Cerebellar Dysfunction in Multiple Sclerosis

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Multiple sclerosis (MS) commonly affects the cerebellum causing acute and chronic symptoms. Cerebellar signs contribute significantly to clinical disability, and symptoms such as tremor, ataxia, and dysarthria are particularly difficult to treat. Increasing knowledge concerning the pathophysiology of cerebellar disease in MS from human postmortem studies, experimental models, and clinical trials has raised the hope that cerebellar symptoms will be better treated in the future.

Keywords: multiple sclerosis, demyelinating diseases, ataxia, cerebellar diseases, purkinje cells

INTRODUCTION TO MULTIPLE SCLEROSIS (MS)

Multiple sclerosis is an inflammatory disease of the central nervous system of unknown etiology. Typically patients have an initial relapsing and remitting course [relapsing remitting MS (RRMS)] followed by, in the majority of case, secondary progressive MS (SPMS) during which patients develop slow, insidious accumulation of disability (1). A small percentage of patients with MS have progressive disability from onset [primary progressive MS (PPMS)]. Despite extensive research and increasing knowledge of pathophysiological mechanisms, there is still no cure for the disease. Significant advances in therapeutics have occurred, and novel therapies, including natalizumab, alemtuzumab, fingolimod, and dimethyl fumarate have important disease-modifying effects (2).

The cerebellum and its efferent and afferent pathways are commonly affected in MS; and cerebellar ataxia is a common symptom of the disease, particularly in progressive disease (3, 4). Despite affecting the entire central nervous system, there are aspects of cerebellar involvement in MS that warrant specific attention and may give important insights into potential mechanisms and treatments for progressive disease. Within this review clinical aspects, pathological changes, monitoring of cerebellar changes in MS, and treatments for cerebellar disease will be discussed. In addition, various experimental models of cerebellar inflammation will be reviewed and how they may inform on future potential therapies.

CLINICAL FEATURES OF CEREBELLAR DYSFUNCTION IN MS

Coordination problems are common in MS and occur predominantly due to pathology within the cerebellum itself or impairment in cerebellar connections, including proprioceptive afferent inputs. Patients with MS may present with either acute cerebellar dysfunction relating to acute relapse or chronic cerebellar problems in progressive disease. Cerebellar pathology may lead to limb, gait, and truncal ataxia, dependent on precise lesional site, as well as other cerebellar features such as gaze-evoked nystagmus, dysarthria, and tremor.

Involvement of the cerebellum and brainstem connections occurs during MS relapse not infrequently. Cerebellar relapse at disease onset also seems to be associated with increased risk of cerebellar involvement during subsequent relapse (5). Multivariate analyses have suggested that involvement of the cerebellum at onset of the disease is associated with worse prognosis [shorter...
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In summary, cerebellar dysfunction is a feature of both RRMS and progressive MS leading to a range of neurological manifestations. In general, involvement of the cerebellum is linked to increased disability and worse prognosis, which, given the important role of cerebellar connections in motor control, is perhaps not surprising.

CEREBELLAR PATHOLOGY IN MS

Cerebellar white matter lesions are well described in the literature and often apparent on MRI scans of patients with MS. Cerebellar peduncles are common lesional sites. Demyelination commonly occurs in cerebellar white matter (Figure 1). In addition, observations concerning gray matter demyelination in cerebral cortex has led to studies evaluating gray matter disease in the cerebellum (18, 19). Indeed, the cerebellar cortex appears a major site for demyelination with one study reporting 38.7% of the cerebellar cortex being affected in a cohort of PPMS and SPMS patients (20). Interestingly, in this study, the majority of gray matter lesions appeared independent of white matter lesions, with some tissue blocks showing very extensive gray matter demyelination in the near absence of underlying white matter disease. Cerebellar cortical lesions are typically classified as leukocortical (extending from white matter into adjacent gray matter); intracortical (purely gray matter) arising around inflamed veins and venules; and subpial lesions that are “band-like” and parallel to the meningeal surface (20). The latter are abundant in patients with progressive MS. Observations concerning the relationship of subpial demyelination and meningeal inflammation have been suggested a possible driver for neurodegenerative processes in progressive MS, although causative proof is needed (21).

Neuronal pathology in the MS cerebellum is less well defined. The study by Kutzelnigg et al. (mentioned above) showed neuronal pathology with some reductions in Purkinje cell density in lesions (compared to control) (20). No significant reductions in Purkinje cell densities were seen in non-lesional cerebellar gray matter. Our own study confirmed these observations and also showed changes in neurofilament phosphorylation states in Purkinje cells (22). We noted abnormal neurofilament phosphorylation with loss of dephosphorylated neurofilaments and increased expression of hyperphosphorylated neurofilaments. In addition, axonal spheroids (representing transection of the Purkinje cell axon) were not within leukocortical lesions, indicating significant Purkinje cell pathology within the MS cerebellum. Alterations in neurofilament phosphorylation states and

FIGURE 1 | Proteolipid protein staining of human cerebellum of a patient with multiple sclerosis showing extensive white matter demyelination (red), which extends into the gray matter.

time to expanded disability status scale (EDSS) score 6] (6). In a recent database study of approximately 15,000 patients who experienced a total of nearly 50,000 relapses, cerebellar relapses accounted for approximately 10% of all relapses; being more frequent in men and in those with longer duration of disease (7). Cerebellar/brainstem relapses are also associated with poor relapse recovery which, in itself, is associated with earlier onset of progressive disease (8).

In patients with established MS, ataxia is thought to occur in about 80%, with symptoms particularly prevalent in those with progressive disease (3, 4). MS tremor is thought to arise predominantly as a consequence of cerebellar and/or thalamic disease (9). Tremor may affect limbs, trunk, vocal cords, and head (titubation). Intention and postural tremors are the commonest types, although rest and rubral tremors occur rarely. Ataxia has been reported in MS and is thought to occur due to midbrain pathology (13, 14).

The intriguing role of the cerebellum in cognitive processing is a subject in intense research, and there is some suggestion that involvement of the cerebellum may be linked to deficits in cognitive processes in MS (15). Injury to the cerebellum (from whatever cause) has been linked to deficits in verbal fluency, working memory, and attention, as well as executive dysfunction (16). Interestingly, the cognitive profile of MS patients with cerebellar lesions differs to those without cerebellar involvement (17). Reduced total cerebellar volume scores on magnetic resonance imaging (MRI) are associated with worse performance on cognitive tests (15). Specifically, volume loss of the posterior–inferior cerebellum is associated with poor cognitive function, whereas volume loss to anterior cerebellum is associated with motor dysfunction (12).

In summary, cerebellar dysfunction is a feature of both RRMS and progressive MS leading to a range of neurological manifestations. In general, involvement of the cerebellum is linked to increased disability and worse prognosis, which, given the important role of cerebellar connections in motor control, is perhaps not surprising.
axonal transection are reported within cerebral white matter
(23). In a further study of MS cerebellum, we showed reductions
in neuronal and myelin markers with evidence for increased
lipid peroxidation end products (end products of oxidative
injury to polyunsaturated fatty acids found in cell membranes)
(24). We also found elevation in superoxide dismutase enzyme
expression in MS cerebellum (compared to control) but a rela-
tive lack of other antioxidant enzyme expression, suggesting a
possible mechanism for the extensive oxidative stress-related
injury seen.

Other changes in Purkinje cell phenotype have been docu-
mented in MS, notably changes in ion channel expression and
receptor profiles. The Na\textsubscript{1.8} sensory neuron-specific sodium
channel is normally expressed at very low levels in Purkinje
cells, but its expression is markedly upregulated in MS (25). In
addition, annexin light chain (p11) that facilitates the functional
expression of this ion channel is also upregulated in Purkinje cells
(26). In experimental models, aberrant expression of Na\textsubscript{1.8} in
Purkinje cells causes significant abnormalities of firing patterns
of these cells (27, 28).

The interesting phenomenon of Purkinje cell heterokaryon
formation has been noted to occur in MS cerebellum (29). This
process appears to occur almost exclusively in Purkinje cells
within the central nervous system and denotes the appear-
ance of binucleated cells. Binucleation is thought to occur by
cell fusion, a process by which the nucleus of a non-Purkinje
cell is donated and integrates with the Purkinje cell (30). The
commonest type of “donor” cell in experimental paradigms of
cell fusion are bone marrow-derived cells, and researchers have
postulated the donation of a foreign nucleus to the Purkinje
cell may be a mechanism of protecting the cell from injury,
since expression of donor cell genes occurs. In MS, binucleated
Purkinje cells are significantly increased compared to control,
suggesting a possible adaptive process occurring in response to
the disease.

There still remains much to be understood about the pathol-
ogy of the cerebellum in MS; for instance, there is a paucity
of information on processes occurring in the molecular or
granular layers; and histopathological correlates of cerebellar
symptoms are lacking. Many of the pathological findings in
MS cerebellum are shared with other neurological conditions
affecting the cerebellum (e.g., neurofilament changes; Purkinje
cell injury). Understanding which pathological features are
specific to MS may help design future therapies. Furthermore,
understanding why cerebellar pathology is linked to a worse
prognosis in the disease may help design therapies for cerebel-
lar dysfunction in MS.

**MONITORING CEREBELLAR DISEASE IN MS**

The major disability rating scales used in MS incorporate esti-
mations of ataxia to a variable degree. These scales are used to
provide a semiquantitative description of disability in the disease
and remain prominent outcome measures of new therapies in
clinical trials. The EDSS is the most commonly used disability
scale in MS trials, and its quantification of disability relies heavily
on motor dysfunction (31). As part of the grading, disturbances
of functional systems (FS), such as visual function or bowel and
bladder function, are assessed and can contribute to the overall
score attributed. The cerebellar system is one of the FS tested,
and increasing levels of ataxia may contribute to the overall
EDSS level. Other scales also incorporate cerebellar function. For
instance, the multiple sclerosis functional composite has a 9-hole
peg test as one of the three tests involved, which relies heavily
on the integrity of cerebellar functioning (32). The relevance
of specific ataxia ratings scales has not been studied in MS in
detail. Primarily these scales have been developed for assessment
of “pure” cerebellar or spinocerebellar syndromes, typically the
inherited ataxias.

Magnetic resonance imaging studies have confirmed exten-
sive cerebellar involvement in both RRMS and progressive MS
(33, 34). The revised McDonald criteria for the diagnosis of MS
recognize infratentorial regions of the central nervous system to
be a typical lesional site in MS (35). Recent diffusion-weighted
MRI (DWI) techniques using a diffusion tensor model have
changes in white matter tracts in MS, notably cerebellar pedun-
cles. One study revealed abnormalities in tractography signals
within the superior cerebellar peduncle, which correlated with
upper limb dysfunction in patients with PPMS (36). Indeed,
the presence of T2 lesions within cerebellar peduncles on MRI
was associated with cerebellar and ambulatory symptoms in a
large imaging study (37). As stated above, cerebellar gray matter
disease may be extensive in pathological samples. A combined
MRI and posturography study showed that gray matter atrophy
of the superior lobules of the cerebellum (IV, V, VI), and lobules
VIII correlated with worse posturometric values (38). Magnetic
resonance spectroscopy (MRS) has been used to study neuronal
markers, typically N-acetyl aspartate, in patients with MS look-
ing for evidence of neuronal/axonal loss in progressive disease.
 Persistent cerebellar dysfunction has been linked with MRS
markers of axonal loss in the cerebellum of patients with progres-
sive MS (39).

Other cerebellar monitoring techniques, such as saccadic
movements using eye-tracking technology, may become useful
methods of monitoring cerebellar disease in MS (40). Defining
suitable measures of cerebellar disease burden may be of impor-
tance in the future should disease modifying or symptomatic
therapies be developed for specific cerebellar issues related to MS.

**EXPERIMENTAL MODELS OF CEREBELLAR INFLAMMATION
AND NEURODEGENERATION**

The most well-established animal model for MS is experimental
autoimmune encephalomyelitis (EAE). It is, however, not without
its limitations, since many of the features of MS fail to be repli-
cated in the model. Cerebellar inflammation occurs commonly
during the course of EAE. The specific mode of inducing EAE
may influence the predilection for cerebellar involvement (41).
Purkinje cell dysfunction has been show to occur in EAE with
reduced synaptic function (42). Loss of Purkinje cell and gray
matter pathology have also been documented in EAE (43).
Several models of dysmyelination and non-immune myelin loss have been studied. A model of slow, progressive myelin loss in a naturally occurring rat mutant is associated with extensive axonal changes (spheroid formation and neurofilament dephosphorylation) within the cerebellum (44).

Ion channel abnormalities in EAE have been noted for some time and form the basis for ongoing trials of sodium channel blockers in MS. Persistent sodium ion influx into the axon (associated with failure of ATP-dependent sodium/potassium exchange mechanisms in the context of reduced energy availability) leads to reversal of the sodium/calcium exchanger and thus excess intra-axonal calcium ions accumulate. Sodium channel blockade certainly appears to ameliorate axonal injury in EAE, but so far human trials have been less impressive (45). Indeed, ion channel abnormalities in Purkinje cells of mice affected by EAE have been documented, with aberrant expression of Na.1.8 channels causing alteration of electrophysiological properties of Purkinje cells (28).

The finding of cell fusion in MS cerebellum noted above has been postulated as a potential mechanism for Purkinje cell rescue in various neurological diseases. Transplantation of human menenchymal stem cells into mice which have subsequently been given EAE leads to expression of human markers within the cerebellum of the mice, suggesting that in EAE cell fusion occurs (46). Whether this leads to some form of neuroprotection in the cerebellum is unclear.

TREATMENTS FOR CEREBELLAR DISEASE IN MS

Treatments for MS fall into the categories of symptomatic treatments for established symptoms and disease-modifying therapies (DMTs), which aim to reduce the burden of disease. The design of DMT trials has not sought to determine whether these drugs improve cerebellar function specifically. The majority of trials have used reduction in annualized relapse rates and EDSS scores (as a marker of disability progression). As mentioned earlier, the EDSS does have cerebellar function as part of the assessment, but its utility in picking out specific effects of drugs on cerebellar function is limited. However, cerebellar relapses are associated with an increased risk of disability accrual (relative to some other relapse types, such as sensory) (47). Thus, reducing relapses by DMTs is likely to reduce the burden of cerebellar disease in the long term. Interestingly, analysis of an alemtuzumab trial (CAMSS223 versus beta-interferon) demonstrated improvements in the cerebellar FS score of the EDSS (48).

Non-pharmacological approaches to MS ataxia are commonly employed, of which physiotherapy regimens are the most widely accepted. Balance-specific exercises involving somatosensory and motor strategy facilitation are generally employed to varying degrees (49). Improving core stability in patients with balance problems may be effective, and lumbar stabilization exercises (which improve core trunk muscles, leading to effects on postural control, ambulation, and skilled motor function) are often incorporated into MS rehabilitation programs (50).

In addition, task-oriented training enhances ambulation and postural control in MS patients due to the promotion of motor learning (51). In general, combinations of these physiotherapy approaches are thought to be most beneficial (52).

Pharmacological approaches to improving ataxic symptoms are generally disappointing, and newer therapies are needed. The most recent Cochrane review of treatments for ataxia in MS (which reviewed six randomized placebo-controlled trials) concluded that absolute and comparative efficacy and tolerability of pharmacotherapies are poorly documented and no recommendations could be made (53). Small open label studies or case reports have suggested benefits for a range of drugs for the treatment of tremor. Isoniazid, propranolol, and levetiracetam have been studied, although the data on their use are not convincing (the number of patients involved in these trials was generally very low, and limited conclusions could be drawn) and they are not widely used (54–59). Several randomized controlled trials of cannabis extracts have concluded that cannabinoids appear to have no beneficial effect on MS tremor (60–62). Paroxysmal ataxia and dysarthria have been reported in MS, albeit rarely, and there is some suggestion that they may respond to carbamazepine, in a similar way to other paroxysmal symptoms of MS, such as tonic spasm (63).

The majority of reports of stereotactic surgery in MS tremor have targeted thalamic structures with variable results (64, 65). In the Cochrane review, one neurosurgical study of thalamotomy versus thalamic stimulation was included (53). Tremor was abolished by both thalamotomy and thalamic stimulation in all patients immediately postsurgery (66). However, tremor returned in almost all MS patients after 6 months (albeit of less severity than preoperative levels) and general disability scores were unchanged. The short-lived nature of the response is seen in other studies of surgical treatments of MS tremor (67). Improvement in quality of life measures following thalamic stimulation, including improvement in ability to feed oneself, has been demonstrated in selected cases (68).

There is a great need to improve therapeutic options for symptomatic treatments of cerebellar symptoms and to provide neuroprotection within the cerebellum. The design of trials aimed specifically at cerebellar protection in MS will be challenging due to the paucity of good outcome measures, although improvements in imaging techniques will help. Refinements in neurosurgical techniques may help patients with severe ataxic tremors. Increased understanding of the pathophysiology of cerebellar disease in MS will aid the search for new drug therapies. A number of trials are now being pioneered specifically aimed at slowing disease progression. Newer therapies, such as stem cell therapies, are being developed. The observation of Purkinje cell fusion as a potential neurorestorative mechanism makes the prospect of stem cell treatments for MS cerebellar disease particularly attractive.

AUTHOR CONTRIBUTIONS

The author confirms being the sole contributor of this work and approved it for publication.
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