Marchiafava-Bignami’s Disease, as Etiologic Diagnosis of Athetosis

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Abstract

The Marchiafava-Bignami disease, characterized by demyelination and necrosis of the corpus callosum, has typically been associated with chronic alcohol intake, and clinically has various symptoms and signs. However, several cases have been reported without alcohol association, and these – according to several publications – have some common points, such as preference for female, related to malnutrition, and radiological involvement of the splenium of the corpus callosum. We report a case of a patient with the characteristics described above and whose clinical manifestation was Athetosis. The authors associate this manifestation with the somatotopic distribution of the corpus callosum, and contribute to the etiologic diagnosis of Athetosis as a manifestation of the Marchiafava-Bignami disease, which has not been reported in the medical literature according to our review of the database.

Keywords
Marchiafava-Bignami disease · Athetosis · Corpus callosum

Case Report

A 34-year-old woman, with no previous morbid history, presented last year with a quick 1 week installation of restlessness followed by involuntary movements of hands and legs, predominantly in the upper extremities, which disappeared completely during sleep. She also complained of development of gait difficulty, falls, and minimal dysarthria. The severity of symptoms was constant over time. When interrogated about the previous year, a history of weight loss of 10 kg and an anemic syndrome...
that reverted with iron supplements and folic acid came to our knowledge. On physical examination, athetosis was found predominantly in the upper extremities, also hypotonia and hyper-reflexia with an increase of the reflexogenic area in the 4 limbs. The complete blood count, liver biochemical profile, syphilis, VIH test, and serum ceruloplasmin were within normal ranges and/or negative.

The study was completed with CT (Fig. 1), which showed a hypodense region in the splenium of the corpus callosum, and nuclear MRI was compatible MBD (Fig. 2).

By ethical considerations, the patient was asked to consent to publish her clinical case in a medical journal, with affirmative answer.

**Analysis**

Prior to the implementation of imaging studies, the diagnosis of MBD was done based on postmortem pathological findings, because the manifestations were diverse and no clinical diagnostic criteria were defined for the disease [4]. Currently, the CT images and MRI allow “in vivo” diagnosis of this condition [5], allowing the initiation of early treatment in an attempt to achieve an improvement in the living conditions of patients.

The classic features described for imaging in MRI diagnosis of MBD are diffuse atrophy of the corpus callosum and areas of focal necrosis, which present as hypointensity on T1 and hyperintensity on T2 [6]. Most of the findings are distributed throughout the corpus callosum (splenium, body, and knee), and injuries seen in the acute phase usually persist in the chronic phase and are often described as gadolinium uptake, without the latter being an exclusive criterion [7].

Among the reported cases of non-alcoholic MBD, evaluated by MRI, none presented with gadolinium contrast enhancement. In turn, most of them showed selective compromise of the splenium of the corpus callosum [8]. So far both findings could be described as common characteristics of MRI in non-alcoholic MBD.

With the advancement of functional imaging studies, it has been shown that the corpus callosum has a topographic distribution of different functions that connect from one hemisphere to another [9]. Wahl et al. [10] showed that motor human corpus callosum fibers have a somatotopic organization, where hand motor fibers are located above the feet. Although it is an initial approximation to the study of coordination of interhemispheric movement, in this series, 2 of 12 cases presented a variety of this distribution, finding the feet fibers in front of the hands fibers. Greater accuracy of functional localization of the corpus callosum could eventually help correlate the clinical manifestations of the MBD.

Common features of non-alcoholic MBD – according to a recent review – are female patients, young population, and those associated with malnutrition [7], all of which were present in our patients, as well as the imaging study which shows selective engagement of the splenium of corpus callosum, also described in other cases of the disease without alcohol involvement [8].

In our case report, the patient did show abnormal involuntary movements of hands and feet, consistent with athetosis. Subsequently, the CT and MR imaging findings presented MBD. So we postulate MBD as the possible etiology of acquired athetosis.

The term athetosis, comes from a Greek word meaning “without attachment” or “changing.” The disease is characterized by the inability to maintain a single position in the fingers and toes, tongue or any other part of the body where no muscle group is respected and movements are slower than in chorea, and with distal predominance [11]. This seems to be related to a failure of the striatum to suppress the activity of unwanted muscle groups [3]. Among the etiologies of athetosis, are postnatal congenital hypoxia, head trauma, and perinatal ker-

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**Fig. 1.** TC showing hypodense corpus callosum splenium.
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With icterus; with neuronal loss and demyelination that affect the basal ganglia in the latter [3, 12]. In adults, other causes of athetosis have been described, such as hepatic encephalopathy, Wilson disease, Hallervorden-Spatz Syndrome, Leigh disease, and mitochondrial disease [3]. While there is no direct evidence that MBD may be an etiological cause of athetosis, there are reports of MBD with impaired basal ganglia structures, athetoid cerebral palsy and multiple subcortical lesions that include the corpus callosum; this, despite not being compromised in classic description of both diseases [13, 14]. Histopathological studies of Lechevalier et al. [13] in 1977 described injuries in caudate putamen and interhemispheric disconnection in a patient with clinical MBD. Yoshida et al. [14] evaluated the anatomy of the gray and white matter in children with cerebral palsy and athetoid characteristics with MRI findings, and described a case of slight thinning of the corpus callosum. As the somatotopic distribution of motor fibers in the callosum of hands and toes are next to each other, as demonstrated in studies of functional imaging [10]; this could explain that any damage located in that area could compromise the movement of hands and feet as in athetosis.

Conclusions

From the case report and review of literature, it can be concluded that in the low frequency of non-alcoholic MBD, this presented as common characteristics: is most common in female patients, young people, those associated with malnutrition, and the studies by MRI without gadolinium contrast enhancement show a preferential involvement of the splenium of the callosum corpus.

Our patient presented as an acquired athetosis; without other possible causes of extrapyramidal disease; with neuroimaging compatible with MBD, and after a review of the literature, with an etiological relationship to MBD which has not been previously described in literature.

An involvement of basal ganglia in MBD, and damage to the corpus callosum in athetoid cerebral palsy has been reported. Also, there is evidence of a somatotopic distribution in the corpus callosum described as an adjacent location of fibers of coordination and toes movement in the corpus callosum, which could explain that any alteration in that area could produce athetosis. Therefore, one cannot rule out the possibility that the MBD can be a cause of acquired athetosis and there is a need for more studies or other reports to assess the direct relationship between MBD and athetosis.

Authorship Contribution

A.V.C. involved in the conception, design, acquisition of data, analysis, interpretation of data, and final approval. Critically revising, analysis and final approval done by M.R.L. and R.G.T. conception, design, acquisition of data, analysis, interpretation of data done by M.F.F.C.

Disclosure Statement

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