Lhermitte-Duclos Disease and Cowden Syndrome: A Case Report and Literature Review

Doença de Lhermitte-Duclos e Síndrome de Cowden: relato de caso e revisão da literatura

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Keywords
► Lhermitte-Duclos disease
► Cowden syndrome
► hamartoma syndrome
► cerebellum

Abstract
Lhermitte-Duclos disease (LDD), also known as dysplastic gangliocytoma of the cerebellum, is a rare, usually benign, slow-growing tumor, that commonly affects patients aged 30 to 50 years-old. The manifestations of dysplastic cerebellar gangliocytoma are nonspecific and are related both to the mass effect produced by its growth and to the location of the lesion. Cerebellar symptoms such as ataxia are often present. In 40% of cases, the tumor is associated with Cowden syndrome, which is part of a group of genetic disorders called polyoid hamartoma complex. In this case report, the patient presented expansive lesion in the posterior fossa, compatible with LDD, associated with macrocephaly. These findings are considered major criteria for Cowden syndrome. When together, they confirm the diagnoses. To our knowledge, this is the first report of the association of LDD and Cowden syndrome in Brazil.

Palavras-chave
► doença de Lhermitte-Duclos
► doença de Cowden
► síndrome do hamartoma múltiplo
► cerebelo

Resumo
A doença de Lhermitte-Duclos (DLD), também conhecida como gangliocitoma displásico do cerebelo, é um tumor raro, geralmente benigno e de lento crescimento, que geralmente afeta pacientes entre 30 e 50 anos. Suas manifestações são inespecíficas e se relacionam ao efeito de massa produzido por seu crescimento e pela localização da lesão. Comumente, observam-se sintomas cerebelares, como ataxia, dismetria e disdiadococinesia. Em 40% dos casos, a doença encontra-se associada à síndrome de Cowden, a qual faz parte de um grupo de enfermidades genéticas chamado complexo do hamartoma polipoide. No caso relatado, o paciente apresentou lesão expansiva em fossa posterior compatível com DLD, associada à macrocefalia. Esses achados constituem dois critérios maiores, os quais, em conjunto, determinam o diagnóstico de Síndrome de Cowden. Este é o primeiro relato da associação entre DLD e Síndrome de Cowden no Brasil.
Introduction

Lhermitte-Duclos disease (LDD), also known as dysplastic gangliocytoma of the cerebellum, is a rare condition. Only ~300 cases have been reported in the literature since its description in 1920.\textsuperscript{1–3} This tumor, which is composed by atypical ganglion cells, is considered benign and slow-growing.\textsuperscript{4,5} It usually presents in patients aged between 30 and 50 years. The non-specific clinical picture is related to the location of the tumor and to the mass effect produced by its growth. Therefore, the most common symptoms result from cerebellar involvement. Headache, motor disorders, intracranial hypertension, and ataxia, as well as others, are frequently observed.

The crucial point of LDD is the possibility of association with Cowden syndrome (CS), which participates in a group of genetic syndromes called polypoid hamartoma syndrome.\textsuperscript{6} This rare and autosomal dominant disorder is characterized by the presence of multiple hamartomas, which confers an increased risk to developing other neoplasms. The features of CS include benign mucocutaneous lesions, such as trichilemmomas, acral keratoses, and papillomatous lesions, in addition to LDD—which was recognized as a major criterion for Cowden syndrome by the International Cowden Consortium Criteria, in 2004.\textsuperscript{7,8}

As far as we know, there is no report of LDD associated with Cowden syndrome in Brazil. In the present work, we not only report this condition but also present the findings of our systematic review of the literature.

Case Report

A 26-year-old male, living in Manaus, Amazonas, presented at our hospital with progressive unsteadiness. At the moment of admission, he was unable to walk. The patient also reported persistent and gradually increasing headache. Neurological examination revealed preserved motor strength, as well as an ataxic gait, dysdiadochokinesia, hypermetria, and dysarthria. A left nystagmus was further noticed.

Computed tomography (CT) of the skull showed a hypodense lesion in the posterior fossa on the right cerebellar hemisphere. A T1 gadolinium-enhanced magnetic resonance imaging (MRI) demonstrated a lobulated non-enhanced mass lesion, characterized by alternated linear hyposignal and isosignal areas.

In T2-weighted sequences, the hyposignal and hypersignal bands formed a “tiger-striped” pattern, an aspect classically described in cases of LDD. The lesion compressed the fourth ventricle but did not determine hydrocephalus (\textsuperscript{►}Fig. 1).

In light of the suspected diagnosis, it became valid to investigate the possibility of CS. A thorough physical examination revealed macrocephaly (62 cm of circumference, 95th percentile), as well as asymptomatic papillomatous lesions on the penis glans. Sexually transmitted diseases were discarded.

After diagnosis, surgical excision of the tumor through right suboccipital craniotomy was performed. Following dural opening, the cerebrospinal fluid was drained next to the cisterna magna, and we performed a small corticectomy. The non-bleeding hardened lesion resection was subtotal,

\textbf{Fig. 1} Magnetic resonance imaging shows a non-contrasting expansive lesion in the right cerebellar hemisphere, with lamellar areas of intense hyposignal and isosignal in T1 with gadolinium (A). In T2 (B), the hyposignal and hypersignal bands reveal the striated pattern, referred to as “tiger-striped” sign, which is classically described in the cases of Lhermitte-Duclos disease.
due to the absence of cleavage plane and difficult differentiation from the normal tissue.

In the histopathological study, proliferation of polygonal cells with pleomorphic nuclei, and broad and eosinophilic cytoplasm associated with vacuoles was observed. Hypertrophy of the cerebellar granular layer compatible with the diagnosis of dysplastic gangliocytoma of the cerebellum was further noticed. (Fig. 2)

After treatment, the patient presented significant clinical improvement, with no cerebellar complaints and reduction of the headache. He has now completed 12 months of postoperative follow-up.

Discussion

For the literature review, we used the Medline and Lilacs databases to search relevant articles from the last review of the literature on the subject from 2006 until March 2019. We use the terms Lhermitte-Duclos disease, Lhermitte-Duclos disease, Dysplastic gangliocytoma Cowden Disease, Dysplastic Gangliocytoma of the Cerebellum, and Cowden Syndrome. We identified 136 articles, and, of these, most were case reports (77 articles), and 2 articles were case series. So far, there are ~300 cases published in the literature. Of these, 5 reports are Brazilian. Therefore, this article is the 6th report of LDD and the first to report the association with CS in Brazil.

First described in the 1920s, there still is considerable controversy concerning the origin of LDD. It is known, however, that LDD is considered a major criterion of CS, an autosomal dominant genetic disorder.

Lhermitte-Duclos disease, or dysplastic gangliocytoma of the cerebellum, is a slow growing lesion of the cerebellum cortex, and it is usually diagnosed between the third and fourth decade of life, as identified in the present case, with no gender predominance. The expansive effect, associated with obstruction of cerebrospinal fluid flow, may result in noncommunicating hydrocephalus and intracranial hypertension. In addition, cerebellar symptoms, as in the case described, and cranial nerves involvement may occur.

The imaging examination is of great value during the diagnostic investigation of LDD. Although CT has significant limitations—showing a hypodense image without contrast enhancement—, the MRI associated with diffusion sequence and spectroscopy, shows a classic expansive lesion with hypo-

signal on T1 and hypersignal on T2-weighted sequences. Another feature observed in T2-sequences is the characteristic “tiger-striped” pattern. These changes result from both the white matter atrophy and the granular cell layer thickening.

The preoperative diagnosis is given by the tiger striped pattern present in T2-weighted MRI scans, associated with expansive cerebellar lesion affecting a single hemisphere. The lesion is characterized by hyperdense parallel grooves that occur due to cerebellar foliation thickening secondary to cortical cell enlargement, as well as groove dysplasia, which are considered to be practically pathognomonic radiological signs of LDD.

Cowden syndrome, characterized by Lloyd and Dennis in 1963, should be investigated in patients diagnosed with LDD. The syndrome is a genetic disorder due to a mutation in the phosphatase and homologous tensin (PTEN) gene or in its promoter region. The disorder affects the PTEN gene in the 10q23.2 locus in 80% of the patients. In addition, most LDD patients have a deletion-type mutation of the PTEN gene, resulting in abnormal growth of granular cells. The clinical diagnosis of this condition is based on the major and minor criteria described below (Table 1). The pathognomonic CS criteria are: mucocutaneous lesions (facial trichilemmomas, papillomatous lesions, acral keratosis and mucosal lesions). The major criteria include: breast cancer, thyroid carcinoma (especially follicular type), macrocephaly (occipital-frontal circumference ≥ 97th percentile), endometrial carcinoma and LDD. Minor criteria include: other thyroid lesions, decreased intellectual capacity (IQ ≤ 75), hamartomatous intestinal polyps, fibrocystic breast disease, lipomas, fibroids, tumors of the genitourinary tract (mainly renal cell carcinoma), genitourinary malformation, and uterine myoma. Therefore, the patient reported in the present case presents CS since it has 2 major criteria (LDD and macrocephaly), which define the diagnosis. The importance of the association relies on the fact that this disease determines a high risk of both benign and malignant neoplasms development. The thyroid, breast and endometrium are the main organs affected by the aforementioned neoplasms. Therefore, those patients should be closely monitored by a multidisciplinary team for surveillance of possible neoplasms that may occur.

The definitive treatment of LDD is surgical resection with decompression of the posterior fossa by total or subtotal tumor removal. Complete resection of the tumor is difficult due to the

![Fig. 2](image) Histopathological examination of the tumor (hematoxylin and eosin staining, original magnification of 200x). A and B: Polygonal cells with pleomorphic and cytoplasmic broad, eosinophilic nuclei, placed diffusely associated with vacuoles. C: normal glial cells of the same patient.
impossibility of clearly defining its margins.19,20 In the largest series published to date, Jiang et al19 evaluated 18 patients with LDD. Complete resection was possible in 9 of the 17 operated patients, and there was no recurrence in the late follow-up. Cowden syndrome was diagnosed in 11 of these 18 patients. Despite the degree of resection (partial or subtotal), the outcome is favorable, and recurrence is rare.19,20

In the second largest published series, Wang et al20 evaluated 12 patients, and a complete resection was achieved in 3 patients. Only one recurrence was reported. In cases of asymptomatic patients incidentally diagnosed by MRI, conservative management may also be assumed.21,22 The real effectiveness of posterior fossa irradiation therapy is unknown. However, assuming the non-neoplastic etiology of LDD, irradiation is unlikely beneficial, and, therefore, it is not recommended—even in subtotal resections.19,23

Moreover, in the histopathological examination, which is the gold standard for diagnosis, the white mass atrophy and the granular cell layer thickening can be observed.2,6 The absence of the Purkinje, the laminar cytoarchitecture destruction of the cerebellar cortex, and the presence of hypertrophic and dysplastic neurons in the internal granular layer are common findings. On the other hand, the absence of mitotic activity, necrosis, and endothelial proliferation indicate the benign nature of the lesion.

### Conclusion

Lhermitte-Duclos disease is considered a rare cause of progressive headache, associated with cerebellar symptoms. When it comes to preoperative diagnosis, the T2-weighted MRI, which can demonstrate the classical “tiger-striped” pattern, is the preferential exam. The definitive diagnosis, nonetheless, is histopathological, and it results from the demonstration of a laminar cytoarchitecture destruction, as well as from the presence of hypertrophic and dysplastic neurons in the internal granular layer, and the absence of the Purkinje cell layer. The treatment can be either surgical or not. While tumor resection is recommended for all symptomatic patients, a conservative management may be considered for incidentally diagnosed asymptomatic patients. Physicians must remember to search the association with CS, which justifies a regular follow-up of the patient due to the risk of developing malignant lesions in other organs, such as the breast, the colon, and the thyroid.

### Conflict of Interest

The authors declare that there is no conflict of interest.

### References