Immune Responses in Neurodegenerative Diseases
Shrestha R,1 Shakya Shrestha S,1 Millington O,2 Brewer J,3 Bushell T2

1Department of Pharmacology
Dhulikhel Hospital - Kathmandu University Hospital
Kathmandu University School of Medical Sciences
Dhulikhel, Nepal

2Strathclyde Institute of Pharmacy and Biomedical Sciences, University of Strathclyde Glasgow, UK

3Institute of Infection, Immunity and Inflammation
College of Medical, Veterinary and Life Sciences
University of Glasgow, UK

Corresponding Author
Rajeev Shrestha
Department of Pharmacology
Dhulikhel Hospital - Kathmandu University Hospital
Kathmandu University School of Medical Sciences
Dhulikhel, Nepal
Email:rmaleku@hotmail.com

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ABSTRACT
Neurodegenerative disease is a progressive loss of neurons from central nervous system and has a huge impact on health care system. Various causes have been proposed of which inflammation has been suggested to be a probable key factor in the most of such conditions. The involvement of immune cells including lymphocytes in such diseased condition of the CNS supports this notion. The effective therapy for these diseases has been sought for more than a half century but still lacking such therapy. On such basis this review article has mainly focussed on evidence of the involvement of immune cells in various neurodegenerative diseases including Alzheimer’s disease, Parkinson’s diseases and Multiple sclerosis and suggests a possible therapy of such diseased conditions of the CNS by the modulation of immune system.

KEYWORDS
Alzheimer’s disease, lymphocytes, multiple sclerosis, neurodegenerative diseases, parkinson’s disease, stroke, t cells

INTRODUCTION
The global burden of neurodegenerative diseases is huge and accounts for at least 15% of the global burden of diseases. Although cases of neurodegenerative disease are increasing every year, reliable therapeutics are still being sought and a complete understanding of the underlying biology is lacking.

Neurodegenerative disease is a progressive loss of neurons from the CNS, associated with a deficit in the function of the affected region. Cell death during neurodegeneration can be either via apoptosis or necrosis or both. Various conditions have been suggested for the neurodegeneration of the CNS including ageing, inflammation, stress and trauma and genetic predisposition. Recent studies have shown a strong link between inflammation and neurodegeneration but the exact role for inflammation in neurodegeneration is still elusive. It is not clear whether inflammation causes neuronal death in neurodegenerative diseases or whether the inflammatory infiltrate is simply a manifestation of the disease process. However, several possibilities have been proposed for the link between inflammation and neurodegeneration: 1) inflammation induces neurodegeneration, 2) neurodegeneration causes inflammation, 3) other factors cause either inflammation or neurodegeneration or both, 4) inflammation and neurodegeneration occur as a cycle which amplifies each others response and 5) inflammation can be neuroprotective in neurodegeneration. The key features of CNS inflammation are glial cell activation, local production of inflammatory mediators, expression of MHC and adhesion molecules, release of free-radicals and...
recruitment of immune cells. During neurodegenerative diseases, either peripheral immune cells, such as T cells, initiate inflammation in the CNS or CNS resident immune competent cells such as microglia as well as neurons, astrocytes and oligodendrocytes, release inflammatory mediators to recruit more peripheral immune cells including lymphocytes leading to CNS inflammation. Most commonly, inflammation starts within subarachnoid space which disseminates to other regions of the brain. 

During inflammation of the CNS, endothelial cells of the BBB express various selectins and adhesion molecules that increase the migration of lymphocytes from the systemic circulation to the perivascular spaces of the brain. Further, activated lymphocytes also express various receptors including chemokines receptors, integrins and selectins that help to interact with their respective ligands expressed on the surface of endothelial cells during neuroinflammation. Activated lymphocytes and cells of the CNS including microglia, astrocytes, neurons and oligodendrocytes release various pro-inflammatory cytokines such as IL-1, TNF-α, IL-23, INF-γ and chemokines including various neurotrophic factors which can contribute in the outcome of the CNS inflammation. 

There are several neurodegenerative diseases including AD, MS, PD and stroke in which lymphocytes are actively involved and believed to be a key player in the initiation of CNS inflammation. Some examples of the most common neurodegenerative diseases are briefly explained in subsequent headings.

**Alzheimer’s disease**

Alzheimer’s disease (AD) is a progressive neurodegenerative disease that mostly affects patients in their later stage of life. Typical symptoms of AD are loss of cognitive functions including emotion, learning and memory processing skills leading to dementia. The pathological impression of AD can be characterised by the deposition of amyloid-beta (Aβ) protein plaques in the brain parenchyma and accumulation of tau proteins within neurons. These protein plaques are thought to interfere with synaptic transmission and neuron-neuron communication leading to neuronal death. Further, high levels of tau proteins within neurons form tangles and block transportation of nutrients or other vital cellular factors throughout the cell which has been suggested to be one of the reasons for cell death in AD.

In AD, amyloid-beta plaques and tau proteins are considered to be crucial in the pathology. Recently, inflammation has been proposed to be one of the key players in AD. The inflammatory responses in AD can be characterised by the up-regulation of cytokines and chemokines along with activation of microglia. The activated microglia clusters can be seen near amyloid-beta deposition site and these cells also express high levels of MHC-II, cytokines and chemokines contributing to disease progression. However, these microglial cells also involve in clearing of amyloid-beta and this function has been shown to be enhanced in the presence of TGF-β. In addition, reactive astrocytes are also clustered at the site of amyloid-beta deposition but high levels of these cells also accumulate at neuritic plaques. These reactive astrocytes are capable of expressing various cytokines, growth factors, adhesion molecules and prostaglandins. Furthermore, astrocytes are suggested to be involved in inhibition of microglial ability to clear amyloid beta. The analysis of brain autopsy has also revealed that there is a significant increase in inflammatory markers as well as increase in complement activation and lysis of neurites in AD subjects when compared to non-demented subjects, which strongly suggests that inflammation has a role in AD. With these findings, it can be suggested that this inflammatory responses might contribute in recruiting lymphocytes from the systemic circulation into the brain. Further, T cells have been detected in the brain of AD patients. Recent studies have also demonstrated the up-regulation of T cells in the CNS of AD when compared to healthy controls. These studies have revealed an increased activity of various subsets of T cells such as Th-17 and Th-9 in AD and cytokines including IL-9, IL-21 and IL-23 released from these T cells are also increased in AD which have been suggested to be one of the factors in AD-associated neuroinflammation. The beneficial effect of T cells has been demonstrated in AD however, these T cells lose their protective effect in severe condition of AD. With these findings, it is evident that lymphocytes are present in the brain of AD and may have an important role in AD. However, the role of lymphocytes in AD is still poorly understood and a better understanding of this phenomenon could help in search of novel drug targets.

**Multiple sclerosis**

Multiple sclerosis (MS) is a chronic neuroinflammatory disease of the CNS characterised by demyelination, axonal damage and autoimmunity affecting people mostly between 20 and 40. It affects both white and gray matter of the CNS and scattered focal demyelinated lesions can be seen throughout the white matter of the CNS. There is an episodic exacerbation followed by remission during the course of the disease and this relapsing-remitting course is suggested to be immune-mediated which is characterised by activation of microglia and infiltration of peripheral immune cells into the CNS. This disease process leads to secondary MS having marked degeneration of neurons and axons along with massive cortical demyelination. Genetic predisposition, environmental factors and viral/microbial infections have been proposed as risk factors in MS but how these factors contribute to the aetiology of the disease is still under investigation. However, it is now well accepted that MS is a T cell mediated autoimmune disease. The concept of MS as an autoimmune disease arises due to its similarities in clinical symptoms with EAE, an animal model for this disease.
induced in the animal by immunising with myelin-derived protein such as myelin-basic protein, proteolipid protein or myelin oligodendrocyte glycoprotein and the disease is mainly initiated by myelin-specific autoreactive T cells. Autoreactive T cells have been identified in MS patients, as well as being present in healthy people. However, these T cells are more activated and have a memory phenotype in MS when compared to healthy subjects. These activated T cells express various chemokines, cytokines and adhesion molecules which help them to interact with the BBB and migrate into the CNS to initiate immune response against myelin-derived proteins. Although, CD4+ T cells are thought to be initiators of EAE, analysis of MS lesioned brain tissues has shown predominance of CD8+ T cells. CD4+ T cells can be either neuroprotective or pathogenic depending on the types of cytokines or neurotrophic factors they release. As described above, different subsets of CD4+ T cells release their functional cytokines: Th1 cells release inflammatory mediators including IFN-γ and TNF-α whereas Th-2 cells release anti-inflammatory mediators like IL-4. IFN-γ and TNF-α have shown a contrast in response in EAE and MS since IFN-γ exhibits neuroprotection in rodent models of EAE whereas it exacerbates the disease in MS. Similarly, blocking TNF-α function in EAE is neuroprotective but not in MS. This finding therefore suggests that caution should be taken while comparing data from the EAE model with MS. The major concept of immunopathogenesis of MS has been connected to the balance between Th-1 and Th-2 functions. However, there are other subsets of CD4+ T cells such as Th-17 and Treg cells along with CD8+ T cells cannot be excluded. Th-17 cells have been found to express more activation markers, co-stimulatory and adhesion molecules than Th-1 cells suggesting they are more pathogenic. Further, the pathogenic T cells suppressor capacity of Treg cells is found to be attenuated in MS. Moreover, CD8+ T cells have been directly linked to the demyelination of axons in MS and are pathogenic in the immune-mediated demyelination of axons. Nevertheless, there is no exact mechanism how these cells are contributing to the disease process or neuroprotection in MS and further research is required to understand MS pathology.

**Parkinson’s disease**

Parkinson’s disease (PD) is an age-related chronic neurodegenerative disease clinically characterised by tremor, rigidity, bradykinesia, postural instability, dementia and autonomic dysfunction while pathologically by loss of dopaminergic neurons in the substantia niagra and the presence of Lewy’s bodies which are aggregated proteins such as α-synuclein. The accumulation and misfolding of α-synuclein induce toxicity leading to the loss of dopaminergic neurons. This results in a reduction of dopamine production causing gait and movement impairment because dopamine is required for a normal motor function of the brain. Further, proteosomal and lysosomal system dysfunction and reduction in mitochondrial activity due to genetic mutations are also proposed to be causative factors in neuronal death during PD. Various risk factors including environmental genetics and age have been related to pathogenesis of PD. Furthermore, there are several studies suggesting the relation between inflammation and pathology of PD. However, it is still not clear whether inflammation observed in PD can be considered as classical inflammation or not. Therefore, the term ‘neuroinflammation’ has been coined in the pathology of PD. The upregulation of MHC expression is one of the first signs of neuroinflammation in PD with an increase in MHC-II expressed microglia in the substantia niagra. Similar upregulation of MHC molecules has been reported in 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP)-intoxicated animals, an animal model for PD and is also associated with the infiltration of lymphocytes in substantia niagra. The other hallmarks for neuroinflammation are the presence of reactive astrocytes, activated microglia, increase in cytokines, chemokines, prostaglandins, and reactive oxygen and nitrogen species. Microglial activation has been related to accumulation of α-synuclein protein and proteosomal and lysosomal system dysfunction and these activated microglial cells have been reported to induce cell death in dopaminergic neurons. Furthermore, neurons over-expressing α-synuclein protein have demonstrated early activation of microglia and release of various inflammatory mediators such as IL-1, IL-6 and TNF-α and inflammation-related enzymes including cyclooxygenase-2 (COX-2) and inducible nitric oxide synthase. In addition, pro-inflammatory factors including IL-1, TNF-α, reactive oxygen/nitric oxide species and prostaglandins released from the activated microglia induced by α-synuclein protein can enhance oxidative stress and trigger cell-death pathways. These locally released inflammatory mediators also induce expression of cellular adhesion molecules in the endothelium of the BBB which help in the subsequent recruitment of immune cells from the periphery into the inflammatory site. Several studies have shown the infiltration of lymphocytes into the CNS in PD. These immune cells including leukocytes, macrophages, B cells and T cells, predominantly, CD4+ T cells and CD8+ T cells are recruited in the substantia niagra region of the brain in PD with CD4+ T cells being suggested to be cytotoxic in PD. However, the other subsets of T cells such as Treg cells are also present in the substantia niagra of PD and have the ability to suppress the cytotoxicity of effector T cells like Th-17 cells. These Treg cells can also induce apoptosis in activated microglia and reduce α-synuclein protein induced neurotoxicity. Although various studies have argued on either the pathogenic or the neuroprotective role of lymphocytes, the precise role of lymphocytes in PD is still elusive. Hence, it is evident that lymphocytes play a key role in PD but, their role in relation to PD pathology still needs to be explored extensively.

**Stroke**

The World Health Organisation (WHO) has defined stroke as the clinical syndrome of rapid onset of focal or global...
cerebral deficit, lasting more than 24 hours or leading to death with no apparent cause other than a vascular one. Various risk factors including smoking, diabetes, hypertension, atrial fibrillation and transient ischaemic attack have been identified for stroke which can lead to either haemorrhagic or ischaemic stroke. Haemorrhagic stroke is due to hypertensive arteriosclerosis and amyloid angiopathy whereas, the major cause of focal or ischaemic stroke is due to the occlusion of major arteries of the brain and accounts for 80% of all strokes. The occlusion in the artery within the CNS leads to a reduction of blood flow leading to insufficient oxygen and glucose supply to the brain. Usually, this occlusion develops due to thrombosis in situ (such as atherothrombosis of large cervical or intracranial arteries), or embolism from heart or blockage of small penetrating arteries due to arteritis or haematological disorders. In stroke, a series of neurochemical processes occurs which is termed an ischaemic cascade. During ischaemic cascade, various events take place including cellular metabolic failure due to hypoperfusion, oxidative stress, excitotoxicity, damage of BBB, microvascular injury, activation of haemostatic system and inflammation. These series of events result in non-selective cell death in the CNS including neurons, astrocytes, microglia, oligodendrocytes and endothelial cells. Apart from various risk factors including smoking, diabetes and hypertension, there are several studies showing a relation between inflammatory status and the risk of stroke which also has an effect on the outcome of stroke. The inflammatory response arises from the series of ischaemic cascades. The hypoperfusion occurs during this cascade causes a failure of energy-dependent ion pump leading to activation of calcium channels and release of excitatory neurotransmitters including glutamate into extracellular spaces. The release of excess excitatory neurotransmitters causes neuronal death which is termed as excitotoxicity. On the other hand, a disturbance in the scavenging of free reactive radicals causes oxidative stress. There is significant evidence illustrating the cytotoxic effects of free reactive oxygen and nitrogen species. This oxidative stress leads to glial cells activation followed by release of various inflammatory mediators including cytokines, chemokines and reactive free radicals, as well as expression of MHC I and II and co-stimulatory molecules. The level of IL-1 has been reported to be elevated after experimental stroke and has been a target for therapy in stroke in reducing inflammation-related damage.

Non-specific activation of peripheral T cells has been reported in stroke patients compared with healthy subjects, and the number of Treg cells is increased in stroke patients, similar to that described in animal models. During inflammation, endothelial cells of the BBB express various adhesion molecules including ICAM-1, ICAM-2 and VCAM-2 and selectins that help lymphocytes migration from the periphery to the CNS. These lymphocytes express chemokine receptors, α4β1 integrins and LFA-1 helping them in capturing, activation and transendothelial migration across the BBB. Several studies have shown infiltration of lymphocytes into the CNS following stroke. Real time in vivo imaging of the mouse brain after experimental stroke revealed a massive number of T cells infiltration into the stroke brain in comparison to sham. This recent study has reported two distinct populations of T cells in experimental stroke, the fast migrating T cells and slow migrating T cells but, the definitive role of these two populations in stroke is still under investigation. Furthermore, various studies have shown the neurotoxic effect in the stroke while at the same time others have argued on their neuroprotective role in the stroke. Taken together, to date there has been little agreement on the precise role of lymphocytes in the stroke and a better understanding of their contribution to stroke is still required.

**Role of lymphocytes in neurodegenerative diseases: pathogenic or neuroprotective?**

Infiltration of lymphocytes into the CNS during neurodegenerative diseases is well established and the molecular mechanisms underlying their recruitment into the CNS has also been well documented. However, the controversy of scientific evidence for the role of lymphocytes during neurodegeneration has raged unabated for more than a half century. It is now evident that after infiltration into the CNS and recognition of cognate antigen/MHC, peripherally activated lymphocytes (see section 1.3.1) can initiate inflammatory response in the CNS which can be either neuroprotective or neurotoxic. The pathogenic role of T cells has been demonstrated in neurodegenerative diseases causing neuronal death. However, the extent of the CNS injury during neurodegeneration has been correlated with the increase in T cells infiltration into the CNS suggesting the greater the infiltration, the greater the neuronal injury. It has been suggested that these infiltrating T cells can also mediate cell death and demyelination in neurodegenerative diseases, affecting other effector cells including microglia and/or macrophages. The adoptive transfer of T cells from spinal cord injury model mice and EAE-induced mice to healthy recipients develops paralytic disease which further supports the pathogenic role of T cells. It has been shown that during neurodegeneration and brain injury, both T cells and B cells are activated which is referred to as auto-reactive T cells or B cells. The number of auto-reactive T cells is increased in neurodegeneration and CNS trauma and they predominantly release IFN-γ and TNF-α. Moreover, these cytokines released by these auto-reactive cells can exacerbate ischaemia and excitotoxicity in the brain during neurodegeneration. Studies have also demonstrated that TNF-α induces cell death via apoptotic pathways and its concentration was also found to be elevated during neurodegenerative disease including PD. In addition,
activated CD4+ T cells express Fas-ligand (FasL), which has been reported to induce cell death via apoptosis in neurodegenerative diseases including EAE.162,163 These Fas and FasL are type I and II transmembrane receptors belonging to TNF/nerve growth factor and TNF families’ protein respectively.164 The infiltrating CD4+ T cells in PD induce apoptosis of dopaminergic neurons via FasL-Fas interaction, mediating FasL-mediated activation of microglia and neurodegeneration.165 The up-regulation of Fas and their ligands have been demonstrated in the CNS during neurodegenerative disease such as EAE leading to the apoptotic cell death.165 In addition, CD8+ T cells or cytotoxic T lymphocytes (CTL) are proposed to be involved in direct killing of neurons in a MHC-I dependent manner.166,167 The induction of MHC-I expression in neurons via IFN-γ has been documented and it has also been reported that the cytotoxicity of CTL in these neurons is mediated via either FasL-mediated neuronal apoptosis or perforin-dependent lysis of neurons.167-169 Moreover, both CD4+ T cells and CD8+ T cells have been reported to be equally necrototoxic and mediated via direct cell contact mechanism involving FasL, LFA-1 and CD40.170

Despite the proposed role of T cells in neurodegeneration, there is growing evidence for a beneficial or neuroprotective role of lymphocytes in neurodegenerative diseases.171-175 Adoptive transfer of auto-reactive T cells from EAE-induced mice to healthy recipient induces pathology.155 However, when these cells are transferred to the mice with partial optic nerve crush, a model for secondary neurodegeneration, they were found to be beneficial.173 Nerve cells from the mice which received auto-reactive T cells were found to survive well and were resistant to secondary neurodegeneration. Further analysis also revealed that only T cells specific to MBP were able to protect retinal ganglion cells from secondary damage and not T cells specific for non-self antigens, such as ovalbumin and heat-shock proteins. They further have suggested that only CNS-specific T cells are activated at the injured site to exert the neuroprotective response while T cells specific to non-self antigens fail to activate because of inadequate antigen recognition.173 In addition, after experimental axotomy of facial nerves, facial motor neurons are found to be severely impaired in severe combined immunodeficient (SCID) mice which lack T cells and B cells.176,177 but are restored up to wild-type controls after adoptive transfer of wild-type splenocytes containing T and B cells. Similarly, intraperitoneal injection of auto-reactive MBP-specific T cells in rat with an experimentally crushed spinal cord show early recovery with greater locomotor function as compared to controls.178,179 Moreover, an in vitro study in murine entorhinal-hippocampal brain slices shows down-regulation of the Th-1 cells induced inflammatory marker, ICAM-1 in microglia while another study demonstrated the neuroprotective role of both Th-1 and Th-2 cells.179,180

It is now well understood that upon activation by their cognate antigen/MHC-II, CD4+ T cells in the presence of appropriate mediators also differentiate into Treg cells and several studies have demonstrated the immunosuppressive role of Treg during neuroinflammation.105,107,114,181-185 Treg cells have an immunomodulatory role in human immunodeficiency virus (HIV)-1-induced neurodegeneration leading to neuroprotection by suppressing microglial activation and secreting neurotrophic factors.185 Treg cells are able to salvage neurons by suppressing the inflammatory response mediated by Th-17 cells.105 It has also been demonstrated that interaction between neurons and pathogenic T cells in EAE-induced mice cause conversion of pathogenic T cells into Treg cells which are able to suppress neurodegeneration induced by pathogenic T cells.184 The conversion of pathogenic T cells into neuroprotective Treg cells has been suggested to be induced by the interaction of CD4+ T cells and neurons via B7-CD28 and TGF-β receptor signalling pathways.184 In addition, IL-10 producing CD4+ T cells are found to be neuroprotective in stroke and it has been demonstrated that IL-10 and TGF-β can modulate immune processes by inhibiting Th-1-induced inflammatory responses as well as general inflammation leading to neuroprotection.147,186,187 These auto-reactive T cells are able to produce various neurotrophic factors including brain-derived neurotrophic factors (BDNF), neurotrophin-3 (NT-3) and glial-cell derived neurotrophic factors which can rescue neurons from neurodegeneration.172,188 It has also been suggested that T cells can instruct microglia to remove neurotrophic extracellular glutamate resulting in neuroprotection.189

Investigating the role of B cells in EAE-induced mice reveals that B cell deficient mice are more susceptible to EAE induction developing severe pathology with delayed recovery and early demyelination in compare to their controls.190,191 Likewise, a decrease or absence of B cells in EAE-induced mice correlates with an increase in severity of disease as well as influx of more pathogenic T cells into the CNS.192,193 In addition, IL-10 secreted by B cells is reported to be neuroprotective in EAE-induced mice since EAE-induced mice having IL-10 deficit B cells failed to recover and manifested persistent inflammatory responses.194 Hence, IL-10 specific B cells are suggested to be playing a crucial role in the recovery and progression of the disease.195 It has also been demonstrated that B cells are able to limit the spreading and severity of disease in EAE-induced mice.195 Furthermore, there is evidence showing that B cells are able to release various neurotrophic factors including BDNF, NT-3 and neurotrophic growth factor which have a possible role in contributing to neuroprotection in neurodegenerative diseases.196-198 In addition, recent paper has shown the neuroprotective role of lymphocytes during neurodegeneration either induced by excitotoxicity or glucose-oxygen deprivation which is mediated through the astrocytic activation and modulation of mitogen activated protein (MAP) kinases.199 IL-6 has been proposed to be one of the potential factors in lymphocyte mediated neuroprotection.199
Thus, it is clear that the notion of the brain as an absolute immune-privileged site is no longer appropriate, rather it can be considered as a site of active immune-surveillance. Recent evidence suggests the existence of bidirectional communication between these two systems and can influence each other via various mediators in both healthy and diseased condition of the CNS. In addition to this bidirectional communication, several studies have reported that lymphocytes are regularly patrolling the normal CNS in low number which can be increased upon recognition of cognate antigen and initiation of inflammation at the site. Whilst many attempts have been made to describe either the pathogenic or the neuroprotective role of lymphocytes, there has been little discussion about precise function(s) and mechanism(s) of these responses during neurodegeneration. Therefore, by defining the role and underlying mechanism(s) of lymphocytes and other related immune cells in neurodegenerative diseases, we certainly can set a milestone for the better understanding of the disease pathology and its therapy. Moreover, on this basis, the possibility of immune-based therapy of neurodegenerative diseases cannot be overlooked, though proper pros and cons should be determined beforehand.

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