Diagnosis and Counseling of Fetal Mild Ventriculomegaly

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ABSTRACT

Mild ventriculomegaly (MVM) is defined as a lateral ventricular diameter measured at the level of the atrium of ≥10 mm but <15 mm. Prenatal evaluation includes targeted sonographic examination for central nervous system (CNS) and extra-CNS abnormalities (present in 41.4% of the cases), and diagnostic amniocentesis for chromosomal analysis (3% of chromosomal abnormalities in isolated cases) and infectious disease studies (1.5% incidence). Individualized patient counseling is based on these test results. Optimal postnatal care involves appropriate pediatric neurologic and developmental specialists.

Learning objectives: After completion of this article, the reader will be able to define the normal appearance and size of the fetal cerebral ventricles, to list the conditions associated with MVM, and to counsel the parents properly.

Keywords: Fetus, Neurodevelopmental delay, Ultrasound, Ventriculomegaly

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Ventriculomegaly (VM) is a condition characterized by an enlargement of lateral ventricles of the developing fetal brain which affects 1 to 2 in 1,000 births.1 Fetal cerebral lateral ventricles are measured during fetal anomaly scan performed in the second trimester to screen for malformations. The measurement is done at the level of the atrium of the lateral ventricle filled by the echogenic choroid plexus, visible in an axial plane of the fetal brain showing also the frontal horns of the lateral ventricles and the cavum septi pellucidi. The calipers are positioned on the internal margin of the medial and lateral walls of the atria, at the level of the glomus of the choroid plexus, perpendicular to the long axis of the lateral ventricle.2,3

In order to standardize the correct technique of measurement, the following criteria should be fulfilled4:

- Midline structures should be equidistant from the proximal and distal calvarium margins
- The cavum septi pellucidi should be visualized as the anterior landmark and the ambient cistern as the posterior landmark
- The measurement should be performed opposite the internal parieto-occipital sulcus
- The measurement should be perpendicular to the inner and outer borders of the ventricle
- The measurement should be done on the inner edges of the ventricular walls.

An atrial width of less than 10 mm is considered normal (Fig. 1A). Ventriculomegaly is diagnosed when the width of one or both lateral ventricles, measured according to the criteria described so far, is ≥10 mm.

Measurements between ≥10 and <15 mm constitute mild VM (MVM) (Fig. 1B), while values ≥15 mm represent

Figs 1A and B: Correct technique of measurement of the atrial width: (A) normal ventricle; and (B) mild ventriculomegaly
severe VM. Mild ventriculomegaly is also defined as borderline VM. Some authors use the term “milder VM” and “moderate VM” to indicate measurements of 10 to 12 and 12.1 to 14.9 mm respectively. Other authors restrict the term of MVM to measurements between 10 and 12 mm.

Mild ventriculomegaly can be unilateral or bilateral, symmetrical or asymmetrical, isolated or associated with central nervous system (CNS) or extraneural anomalies, chromosomal abnormalities, congenital infections, and hemorrhages.

MILD VENTRICULOMEGALY ASSOCIATED WITH OTHER PATHOLOGIES

Neural and Extraneural Anomalies

Mild ventriculomegaly may be associated with neural and extraneural anomalies. For this reason, an accurate neurosonographic examination should be performed by experienced operators in a reference center. The percentage of associated anomalies ranges from 10 to 71% with an average value of 41.4%. As regards the CNS abnormalities, some of them may be easily recognized, such as open spinal defect or Dandy-Walker malformation. Subtle anomalies, such as dysgenesis of the corpus callosum (Fig. 2) or velum interpositum cyst or neuronal migration and proliferation disorders (Fig. 3) may require an accurate evaluation or may even be missed.

A recent meta-analysis shows that 7.4% of neural anomalies is not recognized at the time of the initial presentation, in contrast with the percentages previously reported in two different reviews (10–12% respectively). This lower percentage is mainly due to the improvement of ultrasound diagnostic capabilities and of operator training. The abnormalities which still are missed are mainly represented by neuronal migration and proliferation disorders which usually develop later in the third trimester. For this reason, magnetic resonance imaging (MRI) plays an important role in evaluating fetuses with MVM when this kind of anomalies are suspected.

Chromosomal Abnormalities

Ultrasound finding of MVM should raise the suspicion of chromosomal abnormalities, mainly trisomy 21, since several studies have demonstrated a percentage of association with abnormal karyotype ranging from 0 to 28.6% with an average value of 2.8%. A more recent review by Pagani et al reports a prevalence of abnormal karyotype of 4.7% in the whole cohort of VM, with a value of 8.3% in cases with associated anomalies and 3% in isolated MVM. This wide variation of results may depend on the prevalence of trisomies in the studied population. Given the strong association between MVM and chromosomal abnormalities, invasive testing for chromosomal analysis should be offered to all patients. However, the correlation between MVM and chromosomal abnormalities is still a cause of debate.

Congenital Infections

Congenital infections (toxoplasmosis, cytomegalovirus (CMV) and Rubella) can be a cause of MVM, with a
mean percentage of 1.5%. In fetuses with proven CMV infection, VM is the most common ultrasound finding, being present in 18% of cases. In the majority of cases, other sonographic signs may be observed in the fetal brain, such as small periventricular calcifications or small subependimal cysts (Fig. 4) and in extraneural structures, such as liver calcifications, ascites, hepatosplenomegaly, echogenic bowel, placentomegaly, growth restriction. When VM is diagnosed, screening for infections (usually TORCH) is recommended for the safety, simplicity, and relative low cost of execution of these tests.

In doubtful cases, MRI can provide important additional information with regard to subtle cerebral signs of infections, such as abnormal gyration, cerebellar hypoplasia, or abnormal signal in white matter.

**Feto-neonatal Alloimmune Thrombocytopenia**

Fetal and neonatal alloimmune thrombocytopenia is an alloimmune disorder resulting from platelet opsonization by maternal antibodies that destroy the fetal platelets. This rare condition implies a risk of intracranial hemorrhage and VM of about 10 to 30%. In these cases, VM is usually associated with other sonographic findings, such as hyperechogenicity of the ventricular walls or with the presence of intraventricular echogenic material, as a sign of the hemorrhage. For this reason, the search for anti-platelet antibodies is suggested when VM is associated with signs of intracranial hemorrhage.

**ISOLATED MILD VENTRICULOMEGALY**

Isolated mild ventriculomegaly (IMVM) is a diagnosis of exclusion, characterized by the sonographic absence of other associated malformations or markers of aneuploidy at the time of the initial presentation. The prevalence of IMVM is extremely variable and has been reported ranging from 0.15 to 0.7%. Isolated mild ventriculomegaly can involve one or both lateral ventricles with no significant differences in terms of neurodevelopment outcome. Moreover, some authors classify bilateral isolated forms into symmetrical and asymmetrical, defining the latter as difference in width of >2 mm. These cases seem to carry a worse neurological outcome.

Isolated mild ventriculomegaly remains stable during pregnancy in 55% of the cases, increases in 15.7%, and decreases in 34%. Fetuses with apparent IMVM should be sent to a referral center with the aim of performing a detailed anomaly scan and neurosonographic evaluation.

The timing and frequency of the follow-up depend on the gestational age and on the protocol used by the different centers. A minimal time before performing a follow-up after the first diagnosis is 2 weeks, since a shorter time interval would not allow to evidence the possible variation in the size of the ventricles.

Isolated mild ventriculomegaly may also be the first sign of brain anomalies recognizable only in the third trimester or even after delivery. For this reason, a MRI between 30 and 32 weeks of gestation should be suggested with the aim of adding information about cortical anomalies.

**NEURODEVELOPMENTAL DELAY IN IMVM**

The short- and medium-term postnatal outcomes of fetuses with IMVM show a slightly higher incidence of neurodevelopmental delay as compared with the normal population. Pagani et al report a prevalence of neurodevelopmental delay of 7.9%; this value is sensibly lower as compared with results of previous studies by Melchiorre et al and Devaseelan et al (10.9 and 12% respectively); however, it is just slightly increased in comparison with the 2 to 3% estimated for childhood disability in the general population.

The lower prevalence reported in the most recent review may depend on the improvement of sonographic
techniques that allow the prenatal diagnosis of anomalies associated with MVM, which in the past were often missed leading to a worse prognosis.

Lyall et al. analyzed the natural history of prenatal diagnosed IMVM in early childhood demonstrating a persistent enlargement of the lateral ventricle volume through 2 years of age. These children got a lower score on the fine motor and expressive language subscales of the Mullen Scale of Early Learning in comparison with the control group. In this way, the authors support the hypothesis that prenatal diagnosis of IMVM should be used as early biomarker of abnormal postnatal brain development.

It is very difficult to define with accuracy the neurological outcome in cases of IMVM because the literature shows conflicting results depending on the inadequate qualitative assessment and the different protocols used to assess postnatal neurological development, on the short period of follow-up planned, on the different ages of infants analyzed, and on the absence of distinction between mild, moderate, and severe delay. The evaluation of the neurodevelopmental delay in preschool children should include examination of locomotor activity, hearing, speech capacity, eye and hand coordination, and learning performance.

Although some authors suggested that other factors, such as female sex, asymmetrical bilateral IMVM, early presentation during the pregnancy, may worsen the neurological outcome, all available data prove that in IMVM the most important prognostic factors are the association with other abnormalities not detected at first examination and the progression of ventricular dilatation, both of which are retrospective diagnoses.

The neurodevelopmental delay in cases of isolated progressive VM is 16.7%; in this group of fetuses, there is also the highest incidence of chromosomal abnormalities (22.2%) and associated anomalies (71.4%); the overall abnormal outcome is 44.4%. From the above considerations, it is evident that in order to evaluate exactly the contribution of IMVM to the neurodevelopmental delay in infancy, a large collaborative prospective study using a unified protocol and long-term objective postnatal follow-up is needed.

COUNSELING

The sonographic finding of fetal MVM still represents a challenge in terms of management, counseling, prognosis, and postnatal neurological development because at the time of first diagnosis it is not always possible to rule out other associated anomalies, particularly in the late second trimester of pregnancy (false-negative rate of 13%). The fetuses affected by MVM are typically identified at 19 to 21 weeks, when neurogenesis and migration are not yet complete, with the consequence that migrational and gyration disorders could be too subtle to be detected with ultrasound. For this reason, during counseling the parents should be carefully informed about the limitations of ultrasound imaging in differentiating true IMVM from the associated type, and about the possibility that even IMVM can progress in 15.7% of the cases getting worse prognosis.

When MVM is associated with abnormal karyotype or infection, counseling is less problematic because the concerned cause is known, being able to provide a more accurate prognosis.

In cases associated with CNS anomalies, counseling depends on the type of abnormality carrying different neurological handicaps. However, there are some CNS anomalies, such as agenesis of corpus callosum whose neurological postnatal outcome is extremely variable, generating conflict decision in the parents.

It is established that the postnatal outcome and neural development are better in children with prenatal IMVM than in those with severe ventricular dilation or VM associated with CNS anomalies. In addition, some authors have suggested that the VM of 10 to 12 mm can be considered a normal variant especially when it resolves spontaneously. On the other hand, Lee et al. emphasize the concept of a prudent counseling even when ultrasound indicates that the VM is mild, isolated, and nonprogressive since the literature shows variable results in terms of neurodevelopmental outcome.

Prenatal imaging diagnosis is the pivot for counseling parents whose fetuses have been referred for VM. Because anomalies associated with MVM are not always detected during routine prenatal ultrasound screening, several studies have pointed out the potential role of MRI in the diagnosis of fetal cerebral anomalies and the information that MRI can provide as an adjunctive tool to sonography. In this regard, evidence suggests that the presence of dedicated neurosonography in an ultrasound unit is associated with high sensitivity and specificity for the detection of fetal CNS abnormalities compared with MRI, which however is performed later at 30 to 32 weeks of gestational age. In prenatal counseling, diagnosis of IMVM remains a dilemma: parents should be informed on the need to perform periodic and long-term follow-up extending up to 6 years in the postnatal period to allow early identification of attention deficit and hyperactivity disorders. A follow-up should be planned according to the diagnosis at birth by an expert pediatrician. Some authors suggest that MRI should be arranged after the age of 1 year to rule out lesions of the white matter that are not detectable during intrauterine or early postnatal life.
REFERENCES


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