Lhermitte-Duclos Disease: Literature Review and Novel Treatment Strategy

Sagun Tuli, John P. Proviast and Mark Bernstein

ABSTRACT: Background: Lhermitte-Duclos disease (LDD) is a rare pathologic entity involving the cerebellum. The fundamental nature of the entity and its pathogenesis remain unknown, and considerable debate has centered on whether it represents a neoplastic, malformative or hamartomatous lesion. The cell or cells of origin remain incompletely defined. Previous reports in the English literature have dealt predominantly with the clinical and pathological aspects yet few address issues of treatment. Methods: A case of Lhermitte-Duclos disease (LDD) in a 54-year-old female leading to local compressive symptoms and obstructive hydrocephalus is presented. A craniectomy, in addition to a Cl laminectomy followed by a decompressive duroplasty (using autologous fascia lata graft) was performed. Results: The patient clinically improved and follow-up MRI 11 months post-operatively revealed improvement in hydrocephalus. Conclusion: The histological and immunohistochemical features of the lesion are described, emphasizing the role of an abnormal dysplastic granule cell layer. The evidence in favor of each of the major theories of pathogenesis, malformative and neoplastic is discussed. Based on these facts a form of surgical intervention involving decompressive duroplasty is proposed.

RÉSUMÉ: Maladie de Lhermitte-Duclos: revue de la littérature et nouveau traitement. Introduction: La maladie de Lhermitte-Duclos est une pathologie rare impliquant le cervelet. La nature de la maladie et sa pathogénèse demeurent inconnues. On s’interroge toujours à savoir si la lésion est une néoplasie, une malformation ou un hamartome. La ou les cellules d’origine demeurent mal définies. Jusqu’à maintenant dans la littérature anglaise on s’est intéressé surtout aux aspects cliniques et anatomopathologiques et peu aux aspects thérapeutiques. Méthodes: Nous présentons un cas de maladie de Lhermitte-Duclos (MLD) qui a entrainé des symptômes de compression locale et d’hydrocéphalé obstructive chez une femme de 54 ans. Nous avons procédé à une craniectomie et à une laminectomie au niveau de Cl suivies d’une duroplastie de décompression (au moyen d’une greffe autologue du fascia lata). Résultats: La patiente s’est améliorée cliniquement et la résonance magnétique effectuée 11 mois après l’intervention a montré une amélioration de l’hydrocéphalé. Conclusions: Nous décrivons les caractéristiques histologiques et immunohistochimiques de la lésion tout en insistant sur le rôle d’une couche anormale de cellules microgliales dysplasiques. Nous discutons des observations qui sont en faveur de chacune des théories sur la pathogénèse. En nous basant sur ces faits, nous proposons une forme unique d’intervention chirurgicale impliquant une duroplastie de décompression.


Lhermitte-Duclos disease (LDD) is a pathologic entity involving the cerebellum. The fundamental nature of the entity and its pathogenesis remain unknown, and considerable debate has centered on whether it represents a neoplastic, malformative or hamartomatous lesion. The cell or cells of origin remain incompletely defined although it is generally felt that an abnormality of the granule cells plays a prominent role. Whether Purkinje cells and/or other normal cellular elements of the cerebellar cortex are involved remains to be defined. Reflecting this confusion and limited understanding, numerous descriptive names have appeared in the literature including cerebellar hamartoma, neurocytic blastoma, hamartoblastoma, dysplastic gangliocytoma, Purkinjeoma, Lhermitte-Duclos, gangliocytoma myelinicum diffusum, benign hypertrophy of the cerebellum, and finally diffuse ganglioneuroma of the cerebellar cortex. Each reflects the possible cell of origin, but since that issue is as yet unresolved most refer to it as Lhermitte-Duclos disease (LDD). The first case report by Lhermitte and Duclos in 1920 used the descriptive term “diffuse ganglioneuroma of the cerebellar cortex.” Subsequent papers have added case reports and small series. These have dealt predominantly with clinical and pathologic aspects including immunohistochemistry, ultrastructure and proliferative aspects. By contrast, there are very few papers addressing issues of treatment, the specifics of surgical therapy and their impact on the natural history of the disease.
Case Report

Clinical History

A 54-year-old married female of Romanian descent presented with impairment of gait. The previously healthy patient described imbalance precipitating a number of falls over an 18 month duration. She also attested to episodic pain in the occipital and cervical region.

Physical Examination

Bilateral appendicular dysmetria in addition to truncal ataxia was exhibited. Unsteadiness of gait was also noted. The patient was otherwise neurologically intact.

Imaging

Plain computed tomographic (CT) scan revealed a homogeneous nonenhancing region of low attenuation in the left cerebellum extending to the vermis. Obstructive hydrocephalus was presumably a result of fourth ventricular compression by the lesion.

Figure 1: (a) T1 weighted image reveals a low intensity lesion consisting of prominent septations (arrow) and producing mass effect. Secondary compression of the fourth ventricle and hydrocephalus prevails.

Figure 1: (b) Increased signal intensity on a T2 weighted image reveals a well circumscribed vermian lesion on the right side (arrow) along with a separate lesion involving the left cerebellar cortex (arrows).

Magnetic resonance imaging, (T1 and T2 weighted images) revealed a two part lesion; a well circumscribed vermian lesion on the right side along with a separate lesion involving the left cerebellar cortex. Each consisted of prominent septations and produced mass effect, with secondary compression of the fourth ventricle and hydrocephalus (Figure 1a and b). The areas were of low signal density on T1 weighted (TR 516, TE 11/Fr) images and increased signal density on T2 weighted imaging (TR 4000, TE 90/Ef) and on proton density (TR 750, TE 12/Fr). No enhancement was noted following intravenous administration of gadolinium.

Operative course

A left occipital craniectomy followed by resection of the posterior arch of C1 was performed. A “Y” shaped dural opening revealed enlarged midline vermian tissue of white discoloration along with a separate lesion in the left cerebellar hemisphere. The latter was composed of a poorly demarcated area consisting of obliquely aligned enlarged white gyri. Intraoperative biopsies were obtained using tissue forceps, for pathological evaluation. The site of dural opening was closed using an autologous fascia lata graft. The bone flap was not replaced. This allowed for adequate bony and dural decompression.

Post-operative course

The patient clinically improved with some amelioration in appendicular and truncal ataxia. Minimal unsteadiness of gait persisted. Follow-up MRI 11 months post-operatively revealed improvement in hydrocephalus.

Pathology

Pathologic examination showed multiple small fragments of predominantly abnormal cerebellar cortex. In most of the specimen the normal cortical architecture was altered, the chief abnormality being loss of the normal granule cell layer. This was replaced by a broader zone of altered dysplastic cells with a neuronal immuno-phenotype, all of which were larger than the normal granule cells (Figure 2a and b). The dysplastic cells were generally round, of variable size, with a rare cluster of larger ganglionic type cells (Figure 2c). In these areas there was loss of the normal Purkinje cell layer. The overlying molecular zone was abnormal containing a prominent plexus of myelinated axons (Figure 2d). Small areas of the specimen showed transition to a normal cerebellar architecture.

The immunohistochemical profile showed strong positivity of the dysplastic cells with antisera to neuron specific enolase (Figure 2e). There were multiple areas of patchy positivity, particularly in the surrounding extracellular neuropil, with antisera to synaptophysin. Neurofilament was largely unreactive with an antibody to the 70 and 200 kd subunits (DAKO 2F-11 monoclonal); only the occasional larger dysplastic cell showed some cytoplasmic positivity. Glial fibrillary acidic protein (GFAP) was unreactive in the dysplastic cells although...
Figure 2b: Dysplastic granule cell layer of LD lesion. Note the slightly larger more pleomorphic cells (thin arrow) with lighter nuclear chromatin. A larger cell with a more neuronal phenotype is present (thick arrow). (Hematoxylin and eosin 250x.)

Figure 2c: Cluster of larger ganglionic type cells (arrows) composing a small percentage of the lesion. (Hematoxylin and eosin 250x.)

Figure 2d: Plexus of abnormal myelinated axons in the "molecular zone" of the LD cortex (arrows). (Hematoxylin and eosin 150x.)

Figure 2e: Immunohistochemistry showing strong cytoplasmic positivity of dysplastic cells for neuron specific enolase, indicating a neuronal like phenotype. (IHC, ABC methods 250x.)

the superficial aspect of cortex contained a network of radial glial fibres which were GFAP positive. p53 and MIB-1 immuno-histochemistry showed no nuclear immunolabelling of the dysplastic cells.

DISCUSSION

LDD is clinically manifest as a slowly growing mass in the cerebellar hemispheres. Upon review of the English literature of patients (i.e., non-autopsy cases) with LDD common features are those of increased intracranial pressure, with 80% (40/50) of patients presenting with headaches, 30% with nausea and/or vomiting, 42% with papilledema, and 24% with diplopia. Signs of cerebellar dysfunction were apparent in a lesser number of patients, with ataxia present in 60%, dysmetria/dysdiadochokinesis in 32%, hypotonia in 8%, and tremors in 10%. Less frequently symptoms of long tract impairment (motor dysfunction, spasticity, hyperreflexia, Babinski reflex) and cranial nerve dysfunction (dysphagia, facial numbness, tinnitus, hearing impairment, vertigo, dysarthria) are present. Unusual cases of orthostatic hypotension and apneic spells, attributed possibly to compression of vasomotor centers of the medulla or pons or possibly disturbances of autonomic hypothalamic centers, have also been reported. A large number of asymptomatic cases, presenting at autopsy were previously reported by Ambler in a comprehensive histological review of the subject.

The duration of symptoms ranges from a few months to 30 years prior to severe clinical deterioration. Cases mimicking subarachnoid hemorrhage have been reported. One such case presented with CT evidence of subarachnoid blood outlining the basal cisterns 20 years after a posterior fossa craniectomy and excisional biopsy for LDD. Angiography proved to be negative, with LDD as the only pathological finding. Age distribution at initial presentation of symptoms varies from 3 days to 63 years. Most often patients present in the age range of 30-41 years. Three cases of congenital LDD have been described. The male to female sex ratio in contrast to the previously reported 1:1 was noted to be 2:3 (20:31) not including autopsy cases. This is in agreement with Roski et al.'s recognition of a higher female preponderance.
Pathology

The underlying pathogenesis of this disorder remains uncertain, as indicated by the various descriptive names in the literature. The essence of the disorder is a thickened zone of abnormal cells generally thought to be dysplastic granule cells, which largely replace the normal granule cell layer in the affected areas of cerebellar cortex. This leads to abnormal folial architecture which accounts for the enlarged folia seen on MRI. Within this zone the normal small granule cell neurons are replaced by variably enlarged “dysplastic” cells of a neuronal phenotype. At the margins of the lesion an area of transition from normal granule cells to these dysplastic cells can be seen, indicating that the lesion is not as sharply demarcated as is suggested by MRI. The neuronal phenotype of these cells is confirmed by immunohistochemistry, showing positivity for neuron specific enolase and synaptophysin both in the cells, and in a perisomatic neuropil-like fashion. This is consistent with the known pattern of axosomatic synapses which develop around these cells and confirmed by previous immunohistochemical studies. It has been suggested that altered regulation of the cytoskeletal neurofilament protein plays a role in the abnormal morphology and increased size of the dysplastic cells. In our case neurofilament immunohistochemistry was largely negative in the dysplastic cells. Although the lesion has been referred to as dysplastic gangliocytoma, the number of larger ganglion type cells is often quite focal, as was seen in this case. Whether the ganglion cells represent Purkinje cells or an extreme form of the hypertrophic dysplastic granule cells is not resolved. Evidence in favour of a Purkinje cell origin is suggested by immunohistochemistry which shows positivity for the pan-T cell antibody anti-leu-4 which stains normal Purkinje cells but not granule cells. This is of importance because of the different embryologic origins of Purkinje and granule cell neurons. If both cells are involved and abnormal this would support a complex hamartomatous pathogenesis as opposed to a simple dysplasia of granule cells.

The molecular zone of the affected cerebellum is abnormal with loss of Purkinje cells and associated dendrites and synapses. This zone, which is normally axon and myelin free contains prominent projecting axons of the dysplastic granule cells which undergo aberrant myelination. The presence of abnormal myelination further support a hamartomatous process. Other pathologic changes described in LDD were not present in our case, including dysplastic calcification and, more rarely, a prominent plexus of angiomatous like vessels in the overlying meninges. The morphologic features of LDD have been well characterized, the fundamental pathogenesis remains undetermined. There is considerable controversy as to whether this represents a malformative lesion or a neoplastic process. Although this lesion tends to be very slowly progressive, clinically and radiologically it does slowly enlarge and can even recur following partial surgical resection. The basis of this enlargement and/or recurrence is not clear. It may represent a true increase in the number of cells (dysplastic granule cells) or an increase in size, in conjunction with increasing myelination of the molecular zone axonal plexus. However, the relative contribution of these processes has not been defined. There is no evidence in support of a neoplastic process and much evidence against one. The process does not invade surrounding brain tissue nor does it metastasize. Progression is extremely slow, and there is an absence of significant mitotic activity and cellular proliferation is minimal at most. Kinetic studies have shown little or no proliferative activity as shown by bromo-deoxyuridine labelling, and PCNA (proliferating cell nuclear antigen) or MIB-1 immunostaining. This is consistent with our immunostudies showing no MIB-1 immunolabeling of dysplastic cellular elements (MIB-1 is a paraffin analogue of Ki67 generally felt to be a better proliferation marker than PCNA).

The recent association of LDD with Cowden disease is consistent with a malformative process. Cowden disease consists of multiple hamartomas involving ectoderm, mesoderm and endodermal layers. Numerous cases of LDD have now been reported in Cowden syndrome families (up to as many as 30%) and indeed, from a clinical point of view, all patients with one disease should be screened for the presence of the other. Although a number of cancers occur with increased incidence in Cowden syndrome, chiefly breast, carcinoma, there is no association with, or increased incidence of CNS neoplasia. The presence of LDD arising congenitally in three cases, also supports and is consistent with a malformative nature of the process.

Treatment

Currently, a marked paucity of information regarding treatment strategies exists. Only sporadic case reports with extensive information regarding histology and pathology exist on the subject, yet little recourse for management prevails. Reasoning for resection, degree of resection, or outcomes of therapy, are lacking in the majority of the reports. Current modes of treatment described in the literature have consisted of medical (medications or radiotherapy) or surgical, consisting of VP shunt or biopsy and lesion debulking. No reports of long term medical treatment are described. Only one case of resolution of symptoms within 48 hours with mannitol and acetazolamide is described in the literature. The efficacy of radiotherapy in LDD with essentially no proliferative activity, is unknown. Five cases of such a treatment have been described. One of five actually worsened over a 7 year duration subsequent to the initial intervention, requiring a subtotal resection.

The first case report of surgical intervention was in 1930 by Bielschowsky and Simons and described a 20-year-old who presented with unsteadiness of gait, headaches and paresthesias in all limbs, vertigo, bilateral papilledema and coarse nystagmus. Of the first 12 reported cases of LDD from 1920 to 1955, 10 died. 5/10 cases were deaths on the operating table or immediately post-operatively, and the rest died prior to any surgical resection. In reviewing the English literature on LDD (diagnosed prior to autopsy), reports of surgical intervention have been described in 46/51 cases. Of the remaining 5 cases, two were reviews of prior reports. No information regarding degree of decompression was provided in one case acknowledged in Ambler’s composite review on the subject. Lastly, two cases were treated without resective therapy. Two cases received mannitol and acetazolamide as the primary mode of therapy, while the other was treated with a ventriculocervical shunt and radiation alone on the incorrect presumptive diagnosis of an astrocytoma. CSF drainage (temporary or permanent) has been reported in 20 of the 46 patients who received a biopsy, subtotal resection or a total resection. With
of subtotal resection, and 8/46 of complete resection have been described. Inadequate information, as to the degree of resection, is present for the remaining cases (i.e., 8/46). Little information as to the reasoning for the different options is provided. With evidence of a malformative lesion, LDD requires less invasive treatment. Resection of malformed tissue with minimal or no proliferative activity may not be the most optimal decision. The majority of symptoms, due to intracranial hypertension (i.e., headaches, nausea, vomiting, papilledema and diplopia) and local compression (cranial nerve and long tract findings) require treatment. Resection of abnormal cerebellar tissue does not relieve cerebellar symptoms but does decompress the posterior fossa and reduces intracranial hypertension. Ventriculoperitoneal shunting relieves the obstructive hydrocephalus, but allows for the local compressive effects of the lesion to persist. To incorporate treatment of the latter (space occupying lesion) in addition to the former (increased ICP) a method needs to be devised such that decompression of the malformed lesion is carried out. Subtotal or complete resection, well defined in the literature certainly would resolve both issues. The major flaw that exists is the poorly defined boundaries for resection, leading to resection of normal tissue.

Our case demonstrated a unique method in treatment of the malformation, allowing for resolution of both issues. The method involved decompression of the posterior fossa allowing for resolution of compressive symptoms upon the brain stem, adjacent cerebellum and fourth ventricle (thus also obstructive hydrocephalus). A craniectomy, in addition to a Cl laminectomy followed by a decompressive duroplasty (using autologous fascia lata graft) was performed. Decompression of the bony component in addition to the dura was achieved. The only other report of such a procedure was performed by Oppenheimer upon a 52-year-old presenting with a multitude of symptoms over an 18 month period. A posterior fossa decompression via a craniectomy and non-closure of the dura along with placement of a temporary ventriculostomy, was initially performed. A subsequent operative procedure was carried out a few weeks later due to frequent headaches despite improvement in the rest of the symptoms. Amputation of the obstructive tonsil and dural coverage with a fascial patch was carried out with success. Resolution of all other symptoms long after removal of the ventriculostomy, and prior to the second operation, proves the usefulness of dural decompressive surgery in Oppenheimer’s case. The above is certainly substantiated by our case as in many other malformative lesions of the posterior fossa (e.g., Chiari malformation).

REFERENCES