Ultrasound Markers of Aneuploidy in the First Trimester

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ABSTRACT

Prenatal screening of fetal aneuploidy is a continuously and rapidly evolving area of research; there have been tremendous advancements over the past decades in prenatal screening for aneuploidy, especially during the first trimester. As there is extensive evidence that effective screening for major chromosomal abnormalities can be provided in the first trimester of pregnancy, recently we have changed our practice; the prenatal screening of fetal chromosomal aberrations has been moved and pointed to the first trimester. Besides the nuchal translucency, which is one of the most known ultrasonographic markers, there are other markers, which can be examined during the first trimester of pregnancy. To maximize the quality of sonography, increase the screening sensitivity, and decrease the range of false-positive rate, all of the first-trimester ultrasound markers have well-established criteria for the measurement. With the use of high standards of scanning, the early recognition of sonographic markers of chromosomal aberrations can be helpful in forward prenatal diagnosis. On the contrary, the early diagnosis makes the termination of the pregnancy possible with fewer complications, and there is time for planning of further follow-up and interventions.

Keywords: Aneuploidy, First trimester, Marker, Screening, Ultrasound.


INTRODUCTION

With the evolution of ultrasound technology, a large gain in efficiency of detecting aneuploid pregnancies has occurred. The recent trend is to recognize and establish useful sonographic markers as soon as possible, especially at the first trimester of the pregnancy. Over the last decades, the advancements in medicine and technique have improved the methods of detection of chromosomal aberrations. A significant number of fetal structural anomalies develop by the end of the first trimester; hence, besides the well-known second-trimester markers, there are real opportunities to recognize chromosomal abnormalities at an early gestational age.

During the first trimester, the ultrasound examination can be performed with the transvaginal or transabdominal technique (the recommended procedure is transabdominal sonogram). The most common chromosomal aneuploidy, such as trisomies 21, 18, and 13, might have a characteristic appearance at the first trimester; however, the establishment of the early screening methods can give a real opportunity to decrease the prevalence of neonates born with chromosomal aberrations as soon as possible. The early recognition of sonographic markers of chromosomal aberrations can be helpful in forward prenatal diagnosis. On the contrary, the early diagnosis makes the termination of the pregnancy possible with fewer complications, and there is time for planning of further follow-up and interventions. Prenatal cytogenetic analysis or noninvasive testing should be offered to women with fetal ultrasound markers of aneuploidy diagnosed in the first trimester.

NUCHAL TRANSLUCENCY

In 1985, Benacerraf et al.2 presented the ultrasound sign of enlarged nuchal fold thickness in second-trimester fetuses. In 1992, Nicolaides et al.3 described enlarged nuchal translucency (NT) in first-trimester fetuses with trisomy 21.3 Nuchal translucency refers to the normal subcutaneous fluid-filled area between the back of the embryonal/fetal neck and the overlying skin (Fig. 1).
As a measurement mistake can have a major consequence, it is very important to establish the criteria for NT measurement. The Fetal Medicine Foundation (FMF) has introduced a process of training and certification to help establish high standards of scanning on an international basis. The FMF proposed sonographic criteria (Box 1) to maximize the quality of NT sonography, which were used successfully.4

Box 1: Sonographic criteria to maximize quality of nuchal

Fetal NT can be measured successfully by transabdominal ultrasound examination in about 95% of cases (in the others, it is necessary to perform transvaginal sonography – the results from transabdominal and transvaginal scanning are similar).4 For the sonographic examination of NT, the section obtained is a midsagittal section of the embryo/fetus [this is the section where the measure of the crown-rump length (CRL) is obtained].

Box 1: The NT measurement (FMF criteria)

• The gestation should be 11 to 13 + 6 weeks and the fetal CRL should be 45 to 84 mm.
• A midsagittal section of the fetus should be obtained and the NT should be measured with the fetus in the neutral position.
• Only the fetal head and upper thorax should be included in the image. The magnification should be as large as possible and always such that each slight movement of the calipers produces only a 0.1 mm change in the measurement.
• The maximum thickness of the subcutaneous translucency between the skin and the soft tissue overlying the cervical spine should be measured. Care must be taken to distinguish between fetal skin and amnion.
• The calipers should be placed on the lines that define the NT thickness – the crossbar of the caliper should be such that it is hardly visible as it merges with the white line of the border and not in the nuchal fluid.
• During the scan, more than one measurement must be taken and the maximum one should be recorded.

Only the fetal head and upper thorax should be included in the image for measurement of NT. The optimal gestational age for measurement of fetal NT is 11 weeks to 13 weeks and 6 days. The minimum fetal CRL should be 45 mm and the maximum 84 mm. The procedure of the measurement should be performed exclusively at the level of maximum thickness of the subcutaneous translucency between the skin and soft tissues overlying the cervical spine (Fig. 2A). The calipers will be placed “on to on,” perpendicular to the long axis of the fetal body (Fig. 2B). For adequate measurement, it is practical to zoom into the selected area.

Sometimes, a false-positive result in the measurement can happen if the amnion is not truly separated from the fetal skin (the movement of the fetus can account for these situations).

Furthermore, the umbilical cord or sometimes the presence of amnion band can also distract the measurement. For these reasons, application of the technique and training of the sonographers are very important.

The NT is usually considered abnormal if it is greater than 3 mm, but it is known that NT thicknesses extend with the gestational age. According to the recent most proposed protocol, if the NT is 3 mm or larger, then genetic counseling is advisable.

The mechanism of fluid accumulation in the fetal neck, which produces the enlarged NT, is still unknown.5-8 However, it is possible that the physiopathological process leading to NT may be related to cardiac malformations, early hemodynamic disorders of the affected fetuses, or abnormalities of the extracellular matrix of the skin in fetuses with trisomy 21.9-11

In first-trimester scans (11–14 weeks), approximately 75 to 80% of fetuses with trisomy 21 present an enlarged NT (Fig. 3); however, 5 to 10% of healthy fetuses also show the same sign.12-14 Malone and D’Alton4 evaluated 30 studies on the subject. These studies describe 316,311
patients who were screened with NT sonography during the first trimester of pregnancy. The overall sensitivity for trisomy 21 was 77%, with a false-positive rate of 6% (sensitivities varying from 29 to 100%, false-positive rates varying from 0.3 to 11.6%). The study established that abnormal NT is 13 times more likely when trisomy 21 is present compared with the healthy fetuses.

In conjunction with maternal age, measurement of NT thickness between 11 and 14 weeks demonstrated a detection rate of 77%. A combination of NT thickness measurement, maternal age indicator, and first-trimester serum markers should increase detection rates while avoiding false-positive findings. In this combination, the rate of detection is increased from 90 to 92% with a 3% false-positive rate. Furthermore, the combination of NT test with the maternal age of more than 35 years and ultrasound screening markers in the second trimester (which is practiced in most centers) has shown increase in accuracy of the screening test. Technique in NT imaging is essential for accurate risk assessment (undermeasurement by even 0.5 mm can reduce the test sensitivity by 18%). The NT can be also enlarged in other chromosomal aberrations, such as trisomies 18 and 13, Turner’s syndrome (monosomy X), Klinefelter’s syndrome, and triploidy. Currently, the relatively small experience with three-dimensional (3D) technique of this subject does not show any advancement with the NT measurement.

Cystic Hygroma

Fetal nuchal cystic hygroma (NCH) can be determined as an area of sonolucency in the soft tissue of the occipital region (Figs 4 and 5). The NCH consists of two symmetrical cavities completely separated by a nuchal ligament, with or without internal trabeculae (septated and nonseptated NCH).

The etiopathogenesis of the cystic hygroma is different from NT. The NCH is the congenital malformation of the lymphatic system (lymphatic stasis).

The association with chromosomal abnormalities, such as Turner’s syndrome (the most common), trisomies 21, 18, and 13, and Klinefelter’s syndrome, has been reported. The volume of hygroma and the presence of septa are associated with higher incidence of chromosomal aberrations.

Fetal Growth Disorders

The biometry of embryo/fetus in the first trimester consists of the measurement of CRL (Fig. 6). During this time, biological variation in the fetal size is minimal. Fetal growth disorders in the first trimester could be associated with chromosomal aberrations; CRL, compared with what is expected, is substantially less. The CRL, i.e., smaller or more than 7 mm expected indicates a three times higher risk for chromosomal aberration. Trisomy 18 is associated with severe early-onset growth restriction, which is more severe than in trisomies 21 and 13.
NASAL BONE

The absence of the nasal bone can be a useful ultrasound marker for identifying fetuses with trisomy 21 in the first trimester of pregnancy. Among other chromosomal abnormalities, such as trisomies 9 and 18 and Turner’s syndrome, different defects of the bone and connective tissue have been described, and the absence of nasal bone has been recognized. Between the 11th and 14th weeks of gestation, the fetal nasal bone can be visualized by ultrasonography in 99.5% of chromosomally normal fetuses. This finding is compatible with the results of histological and radiological studies of aborted fetuses, which showed that the intramembranous ossification process of the nasal bone first appears at a CRL of 42 mm and increases linearly with gestation. There is a significant difference in the rate of visualization of the fetal nasal bones in the first trimester in mothers of different ethnic origins.

For visualization of the fetal nose (Box 2), the obtained section is a midsagittal view of fetal profile (Fig. 7), with an angle of incidence between the beam of the ultrasound transducer and the line traced from front to the chin of the fetus at about 45° or 135°. The importance of the angle of insonation cannot be underestimated; nasal bone may artificially appear to be absent if the angle is inadequate. In the midsagittal position, two echogenic lines can be seen, superficial line representing the skin of the nose, and the inner one representing the ossificated nasal bone (Fig. 8). The echoes from the skin of the nose can be misinterpreted as the nasal bone. To avoid this mistake, the ultrasound transducer must be gently tilted from side-to-side to ensure that the nasal bone was seen separately from the nasal skin.

A full view of the fetal profile had a success rate of 94 to 99%, meaning it was not always possible for sonographers to confirm the absence or presence of the nasal bone. For visualization of the fetal nose (Box 2), the obtained section is a midsagittal view of fetal profile (Fig. 7), with an angle of incidence between the beam of the ultrasound transducer and the line traced from front to the chin of the fetus at about 45° or 135°. The importance of the angle of insonation cannot be underestimated; nasal bone may artificially appear to be absent if the angle is inadequate. In the midsagittal position, two echogenic lines can be seen, superficial line representing the skin of the nose, and the inner one representing the ossificated nasal bone (Fig. 8). The echoes from the skin of the nose can be misinterpreted as the nasal bone. To avoid this mistake, the ultrasound transducer must be gently tilted from side-to-side to ensure that the nasal bone was seen separately from the nasal skin.

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The process of ossification is extremely complex and numerous genetic syndromes and chromosomal anomalies can be implicated in the origin of alterations and defects in the development of the cranial skeleton. Among other things, the curiosity of the growing of fetal bone is dependent on the surrounding functional matrix. Immunohistochemical studies of fetuses with trisomy 21 have shown alterations in the composition of extracellular matrix, which might be attributed to gene dosage effects. This can be supported by the biochemical and molecular–genetic changes in trisomy 21, namely trisomy 21 is associated with a substantial increase of hyaluronic acid. This increase could be a consequence of increased superoxide dismutase, which is encoded in chromosome 21 and protects against free radical-mediated degradation of hyaluronic acid. Likewise, the genes for two of the three polypeptide chains (a1, a2, and a3) of collagen type IV are found on chromosome 21 (COL6A1 and COL6A2), and it is known that the dermis of fetuses with trisomy 21 is rich in this collagen.

As a probable consequence of hypoplasia or delayed ossification of the nasal bone (Fig. 9) in fetuses with...
trisomy 21, the nasal bone is not visible in 60 to 73% of first-trimester trisomy 21 fetuses. However, the prevalence of the absent nasal bone among healthy first-trimester fetuses is low (0.3–0.6%).

MEGACYSTIS

During the first trimester of pregnancy, measuring of fetal bladder is the simplest method for screening the fetal urinary system (Fig. 10). The fetal bladder can be visualized at a CRL of 67 mm, as a spherical hypoechoic mass within the fetal pelvis. Megacystis at the first-trimester fetuses is defined as enlargement of the fetal bladder (longitudinal diameter of the fetal bladder is >6 mm) (Fig. 11). The megacystis can be arranged as being either mild-to-moderate (8–12 mm) or severe (>17 mm). The fetal bladder diameter/CRL ratio can be an important sample at the assessment, normal range of bladder diameter/CRL is less than 10%. The association with chromosomal abnormalities has been reported. There is a correlation between the extent of megacystis and risk of chromosomal defects: In the moderate fetal megacystis, the risk of chromosomal aberration is about 25%, in the severe fetal megacystis, this risk is about 10% at 10 to 14 weeks of gestation.

DUCTUS VENOSUS VELOCEMISTRY

The ductus venosus is a primary important vascular structure during the intrauterine life. The importance of this structure is carrying well-oxygenated blood from the placenta and umbilical vein directly to the cerebral and coronary circulation (through the foramen ovale to the left atrium). It behaves as an “arterialized” vessel with a high pulsatile flow and forward velocities throughout the cardiac cycle.

The ductus venosus blood flow can be detected as early as 8 weeks of gestation. For the ductus venosus velocimetry, the obtained plane is a midsagittal section of the fetal trunk (a narrow, trumpet-like structure). The Doppler sample gate must be placed at the initial portion of the ductus venosus where it originates from the umbilical vein (interrogation angle is < 60°). Blood flow in the ductus is characterized by high velocity during ventricular systole (s-wave) and diastole (d-wave) and a presence of forward flow during atrial contraction (a-wave) (Fig. 12A). The measurement of ductus venosus pulsatility index of the veins (DVPIV) happened on three to five consecutive high-quality ductus venosus waveforms.

Overlap signals originating from different adjacent vessels can cause difficulties in distinction between normal and abnormal waveforms. Ductus venosus flow velocity is approximately three times higher than the flow velocity in the umbilical vein or inferior vena cava.

The absent or reversed flow during the a-wave or a DVPIV of more than 95 percentile (Fig. 12B) can be sign of chromosomal aberration; it has been seen in about 70 to 90% of fetuses with a chromosomal abnormality. Among fetuses with trisomy 13, ductus venosus studies were found to be more frequently normal.
PULSATILITY INDEX OF THE UMBILICAL ARTERY

Some studies with umbilical artery pulsatility index (UAPI) have shown that in first-trimester fetuses with trisomy 21, a significantly higher UAPI value is seen than in healthy fetuses. Intriguing is that other studies of the subject did not find significant difference at the UAPI between fetuses with trisomy 21 and normal fetuses. The question is still open and further large prospective studies are needed to explain these differences.

FETAL HEART RATE

Fetal heart rate (FHR) is detectable at the 6th week of gestation by transvaginal sonography. In this gestational time, the mean heart rate is about 100 beats per minute (bpm). In normal pregnancy, the FHR increases from about 110 bpm at 5 gestational weeks to about 160 to 170 bpm at 9 weeks and then slightly decreases. The early increase in FHR coincides with the morphological development of the fetal heart, and the subsequent FHR decrease might be the result of functional maturation of the fetal parasympathetic system. The FHR abnormalities can be found among aneuploid fetuses. In trisomies 21 and 13, monosomy X, the significant mean increase in the fetal heart frequency has been described. Then again, in fetuses with trisomy 18 or triploidy, the mean FHR is significantly reduced.

UMBILICAL CORD

Diameter of umbilical cord can be measured during the first trimester (10–14 weeks scan). There is the observation that fetuses with chromosomal defects, compared with healthy fetuses, have an increased diameter of the umbilical cord (above the 95th centile of reference value). Normally, there are two umbilical arteries around the fetal bladder, which can be visualized during the first-trimester scan (Fig. 13). The presence of two-vessel cord (one vena and one artery) can also be a marker for fetal aneuploidy (singular umbilical artery).

OMPHALOCELE

The return of intestine into the peritoneal cavity normally takes place around the 12th week of gestation. Omphalocele (exomphalos) is a ventral wall defect characterized by the herniation of intra-abdominal organs (bowl loops, stomach, and liver) into the base of umbilical cord, with a covering amnioperitoneal membrane. The omphalocele can be diagnosed during the first-trimester scan (above 12th week of gestation). Association with chromosomal aberrations is 35 to 58%. The 3D first-trimester ultrasonography can be helpful in identification of the omphalocele.

OTHER ABNORMALITIES

Fetal kidney and bladder can be clearly seen in the 12th week of gestation. The size of normal fetal kidney...
pelvis at the end of the first trimester has not yet been established. The fetal pyelectasis is unilateral or bilateral dilatation of the renal pelvis. The proposed 3 mm, as a higher level for normal renal pelvis anteroposterior diameter, can be used for the border of pyelectasis at the end of first trimester.

Presence of the bilateral pyelectasis increases the risk of aneuploidy by about 1.5 to 2 times. The incidence of pyelectasis among fetuses with trisomy 21 is about 17 to 25%, compared with only 2.1 to 2.8% in healthy fetuses.

Choroid plexus cysts (CPCs) are relatively common in the first trimester. The choroid plexus can be visualized as early as 8 weeks of gestation. The size, form, position of choroid plexus do change during the early stage of pregnancy. In the first trimester, the size of the majority of CPCs is 1 to 2 mm. The CPCs are significantly associated with trisomy 18 at midgestation. The association of CPCs with other chromosomal abnormalities is controversial.

Fetal structural malformations are evident in the second and third trimesters. First-trimester studies of fetal anatomy are limited (for the technical reasons and due to the natural development of the organs). Certain structural malformations in the first trimester have recently been described in association with chromosomal abnormalities (holoprosencephaly, ventriculomegaly, facial cleft, micrognathia), etc.

**FUTURE POSSIBILITIES**

With the evolution of ultrasound technique, the advancement of sonographic markers for fetal aneuploidy is expected.

Experiences with the fetal iliac angle measurement at the second trimester can be useful at the first-trimester scan, but this statement is unproven yet. The fetal iliac bones can be visualized during the first-trimester scan; the angle between them can be measured (Fig. 15). In the future, large prospective studies are needed to investigate the usefulness of this marker.

**REFERENCES**


