DANDY-WALKER SYNDROME IN ADULT MIMICKING MYASTHENIA GRAVIS

Juliana Cardoso, Marcos C. Lange, Paulo J. Lorenzoni, Rosana H. Scola, Lineu C. Werneck

ABSTRACT - The Dandy-Walker syndrome (DWS) is a rare posterior fossa malformation. The DWS can occur associated with other brain or systemic malformations, but ocular abnormalities in this disease are rare and clinical findings mimicking myasthenia gravis have not been described to date. We report a 23-year-old woman who presented mild limitation of the ocular movements with progressive palpebral ptosis, which changed in intensity during the day. The investigation showed negative anti-acetylcholine receptor antibody, repetitive nerve stimulation and “Tension test”, but the brain magnetic resonance image reveals DWS with hydrocephalus associated with calosal dysgenesis. The characteristic of disease, clinical manifestations and pathologic features, specially the clinical evaluation of ocular abnormalities in suspicion of DWS, including the MG in differential diagnosis are discussed.

KEY WORDS: Dandy-Walker syndrome, hydrocephalus, myasthenia gravis, palpebral ptosis.

The Dandy-Walker syndrome (DWS) is a rare posterior fossa malformation, characterized by aplasia or hypoplasia of the cerebellar vermis, involving the cortex and deep cerebellar nuclei; the presence of a membranous cyst of the fourth ventricle; and, commonly, hydrocephalus1,2. This malformation was first described by Dandy and Blackfan (1914)3. Taggart and Walker (1942) revised this malformation characterizing the main clinical and pathologic features4. Benda (1954) finally labeled the disease as “Dandy-Walker syndrome”5. Initially, the DWS was reported in Brazil by Almeida (1960)6.

The DWS can occur associated with other malformations of the brain, as calosal agenesis, and other systemic abnormalities7. Several other structured lesions of the central nervous system, such as cerebrovascularto disease, brain stem tumors and Chiari malformation were described as mimicking MG, due to involvement or compression of the brain stem, could easily lead to misdiagnosis and an inappropriate treatment, but the DWS have not been reported to date1,2,6-12.

The clinical presentation of dysfunction of the neuromuscular junction mimicked the DWS have not been reported to date and for this reason we report this case.

CASE
A 23-year-old woman presented mild limitation of the ocular movements with progressive palpebral ptosis, who changed in intensity from day to day and during the day, according to the intensity of physical activity, since the age of 19 years without diplopia. She was born at term and was the first child of non-consanguineous parents and the only...
affected case in the family. The psychomotor development was normal.

Physical examination did not reveal abnormalities. Neurological examination showed intellectually normal; asymmetrical palpebral ptosis (right>left) associated with bilateral ocular lateral rectus muscle weakness; saccades and pupils reflexes were preserved, absence of papilledema and nystagmus; generalized hypotonia; symmetrical muscle strength grade 4 (Medical Research Council Scale) in proximal upper limbs and grade 5 in lower limbs; deep tendon reflexes, gait and all sensory examinations were normal.

The symptoms and signs were thought to be consistent with myasthenia gravis, but the investigation showed the following results: (1) anti-acetylcholine receptor antibody normal (0.15 nmol; normal: <0.20 nmol); (2) needle electromyography, motor and sensory nerve conduction studies were normal; (3) repetitive stimulation of the facial, spinal accessory and ulnar nerves at 3 Hz were normal; and (4) "Tensilon test" (10 mg of edrophonium intravenous preceded by atropine) and use of pyridostigmine (180 mg/daily) did not improve the ocular palsies.

After this initial investigation the patient was submitted to magnetic resonance imaging (MRI) that revealed hydrocephalus, cerebellar vermis hypoplasia, posterior fossa cyst (large fourth ventricle) and dysgenesis of the corpus callosum (Figure).

All studies were done following informed consent.

**DISCUSSION**

The DWS is a very rare congenital malformation of the posterior fossa, with an incidence ranging between 1/25000 and 1/35000 births, usually observed during the prenatal period or early infancy and more rarely in adults1,2. The DWS is characterized by a hypoplasia or agenesis of the cerebellar vermis, involving the cortex and deep cerebellar nuclei, enlargement of the fourth ventricle in continuity with a posterior fossa cyst, and usually, though not exclusively, hydrocephalus3,5. There is some doubt if the malformation is simply due to congenital obstruction of the foramina of Luschka and Magendie, which results in ballooning of the fourth ventricle and deformity of the cerebellum of varying degree, or represents a structural developmental anomaly of the area of the fourth ventricle1,6,12.
Various predisposing factors have been reported such as infections, cranial trauma, chronically disturbance in cerebrospinal fluid pressure, persistence of embryonic tissue, vascular lesions, teratogenesis and maternal diabetes\textsuperscript{1,6,13}. The DWS has been frequently associated with trisomies 3q, 6p, 9p, 11 or 22, and more rarely with chromosome translocations\textsuperscript{13}. More recently, a first critical region associated with DWS, encompassing two adjacent Zinc finger in cerebellum genes, ZIC1 and ZIC4 has been identified\textsuperscript{13}.

In DWS patients focal neurological signs are usually less prominent, but symptoms and signs of hydrocephalus are more frequent\textsuperscript{1,6,12,14}. In most previous series of DWS, approximately 40% of the children were intellectually normal, 40% had mental retardation, and 20% were borderlines, but our case is intellectually normal\textsuperscript{6}. Seizures, cerebellar ataxia and dizziness, when present, become manifest during the first years of life\textsuperscript{1,6,13}. Symptoms suggestive of increased intracranial pressure such as lethargy, vomiting and nystagmus may also appear during the evolution\textsuperscript{1,13}. The associated neurodevelopmental abnormalities were also less severe in adults cases compared with those reported in early infancy\textsuperscript{13}. The patients with minor abnormalities degrees can remain asymptomatic until later in life and then present with hydrocephalus\textsuperscript{12-14}.

In this context, the preserved cortical cytoarchitecture and the rarity of additional neurodevelopmental changes in DWS adults may explain the mild or absence of clinical expression, compared with DWS infants\textsuperscript{13}. Structural imaging in eighth previously described adult cases also show a relative paucity of neurodevelopmental abnormalities\textsuperscript{13}.

The clinical manifestation of the DWS depending of the degree of the malformations and other brain or systemic abnormalities associated, but abnormal eye movements with palpebral ptosis has been rarely reported in DWS. In our case, bilateral abducens nerve palsy can be attributed to the hydrocephalus (traction on the abducens nerves as it is tethered at the petroclinoid ligament). Also, the palpebral ptosis can be attributed to the hydrocephalus, because in chronic cases a change in the cerebrospinal fluid pressure gradient, between the ventricles and the brain, can produce a transient disturbance of the periaqueductal structures (ventral), resulting in the development of palpebral ptosis\textsuperscript{15}. Lepore also reported a series of 13 patients with bilateral palpebral ptosis after acute right fronto-temporo-parietal lobe lesions\textsuperscript{16}. The changed of palpebral ptosis and lateral rectus weakness from day to day and according physical activity can be explained by hydrocephalus (change in the cerebrospinal fluid pressure).

The ocular abnormalities are found more frequent in MG and rarely in DWS, reason for which the initial diagnosis was MG in our patient.

Also, in agreement with the structural imaging data reported, this case suggests that the presence of chronic hydrocephalus in adult form of DWS may mimic dysfunction of the neuromuscular junction, as MG. This finding is important to remember to neurologists that patients with ocular form of the myasthenia gravis and normal investigation or atypical clinical characteristics must be submitted to brain images for differential diagnosis.

REFERENCES