Case Report of Dandy-Walker Malformation and Noonan Syndrome – Correlation of prenatal findings with neonatal evaluations

Caso Clínico de Malformação de Dandy-Walker e Síndrome de Noonan – Correlação dos achados pré-natais com avaliações neonatais

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Abstract

This article reports a case of Dandy-Walker malformation (DWM) and Noonan Syndrome (NS) diagnosed in a fetus at the second trimester. The association between DWM and NS is rare and prenatal diagnosis of DWM generally motivates a medical termination of pregnancy. This is the first clinical case reported of DWM associated with NS with PTPN11 gene mutations and this article allows to correlate prenatal findings with clinical and other neonatal evaluations. On the other hand, it highlights the importance of prenatal evaluation of DWM and it shows the best techniques in order to improve prenatal counseling.

Keywords: Dandy-Walker Malformation; Noonan Syndrome; Prenatal diagnosis.

Resumo

Este artigo apresenta um caso clínico de diagnóstico pré-natal, realizado no 2.º trimestre, de Malformação de Dandy-Walker (MDW) e Síndrome de Noonan (SN) no mesmo feto. A associação entre estas duas entidades é rara e o diagnóstico pré-natal de MDW geralmente leva a uma interrupção médica da gravidez. Posto isto, este é o primeiro caso descrito na literatura de diagnóstico pré-natal de MDW e SN associado a mutações no gene PTPN11, e que permite correlacionar achados pré-natais com a avaliação clínica e outros achados neonatais. Por outro lado, este artigo sublinha a importância do diagnóstico pré-natal de MDW e evidencia as melhores técnicas de forma a otimizar o aconselhamento pré-natal.

Palavras-chave: Malformação de Dandy-Walker; Síndrome de Noonan; Diagnóstico Pré-Natal.

INTRODUCTION

Dandy-Walker malformation (DWM) is a complex malformation involving the cerebellum and the posterior fossa, in which there is a developmental anomaly of the cerebellar vermis and a failure of the normal fourth ventricle closure¹,². This is a rare diagnosis, with an incidence of 1:30,000.³ Among the most common ultrasound findings, there is an enlargement of the posterior fossa with an upward displacement of the tentorium, a cystic dilatation of the fourth ventricle and a variable degree of hypoplasia or aplasia of the cerebellar vermis⁴. The diagnosis of vermis rotation and elevation of the tentorium can be facilitated by finding median planes (sagittal), which is easily obtained by transvaginal ultrasonography in cephalic presentations⁵. Con-
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andhydrocephalus, but the incidence of neurological malformations is unknown. However, NS rarely presents with evident neurologic manifestations and most adults have mild cognitive deficits. Prenatal features are nonspecific and prenatal diagnosis of NS is usually made when there is an affected parent or when there is an enlarged nuchal translucency in first trimester or a cystic hygroma in the second trimester.

The association between DWM and NS was only recently reported in one case of NS with the RRA S2 pathogenic variant p.Q72L. Another study reported a DWM diagnosis in one particular case with de novo missense mutations in PPP1CB, a disease with similar phenotype as NS with loose anagen hair.

CASE REPORT

A 22-year-old primigravida was referred to a prenatal consultation at 22 weeks of gestation due to the detection of a posterior fossa anomaly during her routine second-trimester scan. Her medical history was irrelevant and the male progenitor of the fetus and his mother had a diagnosis of NS, both with normal neurodevelopment.

There were no records of first trimester obstetric assessment and that was the first sonographic evaluation. The detailed transabdominal ultrasound revealed only anomalies confined to the posterior fossa of the brain. On
Genetic analysis was made by Array-comparative genomic hybridization (array-CGH), after amniocentesis, which didn't show any alteration. Type 1 NS, PTPN11 variant (missense variant c.5C>T(p.Thr2Ile) in heterozygosity in PTPN11 gene), the pathogenic variant identified in the affected family members, was investigated and the fetus had the same mutation. An MRI at 25 weeks confirmed the sonographic findings together with hypoplasia of the cerebellar hemispheres and an asymmetry of the lateral ventricles, the right one with 11.4mm. (Figure 3) Additionally, persistence of the left vena cava and a small ventricular septal defect were identified in fetal echocardiography. The severity of the brain malformation and the poor prognosis were explained to the couple in a multidisciplinary-joint consultation with Obstetrics, Pediatrics and Genetics. Medical interruption of the pregnancy was refused and parents chose to maintain pregnancy surveillance. Neurologic prenatal findings remained unaltered throughout the pregnancy and the fetal growth was in the 2nd/3rd percentile.

At 39 weeks, the primigravida was admitted to the hospital with a diagnosis of premature rupture of membranes. A cesarean section was performed by breech presentation and a female infant was born with 2290g and an Apgar index of 6/8/9 in the first, fifth and tenth minutes respectively. Physical examination of the neonate revealed a long forehead and dolichocephalic skull, with normal head circumference and weight and height below the 3rd percentile. Axial views, the cerebellar vermis was not identified and the posterior fossa was enlarged and it had a triangular shape. (Figure 1) Transvaginal ultrasound was performed in order to obtain sagittal planes and it revealed a hypoplasia of the cerebellar vermis, with rotation of the tentorium, with a BV angle greater than 45 degrees. These alterations were also seen in sagittal views at abdominal ultrasound at 30 weeks of gestation. (Figure 2)

![Figure 3](image-url)  
**FIGURE 3.** Cerebral MRI performed at 25 weeks of gestation. There is an agenesis of the inferior portion of the cerebellar vermis with a superior rotation of the superior portion of the vermis and the tentorium, together with hypoplasia of the cerebellar hemispheres and an asymmetry of the lateral ventricles, the right one with 11.4mm. A – Coronal view. B – Sagittal view.

![Figure 4](image-url)  
**FIGURE 4.** Postnatal cerebral MRI. It is evident a chronic supratentorial hydrocephalus, with reduced brainstem, an enlargement of the supratentorial ventricular system and a diffuse reduction of the white matter volume. A – Axial view. B – Coronal view. C – Sagittal view.
tile. Further evaluations revealed frontal erythema, sparse hair, frontal bossing, sparse eyebrows, hypertelorism, flattened nasal bridge with anteverted nostrils and low-set ears with posterior rotation. The prenatal findings were confirmed by postnatal cerebral MRI which additionally revealed chronic supratentorial hydrocephalus. (Figure 4) The head circumference was in the 85-97 percentile, with no indication for neurosurgery treatment. The infant is currently 15 months and she has an axial hypotonia, with no control of the pelvis, no strength in the lower limbs and poor weight development below the 3rd percentile. She is under hospital surveillance in Neurosurgery, Cardiology, Neurology, Otolaryngology, Physiatry and Genetic consultations and she was referred to a national association that was created with the purpose of qualification, rehabilitation and social integration of children with multiple disabilities.

DISCUSSION

This is the first reported clinical case of DWM associated with NS with PTPN11 mutations. Although the neurological development of the infants with NS syndrome is generally normal, DWM is associated with neurodevelopmental complications in 70% of cases, as indicated above. In this case, the association with DWM was the most likely reason for the poor neurological development.

This case highlights the importance of sagittal planes in neurosonographic evaluation, which allow the assessment of the cerebellar vermis with prognostic implications. In this way, sagittal planes are important to make differential diagnosis with other posterior fossa abnormalities, such as the persistent Blake's pouch cyst, an abnormality with better prognosis once the vermis has normal anatomy and size.

On the other hand, this case report allows to correlate prenatal findings with clinical and other neonatal evaluations, such as neonatal MRI. This is of special importance regarding that DWM generally motivates a medical termination of pregnancy and, for this reason, reports of neurological development are rare when it is performed a prenatal diagnosis.

REFERENCES


AUTHORS’ CONTRIBUTION
Author 1: Literature search, Design, Manuscript preparation, Data acquisition, Data analysis, Edition, Review
Author 2: Edition, Review
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Author 4: Data acquisition, Review
Author 5: Data acquisition, Review

DECLARATION OF PATIENT CONSENT
Written informed consent was obtained from the parents for publication of this case report.

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The authors have no conflicts of interest to declare.

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