Mid-Term Neurodevelopmental Outcome in Isolated Mild Ventriculomegaly Diagnosed in Fetal Life

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Abstract
Objective: To analyze mid-term neurodevelopmental outcome in children with isolated mild ventriculomegaly (VM) ≤ 12 mm diagnosed in fetal life, using the Battelle Developmental Inventory Screening Test (BDIST). Methods: 86 cases of mild VM were identified. 68 were excluded due to: other cerebral anomalies (n = 40), extra-cerebral anomalies (n = 3), chromosomal defects (n = 4), dysmorphic syndromes (n = 4), congenital infections (n = 2), termination of pregnancy (n = 9), stillbirth (n = 2) and incomplete follow-up (n = 4). 18 cases (range 1–8 years) of isolated mild VM were included for analysis. Seven neurodevelopmental domains were assessed by BDIST. Results: Routine neuropediatric evaluation detected neurological disorders in five children (28%; 3 with language impairment, one left hemiparesis and one intellectual retardation). BDIST showed some degree of neurodevelopmental delay in higher proportions: 66% in social-personal skills, 56% in gross motor skills, 39% in adaptive behavior and 28% in fine motor skills. Communicative and cognitive areas were the least affected (11 and 22% had moderate-to-severe involvement, respectively). A general trend towards worse outcomes was observed in the group of ≥ 4 years, although significant differences were only found for gross motor skills. Conclusion: Subtle neurological delays may appear during the infant period in fetuses prenatally diagnosed of isolated mild VM. In consequence, adequate measures should be established for early detection and treatment.

Introduction

Ventriculomegaly (VM) is the most common anomaly of the central nervous system diagnosed in fetal life. The prevalence of VM varies between 0.3 and 1.5 per 1,000 births, depending on the technique used for measuring, the evaluation of one or both lateral ventricles and gestational age at examination [1]. This prevalence is 20-fold increased in high-risk populations [2]. Up to 50% of cases are associated with other abnormalities (structural defects 33–61%, chromosomal aberrations 3–9% and congenital infection 5%) [3] and their detection is essential for proper prognosis assessment. This is more difficult when the VM presents in isolation. In fact, although
a relationship between the degree of ventricular dilatation and the neurological impact seems to have been established, the rate of psychomotor retardation in severe forms of VM (≥15 mm) varies widely among series, between 37.5 and 89% [4, 5].

Despite the fact that the majority of cases of fetal VM found in clinical practice fall into the mild category [6], the implications of isolated forms are poorly known, mainly due to the heterogeneity of the criteria used to define mild VM (10–15 vs. 10–12 mm) [6, 7]. Moreover, there is currently little available information about mid and long-term outcome of children diagnosed prenatally of mild VM and, in addition, follow-up studies are often incomplete and subjective [9, 10]. Therefore, prenatal assessment in these cases usually represents a conflicting situation both for parents and clinicians. This means that it is necessary to collect mid and long-term data on the neurological outcome of these children, in order to provide the parents the most accurate counseling following prenatal diagnosis of isolated mild VM. Furthermore, precocious identification of potential psychomotor deficits could allow an early implementation of effective intervention measures designed to minimize their impact on neurodevelopment.

The aim of this study is to analyze mid-term postnatal outcome in a cohort of infants with prenatally detected isolated mild VM analyzing the results provided both by the routine neuropediatric clinical assessment and by an additional structured interview to the parents.

Methods

Prenatal Issues

This is a retrospective study performed at a tertiary care referral center for prenatal diagnosis and management of fetal and neonatal pathology. The study was approved by the local ethics committee. Our database was queried for cases of mild VM prenatally diagnosed in the period from 2002 to 2008. Mild VM was defined as a transverse diameter of one or both ventricular atria between >10.0 and ≤12.0 mm [11]. All measurements of lateral ventricles were obtained according to current recommendations [12]. With these criteria, 86 cases were retrieved and analyzed.

Fetal neurosonography and a thorough sonographic evaluation of the rest of fetal morphology were performed in all cases by a fetal medicine specialist. All ongoing pregnancies underwent sonographic follow-up every 4–6 weeks in order to detect progression of the ventricular dilatation and/or associated defects of late onset. Karyotype analysis was offered in all cases: in 46 cases it was performed by amniocentesis (54%) and in 24 cases by fetal blood sampling (28%). In the rest, the karyotype was studied postnatally according to neonatal clinical findings. In all cases of apparently isolated VM, TORCH analysis was performed. In two cases, maternal seroconversion to cytomegalovirus was detected during pregnancy, and subsequent polymerase chain reaction amplification of cytomegalovirus DNA in amniotic fluid sample confirmed fetal infection. From 2006, we also included fetal magnetic resonance imaging (MRI) after the initial diagnosis of apparently isolated mild VM by neurosonography, in order to confirm the diagnosis and exclude associated CNS anomalies. Ultrasound findings, test results and pregnancy outcomes were entered in the database as soon as they became available.

Postnatal Issues

A detailed neonatal examination of all patients included in the study was conducted in the first week of life, including general and neurological clinical assessment and a cerebral transfonatal ultrasound (TF-US) in order to confirm the prenatal diagnosis. Postnatal MRI was indicated whenever there were concerns regarding the presence of associated defects after performing TF-US. All patients with isolated mild VM underwent clinical follow-up in an outpatient basis by a neonopediatrician specialist, according to the following policy: firstly, an additional TF-US was performed at the end of the first month of life in order to detect a possible progression of the VM. Secondly, a clinical evaluation including a complete physical and neurological examination, measurement of head circumference and psychomotor development assessment by checking if the chronological acquisition of the main neurological milestones are appropriated was performed at 1, 3, 6, 12, 18 and 24 months of age. Thirdly, TF-US was also performed in each visit as long as the fontanelles remained open. Finally, MRI was requested whenever there was progression of the VM or when the clinical examination revealed either psychomotor retardation or neurological deficits. If all these revisions displayed normal results, the infant was discharged from the neuropediatric office. Otherwise, the patient remained under specialized surveillance.

All cases in which cranial or extracranial associated defects, chromosomal abnormalities and/or congenital infections were detected either prenatally or postnatally were excluded. Similarly, terminated pregnancies, stillbirths and patients lost to follow-up were also excluded. Table 1 summarizes all cases of mild VM diagnosed in fetal life and reasons for exclusion from the study.

Information about neurological developmental status of patients included in the study was obtained consulting medical records. Psychomotor development was examined by a clinical psychologist using the Battelle Developmental Inventory Screening Test (BDIST). This questionnaire is the only test to examine pediatric neurodevelopment which has been validated to be applied as a screening tool in Spanish [13]. The BDIST has three administration formats (structured administration, observation, and interviews with parents) which can be used either in isolation or in a complementary manner to get the information [14]. Due to the fact that the majority of our patients were referred from distant areas of our country, we obtained all the data by means of telephone interview with the parents.

Briefly, BDIST is a widely used clinical tool for assessing key developmental skills in children from 6 months to 8 years of age. It is comprised of 96 age-specific items to evaluate the following five domains: personal-social skills, adaptive behavior, psychomotor ability (including fine motor and gross motor skills), communication, and cognition. Mean and standard deviations (SD) from the scores in each domain are calculated to classify patients either as normal, borderline (<1 SD below the mean), mild retard...
dation (1–2 SD) or severe retardation (>2 SD). Adverse neurological outcome is considered if the neurological examination is abnormal or if the results of BDIST are >1 SD [15].

BDIST was transversally applied to our patients at the end of 2009 when all had at least 12 months of life. The study population was subclassified in two groups according to the age at which BDIST was applied (<4 vs. ≥4 years) for further comparisons. We also analyzed the neurological impact of prenatal variables which might be associated with the outcome of isolated mild VM, such as gestational age at diagnosis, fetal sex, and unilateral or bilateral ventricular dilation.

**Statistical Analysis**

Continuous variables were represented as mean (SD) and categorical variables as percentages. Statistical significance (p < 0.05) was determined by Kruskal-Wallis test for continuous variables and by Fisher’s exact tests for categorical variables. Data were analyzed using SPSS software, version 13.0 (SPSS, Chicago, Ill., USA).

### Results

The study group included 18 liveborn fetuses that fulfilled all the inclusion criteria. This figure represents 21% of all mild VM prenatally diagnosed during the study period. All cases were incidental findings in low-risk pregnancies at the routine second or third trimester ultrasound scans. Their most important clinical data are summarized in table 2. At least one additional ultrasound examination was performed in all cases after the initial prenatal diagnosis, and the majority (15/18, 83%) had at least two additional scans. In these follow-up scans, the ventricular dilation remained stable in all fetuses. Postnatally, only one patient showed to have a progressive VM which became severe at 12 months of age and required ventriculo-peritoneal shunting.
Postnatal neurologic follow-up showed a completely normal outcome in most infants (13/18, 72.2%). Neurological disorders were detected in 5 patients, including 3 with mild-to-moderate language impairment requiring a speech therapist, 1 with a left hemiparesis mainly affecting the left hand and which was successfully treated with botulinum toxin and the infant with progressive VM, who suffers from severe intellectual retardation requiring supportive measures.

The mean age of the infants at the time of the neurodevelopment evaluation with BDIST was 4.9 years (range 1–8). BDIST showed some degree of psychomotor delay in a higher proportion of children than the routine clinical neurologic follow-up (table 3).

Disabilities affecting social-personal skills, which includes interaction with adults, expression of emotions, self-concept, peer interaction, and social role, were reported in two thirds of the patients, and in half of the cases this delay was moderate to severe. Similarly, a high proportion of disorders was also found in the adaptive behavior domain (7/18, 39%), which includes attention, eating, dressing, toileting and personal responsibility. One third of the infants ≥4 years showed to have a severe adaptive behavior disorder. Furthermore, a high rate of alterations was detected in the ‘gross motor skills’ section (those affecting the ability of children to carry out activities that require large muscles or groups of muscles) (10/18, 56%), although the 60% of them were minor deviations from normality. ‘Fine motor skills’ disabilities (those affecting the movements of small muscles that act in an organized and subtle fashion to accomplish delicate tasks) were less common (5/18, 28%), but all cases suffered from a moderate-to-severe impairment. Communicative and cognitive areas were the least affected and moderate-to-severe forms of involvement were reported in only 11 and 22% of our patients, respectively.

When comparing the proportion of abnormal findings in each domain between the group of children in whom the BDIST was applied before and after 4 years of age, a general trend towards worse outcomes can be observed in the group of ≥4 years. However, significant differences were only found in the ‘gross motor skills’ area (fig. 1).

We performed a further subanalysis to examine the relationship between the overall development according to BDIST and prognostic variables as gestational age at diagnosis (<22 weeks vs. ≥22 weeks), fetal sex (male vs. female) and laterality (unilateral vs. bilateral VM). This subanalysis showed no significant differences for these variable (p values: 0.65, 0.71 and 1.00, respectively), but in all cases diagnosed after 22 weeks (5/5), the overall development was inside the normal range.

**Discussion**

Prenatal diagnosis of isolated mild VM represents a considerable challenge when counseling about neurological outcome. Firstly, parents should be advised that this is a provisional diagnosis until associated malformations, congenital infections and chromosomal defects can be excluded. Fortunately, these abnormalities can be ruled out prenatally in most cases. The main dilemma relies on the fact that it is not possible to distinguish with certainty which fetuses with isolated mild VM will develop neurological sequelae postnatally. In this sense, fetal MRI...
for further evaluation of isolated mild VM in neurosonography has not been proved to add relevant information beyond the diagnostic confirmation, but this may be important to reassure the parents, specifically in the mid-second trimester of pregnancy. Notwithstanding, it is of paramount importance to provide the parents with the most accurate counseling about the risks of neurological impairment in their child.

Several studies have analyzed the neurological outcome of children with a prenatal diagnosis of isolated mild VM, but there is a wide variation in the reported results [4, 5]. These discrepancies are mainly due to heterogeneity in the definition of mild VM, the use of different methods of neurodevelopment assessment and the lack of distinction on the degree of delay [10]. Few studies have conducted mid or long-term follow-up using objective tests to assess the severity of delay in distinct areas of neurodevelopment. Our study restricted the diagnosis of mild VM to measurements of 10–12 mm, representing a spectrum of cases very close to the threshold of 10 mm, below which the ventricular measurement is considered normal by consensus [16]. It has been stated that ventricles between >12 and ≤15 mm should be studied separately as they are related to more unfavorable outcomes [17]. Although it has even been suggested that lateral ventricular atrium between 10 and 12 mm should be considered a variant of the norm [6], the postnatal neurologic follow-up of our cases revealed abnormal findings in 28% and intelective retardation in one child (5%). Our results correlate well with previous reports [18] and with a recent review which has reported abnormal neurological outcome in 34/288 (12%) of chromosomally normal infants presenting antenataly with an atrial width between 10 and 11.9 mm [7]. Therefore, this collection of results further demonstrates that isolated mild VM should be considered as a separate entity.

In addition, we explored the presence of subtle alterations in the psychomotor development of these children, which may be unnoticed by most parents who do not possess tools to detect non-obvious difficulties, and by pediatricians, who may not be able to observe them in a routine clinical examination. For that purpose, we applied a developmental screening test, as this tool has shown to improve the accuracy for identifying developmental delays in comparison with routine neuropsychiatric clinical evaluation. Certainly, the sensitivity and specificity of these screening instruments are usually reported between 70 and 90% [19].

Of all the tests available, we selected the BDIST for two main reasons. Firstly, it is a widely validated screening test in all its administration formats to be used in children with a wide range of ages between 6 months and 8 years [20], which fitted well with the ages of our patients. Secondly, this test allowed us to inquire into the parental perception about their child, without requiring moving the family from their diverse points of origin. Although we could not conduct a direct observation of the children, parents’ responses were sufficient to complete the test in all cases. Obviously, the scores obtained in this manner are merely indicative of the potential existence of neurodevelopmental delays, and the clinical significance of this information should be established using specific diagnostic tests applied by an expert.

Using this approach, we find out that the most common disabilities in these patients affect the gross motor skills. This is in accordance with two previous reports

Fig. 1. Distribution by group of age (<4 years vs. ≥4 years) of neurodevelopmental outcomes in the seven domains assessed by the Battelle Developmental Inventory Screening Test (BDIST).
Neurodevelopmental Outcome in Fetal Mild Ventriculomegaly

In conclusion, isolated mild VM (10–12 mm) in the prenatal period constitutes an easy-to-detect diagnosis but a difficult-to-assess condition. After excluding other associated abnormalities, information about postnatal outcome should be fundamentally reassuring as the risk of severe cognitive retardation is only about 5%. However, parents and pediatricians should be advised that mild-to-moderate delays in different areas of neurological development may appear during the infant period. Therefore, adequate screening strategies should be implemented to maximize the options of early detection of these delays, in order to establish early supportive measures and minimize their impact in the developing child.

References