Recommendation and Guidelines for Perinatal Practice

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Controversial ultrasound findings in mid trimester pregnancy. Evidence based approach

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Abstract: Mid trimester fetal anatomy scan is a fundamental part of routine antenatal care. Some US soft markers or controversial US signs are seen during the scan and create some confusion regarding their relation to fetal chromosomal abnormalities. Example of these signs: echogenic focus in the heart, echogenic bowel, renal pyelectasis, ventriculomegaly, polydactely, club foot, choroid plexus cyst, single umbilical artery. We are presenting an evidence based approach from the literature for management of these controversial US signs.

Keywords: Fetal cardiac echogenic foci; fetal choroid plexus cyst; fetal club foot; fetal echogenic bowel; fetal mild ventriculomegaly; fetal polydactyly; fetal renal pyelectasis; single umbilical art.

Introduction

Ultrasound (US) has became a crucial tool in the field of fetal medicine.

First and mid trimester scans in particular have become a routine part of antenatal care. These ultrasounds can detect congenital anomalies that are solitary, or part of an underlying chromosomal anomaly. In addition there are a growing number of soft markers that can be seen with US that are associated with chromosomal anomalies or poor pregnancy outcomes.

As US technology has advanced the detection of these soft markers has become easier. With the increased detection of these markers comes confusion regarding the appropriate parental counseling and fetal management. This dilemma is born out of a knowledge gap between the presence of these markers and their clinical significance. We are going to present an evidence-based approach for these markers. We will include the epidemiology, pathophysiology, and underlying risk of associated chromosomal anomalies. We will also include the potential impact on the fetus during the pregnancy and after birth. Our aim is to produce a protocol for the management of pregnancies effected by the presence of these markers.

Echogenic focus of the heart

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An echogenic focus (EF) of the heart is defined as an echogenic area located in the region of the papillary muscles but not attached to the ventricular walls. They move with the atrioventricular valve and can occur in either cardiac ventricle, but are mainly seen in the left ventricle [1].

EF are seen in approximately 4% of obstetric sonograms with the lowest prevalence in Black populations and highest rates seen in Asian women [2]. The exact etiology of these foci is unclear. While some consider them to be calcifications within the fetal papillary muscle, or collections of fibrous tissue with increased echogenicity, others theorize that they are micro calcifications within the cardiac muscle [2].

In the early 1990s it was reported to be associated with aneuploidy; however, the exact pathophysiologic link remains uncertain. This relation was confirmed in multiple studies, however, most of the studies were with women at high risk for aneuploidy [3].

In a metaanalysis evaluating the relation of EF with Down syndrome (T21), including 11 studies with a total of 51,831 patients, the authors concluded that the prior risk of Down syndrome (as calculated by maternal age, history of previous affected pregnancy and prior screening tests) is increased by a factor of 5.4 when EF are present and reduced by a factor of 0.8 when no foci are detected [3].

In a recent study on 12,373 cases, 267 EFs were diagnosed, 149 of them were isolated, and there were no cases of T21 among women less than 35 years of age [4]. In another prospective study of 12,672 women, 479 cases of EF were diagnosed, of which 90% were isolated with only one case of T21 (+LH ratio of 2.66). This study concluded that amniocentesis is not warranted in low risk cases with isolated EF [5].

In the absence of aneuploidy, EF has not been associated with structural cardiac abnormalities or childhood myocardial dysfunction when compared with the general population [6].

Suggested protocol of management of EF:
1. Detailed fetal anatomy to search for any associated anomalies
2. Isolated EF is considered as incidental finding and does not warrant amniocentesis or further evaluation either prenataley or postnataley
3. Amniocentesis should be done to rule out aneuploidy only in high risk cases, such as: >35 years of age, associated abnormalities, other soft markers, or history of chromosomally abnormal pregnancies.

Choroid plexus cysts are cysts that are seen within the substance of the choroid plexus. They can be single or multiple, unilateral or bilateral, and occur with an incidence of approximately 1%. They may result from entrapment of cerebral spinal fluid with tangled villi. As the stroma of decreases with increasing gestational age this fluid is released and the cyst resolves. For this reason more than 95% of these cysts resolve by the end of the second trimester [7].

The presence of a CP cyst has been associated with increased risk of trisomy 21, but even more so with trisomy 18 [8]. Seventy-one percent of trisomy 18 fetuses have CP cysts but these are usually associated with additional sonographic abnormalities [9]. The location, size, and morphology do not affect its relation to aneuploidy [9]. Isolated CP cysts create confusion, as its link to increased risk of chromosomal anomalies is less clear [9]. In a metaanalysis of more than 2000 cases with an isolated CP Cyst showed that trisomy 18 was found in one case out of 128 [10]. Based on the findings of this metaanalysis, the American College of Obstetricians and Gynecologists recommends offering amniocentesis in the case of an isolated CP cyst only if the age is >35 or with abnormal serum multiple marker screen [11]. Another large metaanalysis with 246,545 cases found 1346 cases of isolated CP cyst. The study concluded that only when a CP cyst was present with additional risk factors was an amniocentesis warranted [8].

Suggested protocol for the management of a CP cyst:
1. Careful examination for the fetal anatomy for any additional abnormalities:
   a. careful examination of the hands and feet are very helpful as most of the cases with trisomy 18 are associated with limb posture abnormalities
   b. A FE can be considered.
2. No invasive procedures for aneuploidy except if:
   a. maternal age >35 years
   b. presence of any other sonographic abnormalities
   c. positive serum aneuploidy screen.

3. Parents should be advised that most of these cysts will disappear during pregnancy and that they are not associated with any long-term effects like mental retardation, cerebral palsy, or delayed development.

4. Follow-up ultrasounds are not generally needed, because most CP cysts resolve.

5. Follow up for growth pattern may be suggested in women with high risk who refrain from doing an invasive test since trisomy 18 is usually associated with intrauterine growth restriction (IUGR) [12].

Since the attitude and reaction towards CP cyst and EF has been changing, it was not surprising to see a paper published recently with an interesting title: “It is time to reconsider our approach to EF and choroid plexus” [13].

**Single umbilical artery**

A single umbilical artery (SUA) is the most common anomaly of the cord and occurs in approximately 1% of all deliveries [14]. It may result from primary aplasia of one of the two umbilical arteries or as a consequence of atrophy of one artery [14]. It is more common in the left artery and occurs more often in twin gestation. It is sometimes accompanied with abnormal cord insertion in the placenta, i.e. marginal and velamentous cord insertions [14].

The sonographic diagnosis can be made either by visualizing a transverse section of a free loop of the cord or by using color Doppler to identify both arteries on both sides of the bladder in a longitudinal section of the abdomen. Using the latter method has the benefit of indentifying which side is missing [15]. The diameter of the cord with only two vessels tends also to be larger than a three vessel cord as well, with a diameter of >4 mm or a vein/artery ratio of <2 perhaps being diagnostic of SUA [15]. Lastly, multiple segments should be examined to exclude fusion of the two arteries.

SUA has been associated with fetal anomalies and also with a worse pregnancy outcome when compared to fetuses with two umbilical arteries [16]. In general the increased fetal morbidity and mortality associated with pregnancies with SUA is mainly attributable to increased rates of associated anomalies and aneuploidy [16, 17]. The most common associated anomalies with SUA are cardiac and genitourinary [16]. SUA at 11–14 weeks has a high association with trisomy 18 and other chromosomal defects [17]. Dagklis et al. showed that the finding of an SUA should prompt the sonographer to search for other fetal defects, and if these are found, the risk for chromosomal abnormalities is increased. In cases of apparently isolated SUA there is no evidence of increased risk of chromosomal abnormalities [18].

Suggested protocol of management of SUA:
1. A detailed fetal anatomy scan should be done as well as fetal echocardiography.
2. In cases with isolated SUA and maternal age <35 years, invasive testing is not warranted.
3. Serial growth scans are warranted because SUA has been associated with increased rates of IUGR.
4. Color Doppler of the umbilical artery should be offered whenever there is oligohydramnios or IUGR [17].

**Mild ventriculomegaly**

Ventriculomegaly is a condition caused when there is dilated atrium beyond 10 mm. The mild ventriculomegaly (MVM), or what is called borderline ventriculomegaly, range between 10–12 mm and 10–15 mm [19]. It can be an isolated finding or be associated with an underlying cranial defect or anomaly such as agenesis of the corpus callosum. It is seen in almost 0.1–0.2% of births. In a study of 140 fetuses with both isolated and non-isolated ventriculomegaly only 5% had an abnormal karyotype and only 4.3% of fetuses with MVM were abnormal. Interestingly isolated ventriculomegaly was associated with a higher risk of karyotypic abnormality as compared to those with additional anomalies [20]. The authors therefore recommended karyotype analysis in all cases of ventriculomegaly.

The remaining issue now is parental counseling. The implication of MVM without underlying cranial abnormality or fetal abnormality is slightly confusing due to the lack of good quality postnatal follow-up studies in
the literature. This makes antenatal counseling for this
abnormality very difficult. In one study of 60 cases of
fetuses with isolated MVM of 10–12 mm that were fol-
lowed up to 10 years of life, they found that all of the
children displayed normal neurodevelopment. They
concluded that MVM of 10–12 mm may be considered a
normal variant [21].

In a recent review of cases with MVM up to 15 mm with
postnatal follow-up they found that about 11% of infants
with isolated MVM will display abnormal or delayed neu-
rodevelopment. However, the authors go on to state, in
collection, that given the lack of good quality postnatal
follow-up studies it is difficult to provide accurate antena-
tal counseling [22].

Suggested protocol in cases of MVM:
1. First reassess the atrial diameter using the recom-
mended criteria.
2. Perform a careful survey of the fetal brain to exluce
concominant pathology.
3. Follow-up US should be performed to determine if the
lesion is progressing
4. It is reasonable to offer chromosomal analysis to all
patients.
5. Counsel parents that there is limited evidence on the
long-term neurodevelopmental consequences.

Renal pyelectasis

Dilatation of the renal pelvis is a common finding with
an incidence reported to be between 0.3 and 4.5%. It
is more common in male fetuses, and is often bilateral;
however, if it is unilateral it is more likely to present on
the left side. Although laterality does not seem to useful as
a prognostic indicator, according to one study unilateral
pyelectasis is associated with a higher rate of urinary tract
abnormalities at birth [23].

Mild pyelectasis refers to dilatation of the renal pelvis >4–5 mm and <10 mm in the antero-posterior diam-
ter measured in transverse section of the fetal abdomen. The
cut-off value which is most frequently used in this dimension
is >4 mm in the 2nd trimester and >7 mm thereafter [23].

The relation of mild pyelectasis and aneuploidy,
mainly trisomy 21, was first suggested by Benacerraf et al.
in 1990 [23]. Others studies have subsequently supported
this finding, however, all studies were done on high-risk
populations [23]. Additionally, a large multicenter prospec-
tive observational study of unselected fetuses between 16
and 26 weeks found that its association was strongest in
the presence of other anomalies [24].

Examining cases of isolated mild pyelectasis has
found that there is very little association with aneu-
uploidy. In a retrospective study of 25,582 low-risk
pregnancies, 301 cases of isolated pyelectasis (>5 mm)
were detected and none had aneuploidy [25]. In another
study, Coplen and Jeanty studied 12,672 cases of which
2.9% had mild pyelectasis (>4 mm). Eight-three percent
of these were isolated, and a likelihood ratio for T21 was
3.79 [26]. They concluded that in the absence of other
findings, isolated pyelectasis is not a justification for
amniocentesis. Lastly, another study of isolated pyelec-
tasis was found to have a sensitivity of 0.02 for diagno-
sing fetuses with Down syndrome. In that case it would
be necessary to screen 30,404 women in order to find
one case of Down’s [27].

Pyelectasis, as mentioned above, can also be a
marker for possible postnatal urinary tract abnormal-
ity. In 60–70% of fetuses found to have pyelectasis, the
dilation remains stable. In only 1/3 does the pyelectasis
progress [28]. Wollenberg et al. showed that none of 20
children with a prenatal diagnosis of mild renal pelvis
dilatation (7–9.9 mm) during the 3rd trimester experi-
enced a urinary tract infection or underwent surgery
[28]. In contrast five of 22 (23%) cases with a renal pelvis
diameter of 10–14.9 mm and 23 of 36 (64%) cases with
severe hydronephrosis (>15 mm) had either a UTI or
required surgery after birth.

Suggested protocol in cases of renal pyelectasis:
1. Accurate measurement of the renal pelvis in the
proper plane.
2. Careful search for associated fetal anomalies
abnormalities.
3. Fetal echocardiogram to evaluate the fetal heart.
4. In the absence of other fetal anomalies, soft markers,
or risk factors there is no need for invasive aneuploidy
testing.
5. Thirty percent of cases with mild pyelectasis
will proceed to hydronephrosis, and therefore
follow-up of the renal pelvis diameter in the 3rd trimester is recommended.

6. All infants with persistent mild pyelectasis should undergo postnatal evaluation.

Echogenic bowel

The bowel is considered to be echogenic when it is bright compared to the adjacent bone. A echogenic bowel is reported to be present in 0.2–1.4% of 2nd trimester USs and be diffuse or focal. When present it can be associated with aneuploidy (trisomy 21), congenital infection (CMV, toxoplasmosis, parvovirus), cystic fibrosis, intra amniotic bleeding, IUGR, and thalassemia [29]. The pathophysiology of echogenic bowel differs according to the etiology [29]. In T21, poor bowel motility results in thickened meconium. In fetal infection meconium peritonitis and bowel edema results in perforation and focal calcification at the perforation sites. In IUGR the cause is areas of ischemia due to redistribution of blood flow away from the gut. In cystic fibrosis it is caused by abnormal pancreatic enzymes leading to change in meconium consistency. This leads to diffuse or focal echogenic areas with dilated bowel.

In intra-amniotic bleeding the swallowing of blood causes the increased echogenicity.

The diagnosis is made by comparing the echogenicity of the bowel to that of the liver and adjacent bone, such as the iliac crest. This can be accomplished by turning the gain setting down until other soft tissues are no longer seen and only the bone and bowel is visualized [29]. Many studies have tried to grade the echogenicity of bowel, however, the association with adverse pregnancy outcomes is strongest when the bowel is as echogenic as or more echogenic than bone [30]. To avoid confusion we recommend the use of a strict criteria for the diagnosis (echogenicity similar to the bone) in order reduce over diagnosis of echogenic bowel.

The prognosis of echogenic bowel depends mostly on whether or not it is associated with fetal abnormalities. One study showed that 34% of fetuses with echogenic bowel and IUGR or elevated maternal alpha fetoprotein have a poor perinatal outcome [31]. A larger study with 682 cases of echogenic bowel showed that 65.5% of the pregnancies resulted in the birth of a normal healthy newborn [32].

Protocol management suggested in the case of echogenic bowel:

1. Perform a detailed fetal anatomy scan to evaluate for other abnormalities.
2. Carefully examine the placenta and amniotic fluid to rule out an intraamniotic bleed.
3. Both parents should be tested for cystic fibrosis.
4. Maternal serologic testing for CMV and toxoplasmosis with both IgG and IgM, if amniocentesis is performed the PCR can be performed.
5. Amniocentesis is recommended even when present as an isolated finding.
6. If all are normal, we recommend strict follow-up for the detection of IUGR.
7. Antenatal surveillance with Doppler or BPP seems warranted due to possible association with fetal demise.

Club foot

Club foot (talipes equinovarus) is an abnormal relationship of the foot/ankle to the tibia and fibula. The foot is excessively planter flexed with the forefoot bent medially and the sole facing inward. It occurs in 0.1–0.4% of pregnancies and is bilateral 60% of the time. It is also more common in males.

The best way to detect it with US is when the long axis of foot is in same plane as the long axis of the tibia and fibula [33]. Club foot may occur in isolation or in association with numerous other conditions, such as general musculoskeletal disorders, arthrogryposis, genetic syndromes, neural tube defects and spine defects [33]. In 10–14% of cases it coexists with other structural malformations. First degree relatives of a person with idiopathic club foot are at a higher risk of having a club foot when compared with the general population [34]. The risk of a subsequent pregnancy being affected by club foot is 2% if
a previous male fetus was affected and 5% if the previous affected fetus was female [34].

In 6–22% of cases it is associated with aneuploidy, with trisomy 18 being the most common [35]. Most cases of club foot with concomitant chromosomal anomalies will demonstrate other structural abnormalities. Therefore chromosomal analysis is recommended upon evidence of additional structural abnormalities. In the absence of other structural anomalies, oligohydramnios, and IUGR isolated club foot has not been associated with adverse pregnancy outcomes. Postnatal successful surgery is obtained in 52–91% of cases enabling most children to participate in normal activities [34].

Protocol of management suggested in the case of club foot:
1. Sonographic detection of club foot warrants a detailed fetal anatomic survey, and fetal echocardiography should also be considered.
2. Amniocentesis should be recommended in all cases even though frequency of aneuploidy is low, because ultrasound may fail to detect subtle but significant associated abnormalities and the true frequency of aneuploidy is uncertain.

### Polydactyly

Polydactyly is the presence of an extra finger or toe. It is more common in Black women and two types are recognized: type A is when the extra digit is well developed, whereas in type B the extra digit is rudimentary with only soft tissue and no skeletal structure [36]. The extra digit can be preaxial (radial or tibial side) or postaxial (ulnar or fibular side).

Polydactyly may be present as part of a syndrome or as an isolated finding. It is sometimes familial, so family history is very helpful to exclude the association with other abnormalities. Chromosomal study may be offered if there is no familial history of polydactyly. Patients should be informed that fetuses with an isolated finding of polydactyly usually have a favorable outcome, however, parents should also be informed that at present it is not possible to definitely exclude the possibility of a rare anomaly, such as Bardet-Biedl syndrome [36].

Protocol of management suggested in the case of polydactyly:
1. Perform an anatomic survey, including a detailed examination for other limb abnormalities.
2. If it is an isolated finding and there is a reported family history of polydactyl then no further testing is necessary.
3. If it is an isolated finding and there is no family history reported then it is reasonable to offer a chromosomal analysis.
4. Parents should be counseled about rare possibilities such as Bardet-Biedl syndrome.

### Conclusion

Our increasing ability to detect soft markers with US early in gestation has led to confusion regarding the appropriate management and parental counseling. In this article we have used the best available literature to create a roadmap for these difficult situations. We hope that by following these protocols obstetricians and fetal medicine specialists will be able to more accurately counsel the parents regarding the long- and short-term prognosis of their fetus and direct them to the best evidence-based management.

Lastly, Dagklis et al. showed the presence of some soft markers (EF, echogenic bowel, and hydronephrosis) at 11–13 weeks increases the risk of aneuploidy even in the presence of a normal nuchal translucency [37]. Searching for and identifying these markers can alter the counseling and management of our patients. We believe the routine evaluation for the markers listed in this article can have a meaningful impact on the care we provide.

### References


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