Dandy-Walker Malformation and Down Syndrome Association: Good Developmental Outcome and Successful Endoscopic Treatment of Hydrocephalus

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Key Words
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Abstract
The association of Down syndrome (DS) with Dandy Walker malformation (DWM) is extremely rare, with only 3 cases reported to date. All cases reported have shown a bad life expectancy and a bad developmental outcome. The present case reveals the possibility of a good prognosis. A 19-month-old male patient had successful endoscopic hydrocephalus treatment and a good developmental outcome. He probably had a better outcome because of good DS and DWM prognostic parameters. Our patient suffered from a DWM with vermis identification of 2 fissures and 3 lobes and a DS with a well-preserved tonus, which was not associated with other congenital systemic defects. We may conclude that the prognosis of DS-DWM association may separately depend on the degree of clinical and neurological involvement of each malformation.

Introduction
Down syndrome is the most common genetic cause of mental retardation [1]. Its association with Dandy-Walker malformation is extremely rare, with only 3 cases reported to date.
Case Report

A 19-month-old male patient was referred to our neurosurgical unit because of hydrocephalus that caused progressive macrocraania. He was diagnosed with a prenatal ultrasonographic diagnosis of Dandy-Walker malformation (DWM) and a trisomy 21 karyotype. The neurological examination revealed a child with irritability, a cephalic perimeter of 53 cm and a bulging fontanel. He was active and reactive, mildly hypotonic, but still unable to walk independently or verbalize himself. His CT scan showed tetraventricular hydrocephalus, a posterior fossa cyst and an effacement of his cortical sulcus (fig. 1a, b). His MRI revealed a fourth ventricle dilatation, a hypoplastic cerebellar vermis with an upward displacement and an open aqueduct of Sylvius (fig. 2a). Third ventricle floor and lamina terminalis bulging on his MRI 3D-continuous-flow isotonic solution system sequence suggested the obstructive pattern of the hydrocephalus (fig. 2a), and an endoscopic third ventriculostomy (ETV) was performed. He had an uneventful postoperative outcome and was discharged on the third day. A 99mTc-DTPA ventriculocisternography [5] (injection of the radioisotope inside the lateral ventricle) performed after the ETV revealed a normal cerebrospinal circulation and absorption. A CT scan performed 7 months post-op showed a decreased ventricle size and a reappearance of the cortical sulcus (fig. 1c, d). An 18-month post-op MRI revealed the absence of the third ventricle floor and the lamina terminalis bulging, and the flow void changes in the floor of the third ventricle confirmed an ETV patency (fig. 2b, c). A systematic developmental exam based on the Denver II developmental scale [6] and a full neurological exam was performed at the age of 4 years and 10 months. Gross motor skills, fine motor skills, language and social domains were evaluated. His gross motor skills compared to when he was 3 years and 3 months old, his fine motor skills compared to when he was 3 years and 9 months old, his language was compared to when he was 3 years and 6 months old, and his social skills were those of a 3-year-old. The main qualitative difficulties in acquiring a fine motor performance was related to his deficits in coordination, which restrained the boy from activities such as brushing his teeth and getting dressed. The neurological examination revealed a slightly dysmetric motor pattern with bilateral and symmetrical intentional tremor. The parents referred to this as having only a small impact in their daily life activities. A unilateral right Babinski sign with no other showings of pyramidal dysfunction was verified. Cooperation and behavior were adequate. His tonus was well preserved, with a slightly decreased resistance to passive movement and relative joint hyperextensibility. The boy’s posture and strength were comparable with the expected developmental stage of Down syndrome (DS) patients at the same age. His gross motor function measure raw score was 96.1% and therefore compatible with the expected DS population score [7]. The cephalic perimeter was 55.5 cm.

Discussion

A retrospective study with a prenatal diagnosis of the fetal posterior fossa fluid showed that isolated DWM and vermian hypoplasia were associated with the normal developmental outcome in only 50% of cases [8]. A potentially good clinical and intellectual outcome has been demonstrated to occur in fetuses showing partial vermian agenesis, i.e., with the
identification of 2 fissures and 3 lobes and with an absence of associated intracranial and extracranial anomalies [9, 10]. ETV is considered to be an acceptable alternative for DWM hydrocephalus treatment, especially in older children, with the aim of reducing the shunt-related problems [11]. Cases with third ventricular bowing increase the ETV success rate [12]. DS developmental outcome is characterized by a weakness in communication, combined with daily living and social skills [13]. Delays in motor development of patients with DS appear to be particularly related to the degree of hypotony [14]. The combination of DS and DWN is a bad prognosis for affected subjects. The first patient died at the age of 2 weeks [3] and the second was severely handicapped at the age of 4 months [4]; the third patient, a 37-month-old male, was unable to crawl or ambulate independently [2]. The present case however had a better outcome, probably because of good prognostic parameters. Our patient suffers from DWM with a vermis identification of 2 fissures and 3 lobes (fig. 2c) and no associated intracranial or extracranial anomalies. Although a dysmetric motor profile was constructed, a possible consequence of the cerebellar malformation has a small impact in his daily life activities. His DS was not associated with any other congenital systemic defects and presents a well-preserved tonus and gross motor function scores comparable to isolated DS patients.

Conclusion

We conclude that the prognosis of the DS-DWM association may separately depend on the degree of clinical and neurological involvement of each malformation.

Disclosure Statement

The authors have no personal financial or institutional interest in any of the drugs, materials or devices mentioned in the article.

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Fig. 1. a, b Preoperative CT scan showing the tetraventricular hydrocephalus, posterior fossa cyst and the effacement of cortical sulci. c, d Postoperative CT scan showing a decreased ventricle size and a reappearance of cortical sulci.
Fig. 2. a Preoperative MRI on 3D-continuous-flow isotonic solution system sequence showing the posterior fossa cyst, cerebellar vermis hypoplasia, open aqueduct, the third ventricle floor and the lamina terminalis bulging. b Postoperative MRI on 3D-continuous-flow isotonic solution system sequence with the reversion of the third ventricle floor and the lamina terminalis bulging. c Postoperative MRI T2-weighted sequence showing the flow void signal on the third ventricle floor and the cerebellar vermis hypoplasia with the identification of 2 fissures and 3 lobes.