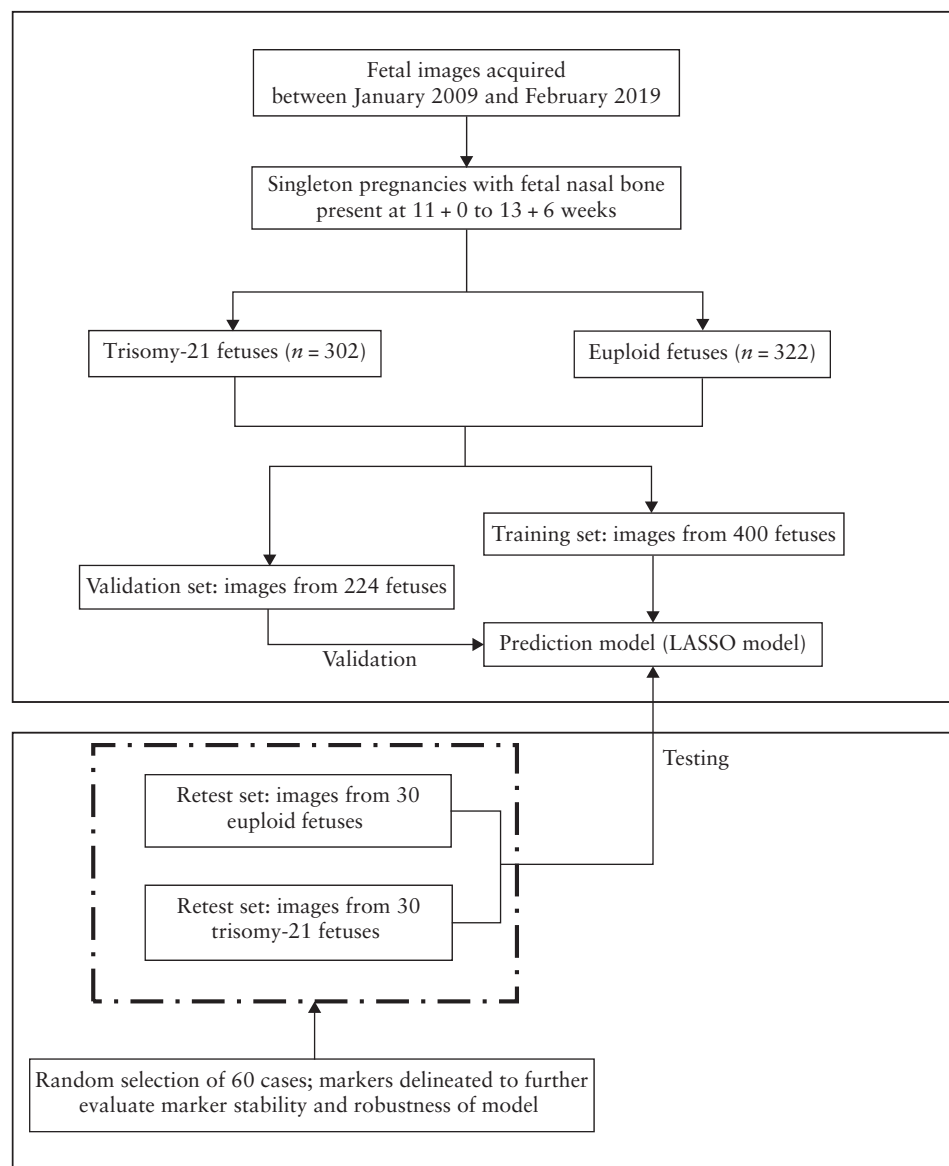


Figure 2 Flowchart summarizing development and verification of prediction model (least absolute shrinkage and selection operator (LASSO) model).

(LASSO) method and multivariable analysis. The LASSO method uses the penalty parameter $\ln(\lambda)$ to shrink automatically regression coefficients of these 15 markers; the greater the value of $\ln(\lambda)$, the more a marker's coefficient will shrink to zero (Figure 3). We excluded any marker whose coefficient was shrunk to approximately zero, and selected for inclusion the remaining markers. Multivariable analysis was used on the markers selected by the LASSO method, and the markers with $P < 0.05$ were selected for development of the prediction model.

Prediction (LASSO) model

In the training set, a prediction model (LASSO model) for screening for trisomy 21 in the first trimester of pregnancy was developed using logistic regression based on the selected markers. In order to validate the predictive performance of the LASSO model, the area under the receiver-operating-characteristics (ROC) curve (AUC) of the LASSO model was calculated for both the training and the validation sets.

In order to examine whether the screening performance of the LASSO model was operator-dependent, 60 fetuses (30 euploid and 30 with trisomy 21) were selected randomly from our dataset of 624, to act as a retest set. Delineation of the ultrasonographic markers in these 60 retest cases was performed by another sonographer (X.L.), who was blinded to the observations of the first sonographer (Y.S.) and to the karyotype results. The predictive performance of the LASSO model for trisomy 21 in this retest set was also assessed.

For the purpose of comparison, two additional models were developed, using logistic regression: (1) a model based on the combination of fetal NT and maternal age (the standard method of screening in many countries); (2) a model combining all 15 markers considered in this study (maternal age, fetal NT, fetal NT ≥ 2.5 mm, prenatal thickness, nasal bone length, prenatal-thickness-to-nasal-bone-length ratio, d1, d2, prefrontal space ratio, frontomaxillary facial angle, maxilla-nasion-mandible angle, frontonasal facial angle, mandibulomaxillary facial angle, facial profile line and the F distance). The predictive performance of the LASSO model for trisomy 21 was compared with that of these two models.

Development and validation of nomogram

In order to provide clinicians with an individualized tool to predict patient-specific probability for trisomy 21, a nomogram was developed based on the LASSO model constructed in the training set. The nomogram provided a visual presentation of the LASSO model. Each marker was assigned a 'point' on the nomogram, based on its predictive ability for trisomy 21. The sum of points for a particular fetus was then converted into an overall risk score for trisomy 21.

In order to validate the performance of the nomogram, the concordance index (C-index) was calculated and a calibration curve for the nomogram was produced.

Statistical analysis

Statistical analysis was performed using R software (Version 3.5.2, <http://www.R-project.org>). Differences for

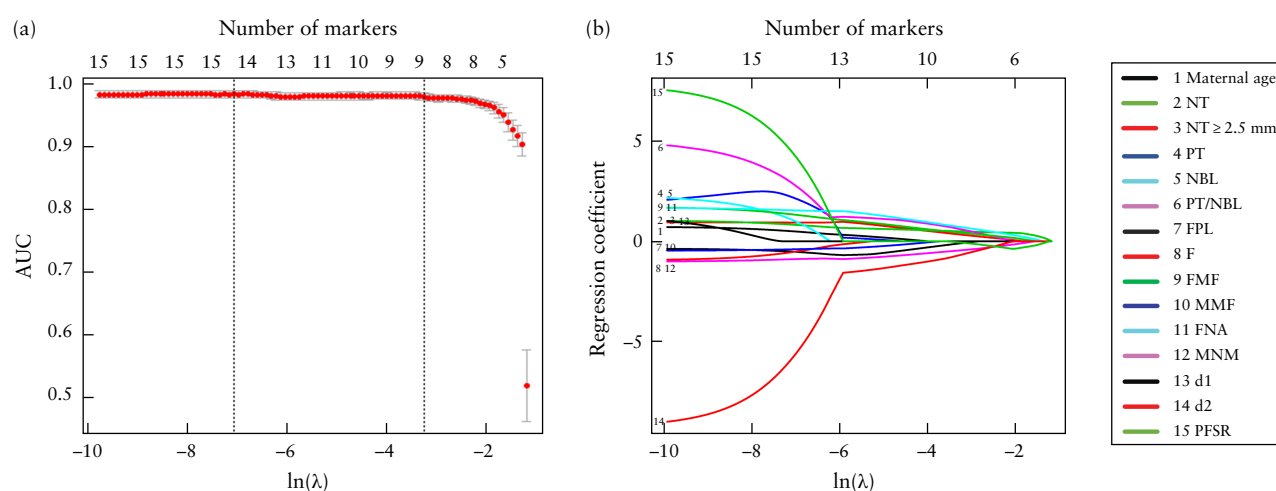


Figure 3 Selection from 15 potential markers of most useful predictive markers for trisomy 21 by least absolute shrinkage and selection operator (LASSO) method. Top x-axes represent number of selected markers and bottom x-axes represent penalty parameter $\ln(\lambda)$. (a) Illustration of relationship between area under receiver-operating-characteristics curve (AUC) of prediction model (LASSO model) and number of markers in the model. Vertical dotted line on right is at optimal value of $\ln(\lambda)$ (-3.44), at which nine markers were selected. Interval between left and right vertical dotted lines represents the range of good values for $\ln(\lambda)$. (b) Illustration of relationship between regression coefficients of markers and penalty parameter $\ln(\lambda)$. d1, distance between anterior edge of frontal bone and anterior edge of prefrontal skin; d2, distance between anterior edge of prefrontal skin (at same point as d1 measurement) and mandibulomaxillary line; F, largest perpendicular distance between facial profile line and outermost part of frontal bone; FMF, frontomaxillary facial angle; FNA, frontonasal facial angle; FPL, facial profile line; MMF, mandibulomaxillary facial angle; MNM, maxilla-nasion-mandible angle; NBL, nasal bone length; NT, nuchal translucency thickness; PFSR, prefrontal space ratio; PT, prenatal thickness; PT/NBL, prenatal-thickness-to-nasal-bone-length ratio.

numerical and for categorical variables were calculated based on the *t*-test, Mann–Whitney *U*-test or chi-square test, as appropriate. The screening performance of each marker and of each of the three models was evaluated by ROC-curve analysis. Youden's index was used to calculate the optimal threshold (cut-off) for prediction of trisomy 21. The DeLong test was used to compare ROC curves. The performance of the nomogram was assessed by C-index and Hosmer–Lemeshow test (a value > 0.05 shows good performance)¹⁶. *P* < 0.05 was considered statistically significant.

RESULTS

Maternal age and first-trimester fetal ultrasonographic markers are presented in Table 1. All markers were significantly different between the trisomy-21 and the euploid fetuses.

In the training set, nine potential markers, including fetal NT, NT ≥ 2.5 mm, prenasal-thickness-to-nasal-bone-length ratio, facial profile line, frontomaxillary facial angle, frontonasal facial angle, mandibulomaxillary facial angle, maxilla-nasion-mandible angle and d2, were identified by the LASSO method. Multivariable analysis led us to exclude fetal NT ≥ 2.5 mm, with a *P*-value of 0.63. Thus, eight markers were used to develop the LASSO model.

LASSO model

The risk scores for trisomy 21 based on the LASSO model are given in Appendix S2 (including Figures S7 and S8). The AUC of the LASSO model was higher than the AUCs of all eight individual ultrasonographic markers included in the model, in both training and validation sets (LASSO model AUC, 0.983 (95% CI, 0.971–0.994) and

0.979 (95% CI, 0.966–0.993), respectively) (Figure 4 and Table 2). The LASSO model showed good discrimination of trisomy 21 in the retest set, with an AUC of 0.997 (95% CI, 0.990–1.000) (Figure 5).

The predictive performance in screening for trisomy 21 of the model based on fetal NT and maternal age, the model based on all markers and the LASSO model are presented in Table 3. The corresponding detection rates of the three models, at false-positive rates (FPRs) of 1%, 3% and 5%, for screening for trisomy 21 are presented in Table 4. The ROC curves of the models are presented in Figure 6. The LASSO model showed significantly higher AUCs in both training (*P* < 0.001) and validation (*P* < 0.001) sets when compared with the model based on fetal NT and maternal age (Table 3), and it showed a lower AUC in the training set (*P* = 0.004), but a higher AUC in the validation set (*P* = 0.001), when compared with the model based on all markers. The model based on all markers showed significantly higher AUCs in both the training set (*P* < 0.001) and the validation set (*P* < 0.001) when compared with the model based on fetal NT and maternal age. For the LASSO model, there was no statistically significant difference in AUCs between the training set and the validation set (*P* = 0.697). For the model based on all markers, the AUC in the training set was higher than the AUC in the validation set (*P* < 0.001).

Nomogram

The predictive ability of each independent marker included in the LASSO model for trisomy 21 is reflected in the nomogram (Figure 7a); an example, based on a trisomy-21 case, is illustrated in Figure 7b. The nomogram showed good performance, with C-indices of 0.983 and 0.981 in the training and validation sets, respectively. The

Table 1 Maternal age and first-trimester fetal ultrasonographic markers in 322 euploid fetuses and 302 fetuses with trisomy 21

Parameter	Euploid	Trisomy 21	P
Maternal age (years)	33 (30–38)	36 (31–39)	< 0.001
Prenasal thickness (mm)	1.2 (1.0–1.4)	1.4 (1.2–1.7)	< 0.001
Nasal bone length (mm)	2.0 (1.8–2.1)	1.8 (1.5–2.0)	< 0.001
Prenasal-thickness-to-nasal-bone-length ratio	0.62 (0.48–0.73)	0.85 (0.71–0.97)	< 0.001
d1 (mm)	1.2 (1.0–1.4)	1.4 (1.2–1.7)	< 0.001
d2 (mm)	1.0 (0.8–1.3)	0.8 (0.6–1.0)	< 0.001
Prefrontal space ratio (d2/d1)	0.86 (0.75–1.00)	0.57 (0.43–0.75)	< 0.001
Mandibulomaxillary facial angle (°)	93.55 (92.15–96.18)	93.15 (91.30–94.57)	0.046
Frontomaxillary facial angle (°)	83.00 ± 4.88	88.84 ± 6.24	< 0.001
Maxilla-nasion-mandible angle (°)	10.02 ± 1.87	7.83 ± 2.41	< 0.001
Frontonasal facial angle (°)	115.75 ± 4.81	121.35 ± 4.90	< 0.001
NT (mm)	1.5 (1.2–1.8)	3.2 (1.9–4.2)	< 0.001
NT ≥ 2.5 mm	26 (8.1)	207 (68.5)	< 0.001
Facial profile line			< 0.001
Negative	0 (0.0)	37 (12.3)	
Zero	250 (77.6)	241 (79.8)	
Positive	72 (22.4)	24 (7.9)	
F distance (cm)*	0.16 (0.15–0.19)	0.30 (0.26–0.32)	< 0.001

Data are given as median (interquartile range), mean ± SD or *n* (%). *In cases with positive facial profile line. d1, distance between anterior edge of frontal bone and anterior edge of prefrontal skin; d2, distance between anterior edge of prefrontal skin and mandibulomaxillary line; F distance, largest perpendicular distance between facial profile line and outermost part of frontal bone; NT, nuchal translucency thickness.

indicators of the Hosmer–Lemeshow test in the training and validation sets were 0.157 and 0.132, respectively. The nomogram also showed a good calibration curve in the training set. However, the calibration curve in the validation set showed that the nomogram had a tendency to overestimate the risk for trisomy 21, in comparison to that in the training set (Figure 8a). The calibration curve of the entire dataset of 624 fetuses is given in Figure 8b.

DISCUSSION

Main findings

In this study, we developed and validated a LASSO model-derived nomogram, which incorporates fetal NT and a series of fetal facial ultrasonographic markers relating to the nasal bone, prenatal thickness and flatness of the face, for the individualized prediction of trisomy

21 in the first trimester of pregnancy. The LASSO model was shown to be superior in the prediction of trisomy 21 to a model based on fetal NT and maternal age, whilst it had a lower AUC in the training set but a higher AUC in the validation set in comparison with a model based on all 15 markers; this may be because the model based on all markers was overfitting due to the use of redundant markers¹⁷. The calibration curve of the validation set showed that the nomogram based on the LASSO model has a tendency to overestimate the risk for trisomy 21; this may be related to the limited sample size.

Comparison with previous studies

Other than fetal NT, only four of the ultrasonographic markers selected by the LASSO method had been evaluated previously for their predictive performance for trisomy 21 in the first trimester: prenatal

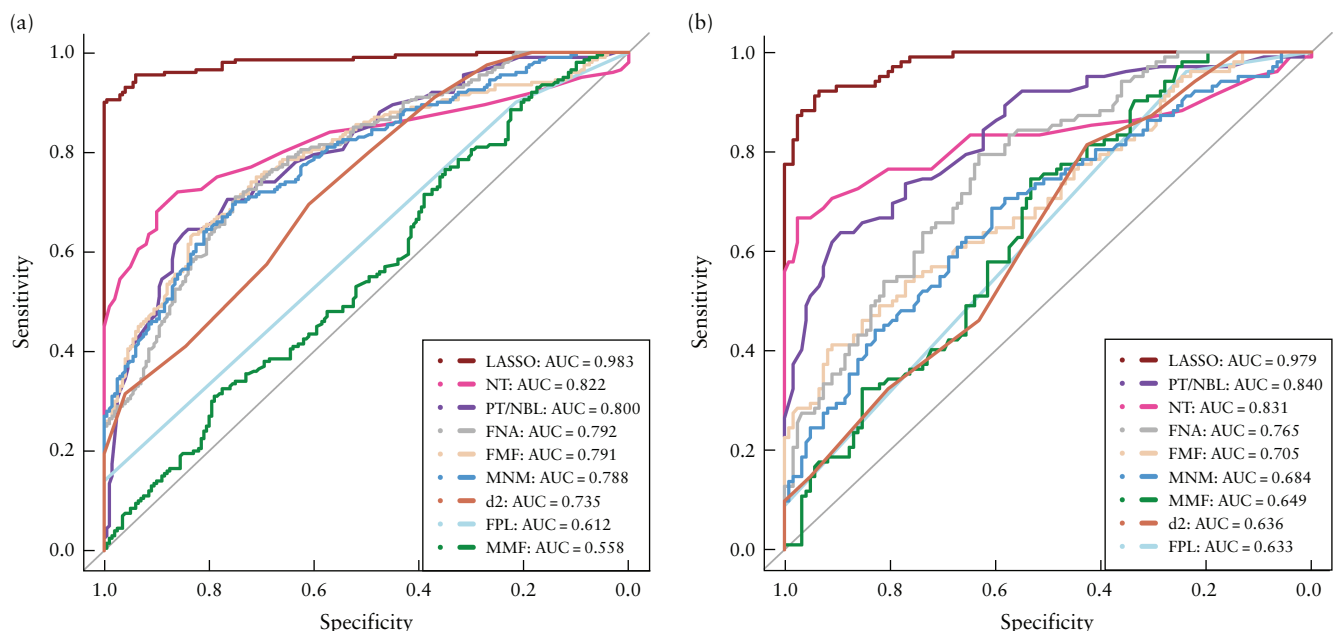


Figure 4 Receiver-operating-characteristics curves for prediction of trisomy 21 by least absolute shrinkage and selection operator (LASSO) model and, individually, eight ultrasonographic markers included in LASSO model, in training set (a) and validation set (b). Areas under the curves are given in Table 2. d2, distance between anterior edge of prefrontal skin and mandibulomaxillary line; FMF, frontomaxillary facial angle; FNA, frontonasal facial angle; FPL, facial profile line; MMF, mandibulomaxillary facial angle; MNM, maxilla-nasion-mandible angle; NT, nuchal translucency thickness; PT/NBL, prenatal-thickness-to-nasal-bone-length ratio.

Table 2 Areas under receiver-operating-characteristics (ROC) curves (AUC) for prediction of trisomy 21 by least absolute shrinkage and selection operator (LASSO) model and, individually, eight ultrasonographic markers included in LASSO model, in training and validation sets

Model/marker	Training set	Validation set
LASSO model	0.983 (0.971–0.994)	0.979 (0.966–0.993)
Nuchal translucency thickness	0.822 (0.778–0.865)	0.831 (0.770–0.892)
Prenasal-thickness-to-nasal-bone-length ratio	0.800 (0.758–0.843)	0.840 (0.788–0.891)
Frontonasal facial angle	0.792 (0.749–0.835)	0.765 (0.704–0.826)
Frontomaxillary facial angle	0.791 (0.747–0.835)	0.705 (0.637–0.773)
Maxilla-nasion-mandible angle	0.788 (0.744–0.831)	0.684 (0.614–0.754)
d2	0.735 (0.688–0.782)	0.636 (0.565–0.707)
Facial profile line	0.612 (0.574–0.651)	0.633 (0.587–0.678)
Mandibulomaxillary facial angle	0.558 (0.501–0.614)	0.649 (0.578–0.720)

Data are given as AUC (95% CI). ROC curves are shown in Figure 4. d2, distance between anterior edge of prefrontal skin and mandibulomaxillary line.

thickness-to-nasal bone length ratio, frontomaxillary facial angle, d2 and facial profile line. Our study confirmed that the first-trimester prenatal thickness-to-nasal bone length ratio in trisomy 21 fetuses is greater than that in euploid fetuses³. Borenstein *et al.*⁵ demonstrated that adding the frontomaxillary facial angle to first-trimester combined screening led to an increase in the estimated detection rate of trisomy 21, from 90% to 94% at a FPR of 5% and from 85% to 92% at a FPR of 3%⁵. We found that the frontomaxillary facial angle was increased in trisomy-21 fetuses. The same research group also reported that trisomy-18 fetuses had smaller mandibulomaxillary facial angle compared with euploid fetuses in the first trimester⁸. Our study proves that the

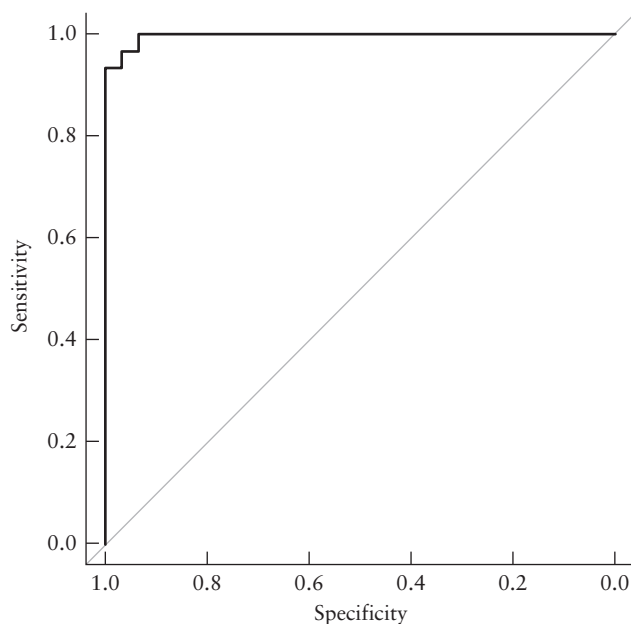


Figure 5 Receiver-operating-characteristics curve for prediction of trisomy 21 by least absolute shrinkage and selection operator (LASSO) model in retest set. Area under the curve = 0.997 (95% CI, 0.990–1.000).

mandibulomaxillary facial angle also has the potential to be an effective marker in the prediction of trisomy 21 in the first trimester. Midfacial hypoplasia is common in trisomy-21 fetuses. However, while measurement of the frontonasal facial angle is feasible at 11 + 0 to 13 + 6 weeks⁷, this has not been reported in trisomy 21. We found that the frontonasal facial angle was increased in these fetuses. Our study also found that the prefrontal space ratio and d2 were both lower in the first trimester in trisomy-21 fetuses compared with euploid fetuses⁴, confirming the suggestion of Bakker *et al.*⁴ that fetuses with trisomy 21 may have increased prenatal thickness, with lagging growth and forward displacement of the maxilla, resulting in a smaller d2 and decreased prefrontal space ratio. Neither prefrontal space ratio nor d1 was included in our LASSO model, while d2 was included. It appears that the prenatal-thickness-to-nasal-bone-length ratio is superior to the prefrontal space ratio and d1 in the prediction of trisomy 21 in the first trimester of pregnancy. In the same study, Bakker *et al.*⁴ found that the facial profile line showed no clear pattern in relation to trisomy-21 fetuses. Interestingly, a second- and third-trimester study

Table 4 Comparison of detection rates of three models in screening for trisomy 21, at false-positive rates (FPR) of 1%, 3% and 5%

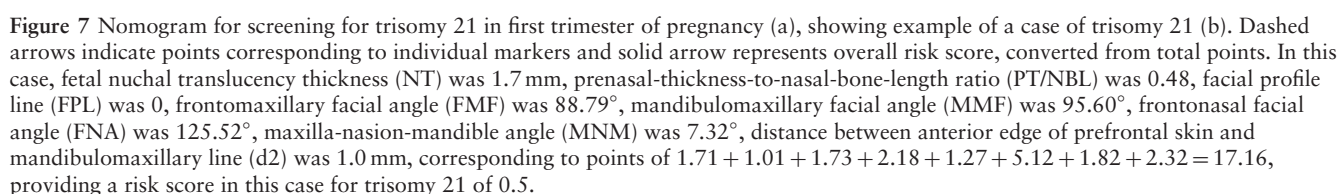
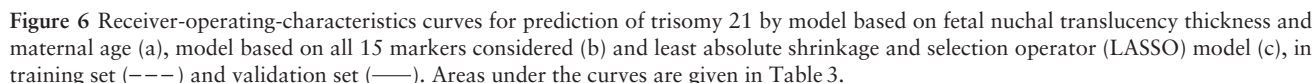
	<i>Detection rate (% (95% CI)) for:</i>		
<i>Model</i>	<i>FPR of 1%</i>	<i>FPR of 3%</i>	<i>FPR of 5%</i>
Fetal NT and maternal age			
Training set	59 (52–66)	66 (59–73)	71 (65–77)
Validation set	76 (67–84)	82 (75–90)	84 (77–91)
All markers			
Training set	92 (89–96)	96 (94–99)	98 (96–100)
Validation set	92 (87–97)	94 (90–99)	95 (91–99)
LASSO model			
Training set	90 (86–95)	91 (87–95)	93 (90–96)
Validation set	92 (87–97)	93 (88–98)	94 (90–99)

For each model, sensitivity in validation set was calculated according to cut-off of training set. LASSO, least absolute shrinkage and selection operator; NT, nuchal translucency thickness.

Table 3 Comparison of predictive performance of three models in screening for trisomy 21

Model	Detection rate (% (95% CI))	False-positive rate (% (95% CI))	Youden index (cut-off)	AUC (95% CI)
Fetal NT and maternal age				
Training set	75.0 (69.0–81.0)	7.0 (3.5–10.5)	0.68 (0.561)	0.883 (0.847–0.918)
Validation set	84.3 (77.3–91.4)	78.7 (71.4–86.0)	—	0.797 (0.729–0.866)
All-markers model				
Training set	96.0 (93.3–98.7)	1.0 (0.1–3.9)	0.95 (0.644)	0.994 (0.988–1.000)
Validation set	93.1 (88.2–98.0)	28.7 (26.8–43.7)	—	0.931 (0.896–0.966)
LASSO model				
Training set	90.5 (86.4–94.6)	0.5 (0–1.5)	0.90 (0.661)	0.983 (0.971–0.994)
Validation set	92.2 (86.9–97.4)	7.4 (2.7–12.0)	—	0.979 (0.966–0.993)

‘All-markers’ model included 15 markers: maternal age, fetal nuchal translucency thickness (NT), fetal NT ≥ 2.5 mm, prenatal thickness, nasal bone length, prenatal-thickness-to-nasal-bone-length ratio, d1, d2, prefrontal space ratio, frontomaxillary facial angle, mandibulomaxillary facial angle, maxilla-nasion-mandible angle, frontonasal facial angle, facial profile line and the F distance. LASSO model included eight markers: fetal NT, prenatal-thickness-to-nasal-bone-length ratio, facial profile line, frontomaxillary facial angle, frontonasal facial angle, mandibulomaxillary facial angle, maxilla-nasion-mandible angle and d2. Receiver-operating-characteristics curves are shown in Figure 6. AUC, area under the receiver-operating-characteristics curve; LASSO, least absolute shrinkage and selection operator.



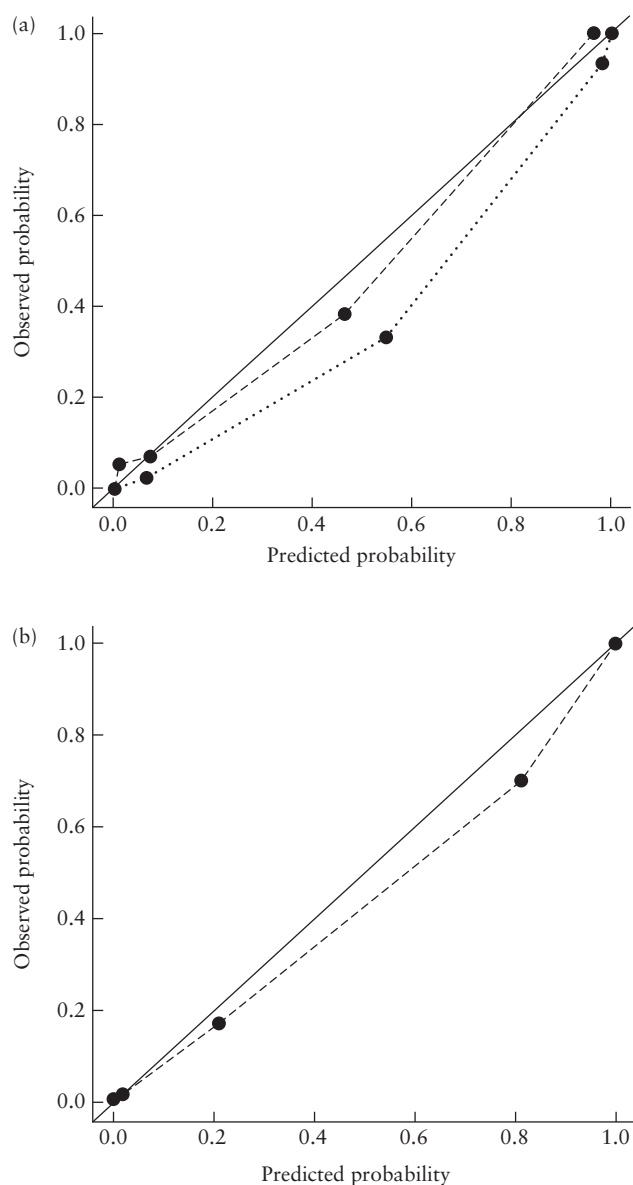


Figure 8 Calibration curves for nomogram for screening for trisomy 21 in first trimester of pregnancy, in training (····) and validation (---) sets (a) and for entire dataset of 624 fetal images (---) (b). Solid lines represent ideal performance.

reported that there were no normal fetuses with a negative facial profile line, which is consistent with our findings¹⁸. Weichert *et al.*¹⁹ found the maxilla-nasion-mandible angle to be smaller in trisomy-21 fetuses than in euploid fetuses in the second trimester of pregnancy. We found a similar relationship in our study, suggesting that this angle could be effective in the prediction of trisomy 21 in the first trimester.

The detection rate based on maternal age alone in screening for trisomy 21 is 30% at a 5% FPR²⁰. With the identification of more sensitive and specific fetal markers, one might expect that the use of maternal age will eventually be phased out. This was indeed the case in the present study, with maternal age being excluded from the LASSO model.

Implications for clinical practice

For over a decade, the first-trimester combined test, including maternal age, fetal NT and maternal serum biochemistry, with a detection rate of 90% at a FPR of 5%, has played a major role in screening for trisomy 21 in the first trimester of pregnancy. In recent years, cell-free (cf)DNA testing has improved significantly the performance of screening for trisomy 21, with a detection rate of > 99% at a FPR of < 0.1%^{21–24}. However, the cost of cfDNA testing is high, making it impossible to offer it to all pregnant women. In some countries, the current recommendation is to offer cfDNA testing contingent on the results of the first-trimester combined test^{25,26}. In China, the primary screening method for trisomy 21 is a combination of fetal NT and maternal age at 11–13 weeks' gestation. Our study has demonstrated that the LASSO model achieves better performance than does a combination of fetal NT and maternal age. This implies that, if the LASSO model is used as the primary ultrasonographic screening method, cfDNA testing could be offered to < 1% of the screened population.

Limitations

This was a retrospective study. Limitations include the fact that fetuses with ineligible images and those with absence of the nasal bone were excluded. This exclusion may have contributed to bias in our findings. In relation to the latter, in clinical practice it is well established that absence of the nasal bone is the second most important ultrasonographic marker in screening for trisomy 21. We excluded 23 of 325 (7%) cases with absence of the nasal bone. Further research is needed for development of a model that would allow evaluation of facial markers when there is absence of the nasal bone.

Conclusions

In this retrospective, case-control study, we have presented a promising, individualized nomogram that incorporates fetal NT and a series of ultrasonographic facial profile markers selected by the LASSO method and multivariable analysis. The nomogram can potentially be utilized as a convenient and effective tool in screening for trisomy 21 in the first trimester of pregnancy and requires prospective validation in future research.

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REFERENCES

- Agarwal GN, Kabra M. Diagnosis and management of Down syndrome. *Indian J Pediatr* 2014; 81: 560–567.
- Salomon LJ, Alfrevic Z, Bilardo CM, Chalouhi GE, Ghi T, Kagan KO, Lau TK, Papageorgiou AT, Raine-Fenning NJ, Stirnemann J, Suresh S, Tabor A, Timor-Tritsch IE, Toi A, Yeo G. ISUOG Practice Guidelines: performance of first-trimester fetal ultrasound scan. *Ultrasound Obstet Gynecol* 2013; 41: 102–113.
- Manegold-Brauer G, Bourdil L, Berg C, Schoetzau A, Gembruch U, Geipel A. Prenatal thickness to nasal bone length ratio in normal and trisomy 21 fetuses at 11–14 weeks of gestation. *Prenat Diagn* 2015; 35: 1079–1084.
- Bakker M, Pace M, de Jong-Pleij E, Birnie E, Kagan KO, Bilardo CM. Prenatal Thickness, Prefrontal Space Ratio and Other Facial Profile Markers in First-Trimester Fetuses with Aneuploidies, Cleft Palate, and Micrognathia. *Fetal Diagn Ther* 2018; 43: 231–240.
- Borenstein M, Persico N, Kagan KO, Gazzoni A, Nicolaides KH. Frontomaxillary facial angle in screening for trisomy 21 at 11 + 0 to 13 + 6 weeks. *Ultrasound Obstet Gynecol* 2008; 32: 5–11.
- Liu W, Qu S, Wang M, Xu W, Zhang G, Zhang C. Maxilla-nasion-mandible (MNM) angle: an indicator to assess fetal facial profile in first-trimester of pregnancy. *Springerplus* 2016; 5: 1335.
- Vicario R, Pirollo LM, De Angelis C, Narcisi M, Pietropolli A, Piccione E. Frontonasal facial angle in chromosomally normal fetuses at 11 + 0 to 13 + 6 weeks. *J Obstet Gynaecol Res* 2010; 36: 1179–1184.
- Borenstein M, Persico N, Strobil I, Sonek J, Nicolaides KH. Frontomaxillary and mandibulomaxillary facial angles at 11 + 0 to 13 + 6 weeks in fetuses with trisomy 18. *Ultrasound Obstet Gynecol* 2007; 30: 928–933.
- Dong D, Tang L, Li ZY, Fang MJ, Gao JB, Shan XH, Ying XJ, Sun YS, Fu J, Wang XX, Li LM, Li ZH, Zhang DF, Zhang Y, Li ZM, Shan F, Bu ZD, Tian J, Ji JF. Development and validation of an individualized nomogram to identify occult peritoneal metastasis in patients with advanced gastric cancer. *Ann Oncol* 2019; 30: 431–438.
- Drukker L, Noble JA, Papageorgiou AT. Introduction to artificial intelligence in ultrasound imaging in obstetrics and gynecology. *Ultrasound Obstet Gynecol* 2020; 56: 498–505.
- Bishop CM. *Pattern recognition and machine learning*. Springer: New York, 2006.
- Brattain LJ, Telfer BA, Dhyani M, Grajo JR, Samir AE. Machine learning for medical ultrasound: status, methods, and future opportunities. *Abdom Radiol (NY)* 2018; 43: 786–799.
- Namburete AI, Stebbing RV, Kemp B, Yaqub M, Papageorgiou AT, Alison NJ. Learning-based prediction of gestational age from ultrasound images of the fetal brain. *Med Image Anal* 2015; 21: 72–86.
- de Jong-Pleij EA, Ribbert LS, Manten GT, Tromp E, Bilardo CM. Maxilla-nasion-mandible angle: a new method to assess profile anomalies in pregnancy. *Ultrasound Obstet Gynecol* 2011; 37: 562–569.
- Ko HS, Lee UY, Choi SK, Park YG, Park IY, Shin JC. Craniofacial inclination at 14 to 39 weeks' gestation in normal Korean fetuses. *J Ultrasound Med* 2012; 31: 569–576.
- Kramer AA, Zimmerman JE. Assessing the calibration of mortality benchmarks in critical care: The Hosmer-Lemeshow test revisited. *Crit Care Med* 2007; 35: 2052–2056.
- Van Calster B, McLernon DJ, van Smeden M, Wynants L, Steyerberg EW, Topic Group 'Evaluating diagnostic tests and prediction models' of the STRATOS initiative. Calibration: the Achilles heel of predictive analytics. *BMC Med* 2019; 17: 230.
- de Jong-Pleij EA, Ribbert LS, Pistorius LR, Tromp E, Bilardo CM. The fetal profile line: a proposal for a sonographic reference line to classify forehead and mandible anomalies in the second and third trimester. *Prenat Diagn* 2012; 32: 797–802.
- Weichert J, Gembicki M, Ribbar-Idel J, Hartge DR. Assessment of Midfacial Hypoplasia in Down Syndrome Fetuses - Validity of a Two-Line Approach and Introduction of a Novel Angle (Maxilla-Mandible-Nasion Angle). *Ultrasound Int Open* 2016; 2: E58–62.
- Kagan KO, Wright D, Baker A, Sahota D, Nicolaides KH. Screening for trisomy 21 by maternal age, fetal nuchal translucency thickness, free beta-human chorionic gonadotropin and pregnancy-associated plasma protein-A. *Ultrasound Obstet Gynecol* 2008; 31: 618–624.
- Gil MM, Accurti V, Santacruz B, Plana MN, Nicolaides KH. Analysis of cell-free DNA in maternal blood in screening for aneuploidies: updated meta-analysis. *Ultrasound Obstet Gynecol* 2017; 50: 302–314.
- Quezada MS, Gil MM, Francisco C, Orosz G, Nicolaides KH. Screening for trisomies 21, 18 and 13 by cell-free DNA analysis of maternal blood at 10–11 weeks' gestation and the combined test at 11–13 weeks. *Ultrasound Obstet Gynecol* 2015; 45: 36–41.
- Galeva S, Konstantinidou L, Gil MM, Akolekar R, Nicolaides KH. Routine first-trimester screening for fetal trisomies in twin pregnancy: cell-free DNA test contingent on results from combined test. *Ultrasound Obstet Gynecol* 2019; 53: 208–213.
- Kagan KO, Maier V, Sonek J, Abele H, Luthgens K, Schmid M, Wagner P, Hoopmann M. False-Positive Rate in First-Trimester Screening Based on Ultrasound and Cell-Free DNA versus First-Trimester Combined Screening with Additional Ultrasound Markers. *Fetal Diagn Ther* 2019; 45: 317–324.
- Nicolaides KH, Syngelaki A, Poon LC, Gil MM, Wright D. First-trimester contingent screening for trisomies 21, 18 and 13 by biomarkers and maternal blood cell-free DNA testing. *Fetal Diagn Ther* 2014; 35: 185–192.
- Nicolaides KH, Wright D, Poon LC, Syngelaki A, Gil MM. First-trimester contingent screening for trisomy 21 by biomarkers and maternal blood cell-free DNA testing. *Ultrasound Obstet Gynecol* 2013; 42: 41–50.

SUPPORTING INFORMATION ON THE INTERNET

The following supporting information may be found in the online version of this article:



Appendix S1 Definition and delineation of ultrasonographic markers (including Figures S1–S6)

Appendix S2 Risk scores for trisomy 21 based on the LASSO model (including Figures S7 and S8)