A Review of Multiple Sclerosis as an Infectious Syndrome

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Abstract

Multiple sclerosis (MS) is a chronic neuroinflammatory disease of the central nervous system (CNS) that is generally considered to be an autoimmune myelitis of unknown etiology. Epidemiological studies suggest that infection may act as a trigger on a predisposing genetic background. A number of causative agents have been considered. This review will focus on the pathogenesis of MS and link this pathophysiology to infectious agents that have been implicated as possible co-factors. By doing so, MS will be viewed as an infectious syndrome that involves a CNS infection that results in a neurodegenerative process as well as an autoimmune disease. Early detection of infectious triggers could allow appropriate intervention and thus improved outcomes.

Keywords: Multiple sclerosis; Pathogenesis; Pathophysiology; Demyelinating disease; Infectious syndrome; Central nervous system toxin; Clostridium perfringens type B; Epstein-Barr virus; Human herpes virus 6; Human endogenous retroviruses; Lyme Borrelia complex; Chlamyphila pneumoniae; Heat shock protein 60

Introduction

Multiple sclerosis (MS) is a chronic neuroinflammatory disease that is characterized by progressive, inflammatory, and multifocal demyelination of the brain and spinal cord [1-3]. MS affects approximately 2.5 million people worldwide, with women being afflicted twice as frequently as men [4-6]. Importantly, young adults are the primary groups afflicted with MS: the average age of diseases onset is 30 years, with half of these patients requiring a wheelchair within 25 years of their diagnosis [7-9]. The onset and progression of MS in young adults places a considerable economic burden on these patients, their families, as well as on health and social resources [10].

Currently, there are three prevalent theories on the pathogenic mechanisms for MS: autoimmune, degenerative, and infectious [1-3,8,11-13]. This review briefly describes the evidence for each of these three proposed pathogenic mechanisms and will point out that these pathogenic mechanisms are not mutually exclusive. In fact, a cogent argument will be made that all three proposed mechanisms are involved in the pathogenesis of MS. Moreover, this review will reinforce the evolving concept that brain infections may be triggers for the autoimmune encephalitis and neurodegenerative process that is known as MS.

It is useful to begin this review with a description of the development and progression of the demyelinating plaque, which is the pathological hallmark of MS. Demyelinating lesions in the white matter were first described by Charcot in 1848 and were based on the gross pathology findings of hardened translucent discoloured plaques of brain tissue distributed randomly throughout the white matter of the brain [14]. Charcot believed that these areas were caused by an unidentified toxin or pathogen or perhaps by a disturbance of the circulation. Dawson in 1916 described the histopathology of these plaques as consisting of demyelination and gliosis around blood vessels [15]. Both early [14,15] and contemporary [16] investigators have noted that these plaques often involve the optic nerve and tend to be located in the periventricular white matter as well as in the cerebellum, the brainstem, and the cervical spinal cord of MS patients. Acute active lesions in MS histologically reveal perivascular infiltration of lymphocytes, plasma cells and macrophages [17-20]. Active lesions are recognized over time as progressing [8,19], with continuous breakdown and regeneration of both myelin [21] and oligodendrocytes [22]. Macrophages are present in the active lesion [20] and are currently thought to be engaged in scavenger activity rather than participating in an autoimmune attack on constituents of the myelin sheath [22]. Heterogeneity of MS lesions has been noted with lesions being distinguished as early active, late active, smouldering, inactive and shadow plaques [23-27]. The histological features of plaques appear to vary over time and also display gender and age-specific differences. These findings suggest the possibility that the targets of injury and the mechanisms of demyelination in MS may be different in different disease subgroups, and that the immune response may shift from adaptive to innate [27]. Finally, the acute MS lesion is associated with increased permeability of the blood brain barrier [27]; such increased permeability of the blood brain barrier has been shown to predict the conversion of optic neuritis to multiple sclerosis [28].

Chronic plaques in MS are seen more frequently with progressive MS and have been classified as smouldering, inactive or shadow plaques [24-27]. Inflammation is present in almost every lesion type although the severity decreases with the age of the patient and the duration of disease. In contrast to acute plaques, the blood brain barrier does not display increased permeability with chronic plaques [29]. Lymphoid follicular structures are found in perivascular locations, and plasma cells increase and persist within the central nervous system. Remyelination can be seen in all stages of plaque and is characterized by the presence of newly formed myelin sheaths as well as by oligodendrocyte precursor cells [30]. The transition of plaque from acute active lesions to chronic inactive lesions is characterized by the decrease in inflammation and the gradual disappearance of macrophages and microglia. Astrocytes actively produce glial fibers, resulting in a glial scar. On gross examination of the MS brain, these glial scars are seen as distinctly demarcated, firm
brownish discolorations and represent the white matter abnormalities that Charcot first noted as "scleroses en plaque" [14].

Although MS typically has been considered a disease of white matter, it has become very clear that gray matter is also involved [31]. Of note is that Charcot in 1877 also described demyelinated lesions in gray matter [32], but these observations were mostly ignored. Histological, immunohistochemical studies, and DIR-based imaging studies clearly demonstrate that cortical gray matter is extensively demyelinated in MS, and that these cortical lesions represent a significant proportion of the pathology of MS [31]. Moreover, the cellular composition of the cortical gray matter lesions is considerably different from that of white matter lesions, suggesting that there is a location-dependent expression of the immunopathological process in MS [31,33]. The most extensive cortical damage is located in the cingulate gyrus and the temporal and frontal cortices. In addition, extensive demyelination is also noted in the hippocampus of MS patients. These areas may be particularly susceptible to inflammation and/or neurodegenerative processes [31]. A currently accepted classification of cortical lesions describes Type I lesions as those that extend through both the white matter and the gray matter. Type II lesions are entirely located with the cerebral cortex and do not interact with the subcortical white matter or the pia mater; these lesions, in general, are perivascular small lesions. Type III lesions are subpial areas of demyelination that do not extend beyond layers 3 and 4 of the cortex. Finally, Type IV lesions typically extend over the entire width of the cortex, but do not extend into the subcortical white matter; these Type IV lesions may spread over several gyri or entire lobes [31]. Of note is the difference in the degree and type of inflammatory response between demyelination in white matter versus that in gray matter, with gray matter demyelination having significantly less lymphocytic infiltration than that seen with white matter demyelination. Another difference in the cellular inflammatory response seen in gray matter demyelination is the presence of activated microglial cells in the demyelinated tissue rather than phagocytic macrophages that are typically seen in white matter demyelination [32,34]. Finally, in those cortical lesions that do not extend into white matter, there appears to be normal blood-brain barrier function [32,35].

In addition to demyelination of both white and gray matter, it has become clear that the pathogenesis of MS also involves widespread synaptic loss that is independent of demyelination and axonal degeneration in the gray matter of MS brains [36]. Although the exact pathophysiology for this widespread synaptic loss is unclear, one possible explanation is that synapses could be a susceptible target of widespread inflammation [36]. For example, secreted inflammatory mediators such as interferon-gamma [37] and TNF-alpha [38] and/or diffusely activated microglial cells induced by focal activation of the complement cascade [39] may contribute to lower-grade, but widespread inflammation that is independent from white and gray matter plaques. This process of low-grade inflammation may be assisted in some MS patients by the presence of B-cell follicle-like structures in the meninges [40].

This brief overview of the pathologic lesions involved in the MS brain strongly suggests that there exists a widespread progressive neurodegenerative disease in which persistent inflammation plays a prominent role. It is very likely that there is an autoimmune component to this persistent inflammation, although specific autoreactive targets have not been well delineated [41] and immunotherapy of MS does not prevent disease progression in a substantial number of patients [42].

Given that both neurodegeneration and autoimmunity appear to be involved in the pathogenesis of MS, what role might infection(s) play? Infections have long been thought to play a role in MS, but that role remains to be clearly defined. The possible roles of infections as triggers for a neurodegenerative process with an autoimmune response in the pathogenesis of MS will be reviewed using examples from the medical literature.

Charcot believed that demyelinated plaques observed in MS brains might be caused by an unidentified toxin. Such a toxin could originate from the GI tract via a colonizing or an infecting microbe. This possibility is nicely illustrated by work done by Rumah et al [42–44]. These investigators initially described the isolation of Clostridium perfringens type B from the stool of a young woman, who presented with actively enhancing lesions on her brain MRI and was diagnosed with MS [43]. C. perfringens type B is an epsilon toxin-secreting bacillus that can be found in the stool of ruminant animals; this 29 kDa epsilon toxin in these ruminant animals can cause a devastating enterotoxemia disease in which there is blood brain barrier dysfunction and white matter injury [44,45]. These investigators also demonstrated that immunoreactivity to this epsilon toxin is 10 times more prevalent in people with MS than in healthy controls [43], and that the epsilon toxin can bind to mammalian cells via the myelin and lymphocyte protein MAL [44]. Moreover, these investigators have shown that in primary murine CNS cultures, the epsilon toxin binds to and kills mature oligodendrocytes, but not astrocytes, microglia, or neurons [45]. For this epsilon toxin to function as a trigger for MS in humans, colonization of a human host by C. perfringens type B would be needed followed by dissemination of this toxin to the bloodstream. Once in the blood stream, this 29 kDa epsilon toxin could reach brain tissue through the third and fourth ventricles via the fenestrated capillaries of the circumventricular organs, which are known to allow direct exchange between the blood and the central nervous system [46]. Conversely, the epsilon toxin could disrupt the blood brain barrier and thus reach brain tissue in a more direct manner. The first of the two possibilities would be consistent with the location of early MS lesions and the progression of these lesions [17,19,20,22].

Another way in which infection could play a role in MS would be a viral infection acting as a trigger for neurodegeneration. This association of Epstein-Barr virus (EBV) with MS was first made when serum antibody titers to EBV were noted in MS patients; the geometric mean titer of antibodies to EBV was significantly higher in MS patients that in controls (107.0 versus 77.1) [47]. This association was further strengthened when Epstein Barr nuclear antigen-1 (EBNA-1) was the target of the oligoclonal bands in the CSF of some MS patients [48]. In 5/15 MS patients, a distinctive oligoclonal band antigen specific banding pattern for EBNA-1 was observed in comparison to 0/12 controls (P=0.037, Fisher’s Exact Probability). The presence of intrathecal IgG in the form of oligoclonal bands is seen in the CSF in greater than 90% of MS patients [49]. Oligoclonal bands are not specific for MS because such bands are also seen in 10% of patients with other inflammatory diseases of the CNS. In most other diseases in which oligoclonal bands are present in the CSF, the bands represent an antigen-driven immune response to infectious agents [50]. There are a number of ways the EBV might be involved in MS [51–53]. For example, EBV may play a role in MS by initially infecting naive B cells in the tonsils, these latently EBV-infected B cells after replicating in germinal centers may enter the circulation as well as the CNS where they could produce oligoclonal IgG. At the same time, T cells would be targeting EBV-infected B cells in the body and also might target them in the CNS; when present in the CNS, these T cells would produce...
cytokines such as interleukin 2, interferon gamma, and tumor necrosis factor beta. These cytokines in turn would result in demyelination as well as axonal injury [54-57].

Another virus that has been associated with MS is human herpes virus 6 (HHV-6) [58]. Challoner et al. detected by PCR a 341-nucleotide fragment that was 99.4% identical to the major DNA binding protein gene of HHV-6 in greater than 70% of MS cases and controls suggesting that HHV-6 is a commensal virus of the human brain [58]. HHV-6 expression was determined by immunocytochemistry with monoclonal antibodies; nuclear staining of oligodendrocytes was seen in MS cases, but not in controls [58]. Moreover, nuclear staining was seen around plaque more often than in uninvolved white matter; these observations suggest an association of HHV-6 with the etiology or pathogenesis of MS [58]. Other investigators confirmed this association of HHV-6 with MS [59-66]. Of interest is one report that demonstrated oligoclonal bands in 38% of MS patients that were reactive to either EBV or HHV-6, suggesting that both viruses may be involved in separate cohorts of MS patients [63].

The role of HHV-6 in the pathogenesis of MS might involve direct infection of oligodendrocytes that could then lead to neurodegeneration with autoimmunity and cytokine production [65,66].

A third group of viruses associated with MS is the human endogenous retroviruses (HERVs) [67,68]. This association began when Perron et al. reported the isolation of a leptomeningeal cell line (LM7) from the CSF of a patient with MS; this cell line was noted to have viral reverse transcriptase activity [69]. Moreover, electron microscopy revealed the presence of viral particles in this LM7 cell line [69]. These investigators then reported isolation of a retrovirus in human monocyes cultures from 18 patients with MS [70]. This group of investigators next used molecular methods to characterize this novel MS-associated retrovirus that had been repeatedly isolated from leptomeningeal cells, post-mortem choroid plexus cells, and EBV-immortalized B cells from MS patients [71]. Other investigators have confirmed the presence of endogenous human retroviruses from patients with MS [72-75], although the exact role of these retroviruses, if any, in the pathogenesis of MS remains unclear. Moreover, it should be noted that endogenous human retroviruses also have been implicated in the pathogenesis of amyotrophic lateral sclerosis (ALS) [76,77]. Clearly, the role of endogenous human retroviruses in neurological disorders such as MS and ALS should be further investigated.

Viruses are not the only microbes that have been associated with MS. One of several bacterial pathogens that have been suggested as possible causes of MS is the spirochete [78-80]. The association of spirochetes with MS began in 1925 when Adams et al. inoculated rhesus monkeys with material from MS cases and observed spirochetes in the ventricular fluid of these monkeys several months later [81]. Steiner in 1952 reported spirochetes in plaque from autopsied MS patients [82], while Ichelson in 1957 reported cultivation of spirochetes from spinal fluids from MS patients [83]. Newman et al. repeated the culture method used by Ichelson and found spirochetes in spinal fluid from MS patients, albeit in a lesser percentage (18.5% versus 78%) [84]. The recognition of Lyme arthritis in 1977 [85] and its spirochetal etiology in 1983 [86] was followed by the association of Lyme disease with chronic neurologic abnormalities [87,88] and the detection of Borrelia burgdorferi DNA by PCR in CSF from patients with Lyme neuroborreliosis [89]. Lyme neuroborreliosis has been well characterized in the medical literature [87-91]. Although antibiotic responsive demyelinating neuropathy related to Lyme disease has been described [92], this is the exception and not the rule [93]. Moreover, the finding of reactive Lyme serology in MS patients has not been found to indicate neurologic Lyme disease [94]. Nevertheless, it should be noted that there are different Borrelia species that cause Lyme disease [91]. In the United States, Borrelia burgdorferi and a newly described spirochete species, Borrelia miyamotoi, [92] are known to cause Lyme disease. In Europe, at least 5 Borrelia species are known to cause human infections [91] with at least 3 of these species (B. burgdorferi, B. afzelii and B. garinii) having been identified by PCR in CSF from patients with neuroborreliosis [95,96] and a fourth species, B. bavariensis [97] recently described as a cause of neuroborreliosis [91]. The Lyme Borrelia complex is expanding and the exact role of newer genomic species as human pathogens remains to be determined [98].
Peripheral blood mononuclear cells as has been demonstrated in an experimental transendothelial migration model [132]. Evidence that this can occur is seen in a report of chlamydial infection of ependymal cells in the ventricular tissues of some patients with multiple sclerosis [133]. These investigators suggest that the circumventricular organs may allow entry of C. pneumoniae-infected mononuclear cells into the CNS. Infection of ventricular ependymal cells by C. pneumoniae could then lead to cytokine-mediated damage to brain tissue surrounding these ventricles. Moreover, infection of ventricular ependymal cells by C. pneumoniae would explain the visualization by electron microscopy of chlamydial elementary bodies in the CSF of MS patients [133]. Such ventricular infection by Chlamydia would also explain the observations that affinity-driven immunoblot studies have demonstrated reactivity of oligoclonal bands in CSF from MS patients against elementary body antigens of C. pneumoniae [134,135]. Finally, ventricular infection by C. pneumoniae would explain the presence of chlamydial DNA and mRNA transcripts in CSF from some patients with MS [136,137].

Chlamydiae are known to produce large amounts of heat shock protein 60 (HSP 60) during chronic, persistent infections such as seen in human atheroma [138]; this HSP 60 is thought to play an important role in the pathogenesis of chronic chlamydial infections [139]. Heat shock protein 60 is an evolutionarily conserved protein in both eukaryotes and prokaryotes that is recognized as potent activator of the innate immune system and is capable of inducing the production of proinflammatory cytokines by the monocyte-macrophage system [140-143]. Importantly, human mitochondrial HSP 60 has been proposed as a danger signal of stressed or damaged cells [144]. Both chlamydial HSP 60 and human HSP 60 contribute to inflammation in human atheroma and are able induce inflammation to levels induced by Escherichia coli lipopolysaccharide [144]. Infection of ventricular ependymal cells by C. pneumoniae or other pathogens could serve as a focus of HSP 60, which in turn would stress and/or damage neurons and oligodendrocytes resulting in neuronal cell death, axonal injury, loss of oligodendrocytes, and demyelination [145]. Such HSP 60 mediated neurodegeneration and demyelination in the CNS has been described in a murine model in which intrathecal HSP 60 is injected [146]. The fact that neurodegeneration and demyelination in the CNS involves TLR4 and MyD88 confirms that this molecular pathway is able to the release of endogenous TLR ligands from injured CNS cells and is a common process in many different types of brain injury. Indeed, once there a brain injury involving release of HSP 60 has occurred, a vicious cycle can occur due to the activation of microglia, the resident CNS monocyte. Such a cycle involving release of HSP 60 from injured CNS tissue with activation of toll-like receptor 4 has been shown to medicate neurodegeneration in the CNS of mice [147] and has been implicated in multiple sclerosis [148] and myalgic encephalomyelitis [149].

This review of the pathophysiology of multiple sclerosis clearly demonstrates that MS is a widespread CNS neurodegenerative process that involves white matter, gray matter, and the spinal cord. This neurodegenerative process seen in MS is similar to many other neurologic diseases such as Alzheimer’s and Parkinson’s diseases, amyotrophic lateral sclerosis, and prion diseases in that it is an inflammatory process [150]. The inflammatory process seen with MS and other neurologic diseases is mediated by HSP 60 from stressed and damaged cells as well as by multiple inflammatory cytokines produced by the immune cells, both local resident cells like activated microglia and by infiltrating immune cells [54]. Hence, this process can be correctly termed an “autoimmune” process. The important role that chlamydial HSP 60 may play as a trigger and human HSP 60 may play as a propagator in this CNS autoimmune process are only now becoming apparent. Because of the numerous ways in which stressed or injured cells in the CNS might contribute to this autoimmune process, it has been difficult to identify a specific “autoreactive target” although HSP 60 might well deserve this designation. Equally difficult is identifying a “trigger” event. A careful review of the medical literature suggests a number of infectious triggers. Of these, Chlamydia pneumoniae is a particularly compelling in that this pathogen could play an important role in several different ways. One way would be to support involvement in the development of chronic cerebrospinal venous insufficiency as recently described [151]. Another way would be to support infection of the ependymal cells of the ventricles where the production of HSP 60 could serve as an inflammatory triggering as described in this commentary. Clearly, the choroid plexus is now recognized as an important component of MS [152,153]. Other toxins or pathogens could enter the CNS via the choroid plexus/ circumventricular organs and/or infect these CNS tissues. Indeed, equally compelling reports in the medical literature strongly suggest that MS is an infectious syndrome with a number of different ways that infection may play a role [103].

Regardless of which infectious triggers may be involved, MS should be considered to be an infectious syndrome that involves a neurodegenerative process resulting in an autoimmune disease. Identification of each potential infectious trigger described in this review early in the course of MS might allow appropriate antimicrobial intervention as has been done with at least one of these triggers, C. pneumoniae [154]. Future studies in MS should identify each of these potential infectious triggers early in the course of MS. Such identification might then allow appropriate antimicrobial therapy. Moreover, stratification of MS based on specific infectious triggers would allow specific clinical trials that are focused on each pathogen.

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