Neurorestorative Roles of Microgliosis and Astrogliosis in Neuroinflammation and Neurodegeneration

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Abstract

The pathophysiological processes involved in neurodegenerative diseases have not been clearly defined. Nevertheless, a significant aspect of the proof focuses directly on the function of several mechanisms of inflammation. The immune system is represented in the central nervous system by the microglial cell capable of detecting harmful or foreign pathogens, and thus initiates self-activation and neuro-inflammatory processes via phagocytosis and cytokines release, to maintain the cellular microenvironment. Then, microglial cells can spawn an emphasis on persistent inflammation via phagocytosis and cytokines release, to maintain the cellular microenvironment. Then, microglial cells can activate the neurodegenerative cycle. The biomechanical properties of the brain, neuronal regeneration, and plasticity can be modified by reactive gliosis. Defining the inception and development of reactive microgliosis is vital for better clinical treatments design.

Keywords: Microglia, Astrogliosis, Neurodegeneration, Inflammation, Neurorestoration


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neurotoxicity and neuroprotection [13].

Introduction

Neurodegeneration is an encompassing word used to define several developments which result in the loss of function, structure, and death of neurons (Fig. 1). Substantial efforts are engaged in clarifying the effect of inflammatory mechanisms in neurodegeneration, which might help in formulating new therapeutics methods aimed at the immune-immune system to stimulate neuro-restoration. Neuroinflammation is known as a vital biological reaction of the body’s tissues to damaging stimuli relating to molecular mediators, blood vessels, and immune cells in the central nervous system (CNS). This process involves movement, accumulation, and attraction of cells supporting a flow of events involved in the impairment and revivification of tissue. In several defects of the central nervous system, the inflammatory response comprises of the activation of the glial cells [1, 2].

The inflammatory responses are prompted to combat infection, moderate injury, and any other stimuli which may include various cell types. The effect of this response depends on several factors released from these cells, possessing self-regulatory capability to repair damaged tissue and eliminate pathogenic elements. However, prolonged response results in chronic inflammation of the environment leading to progressive tissue damage [3]. A shred of important evidence supporting the connection between an increase in age and inflammation is described by an increase in inflammatory mediators for example interleukin (IL)-6 and -8 and also increase in C-reactive protein in humans with age, with the potential to increase in neurodegenerative diseases and obesity [4].

In the brain, microglia and astrocytes serve as the support of the immune response; to be precise, microglia cells constitute 10 to 15 percent of the brain, which varies regionally and preponderate in the midbrain regions for example the substantia nigra and hippocampus [6]. The blood-brain barrier, parted from the systemic immune system by the immune response of the brain depends on the microglia potential to react as a manifold of immune cells, detect toxins, cytokine levels, injury, and pathogens and also react in a neurotoxic or neuroprotective form akin to the macrophages in the systemic immune system [7].

In a neuro-toxic mode (Fig. 2), activated microglial cells respond to injury and insult by inducing pro-inflammatory cytokines to attract other astrocytes and microglia in response to bacterial infection and form extensive and diverse factors such as reactive nitrogen species (RNS), reactive oxygen species (ROS), lipid mediators, cytokines and also eliminate cellular wastes as a form of post-infection feedback via phagocytosis [8]. Thus, microglia self-protect against their personal toxic products via chains of antioxidant proteins been moderated by the nuclear factor erythroid 2-like protein 2 (NFE2L2) effects [9]. Microglia are involved in an increasing number of diseases that are central nervous system (CNS)-related; typically stimulated microglia are found in regions of the brain suffering from AIDS-associated Neurodegenerative disease [10].

Astrocytes give neurons functional and structural support, secrete glio-transmitters to control synaptic activity, and get involved in the formation and remodeling of the synapse [11]. They also have the potential to participate and respond to inflammatory in the brain [12]. Though inflammation and immune reaction are regular mechanisms for defense targeted at brain protection from injury, cellular debris, infection, or irregular protein accumulation. In response to several stimuli, activated astrocytes secrete numerous inflammatory and immune mediators for instance anti- and pro-inflammatory chemokines/cytokines that subsequently elicit neurotoxic or neuroprotective effects. In response to immune stimuli, some other glial cell-type of mesodermal origin activated microglia also partake in neurotoxicity and neuroprotection [13] by releasing anti- or pro-inflammatory chemokines/cytokines. These unique forms of microglia activation are known as functional polarization, with involvements in different disorders of the CNS, such as cerebral ischemia and spinal cord injury [14].

**Figure 1:** Neuro-inflammation and neuro-degeneration model [5].

**Figure 2:** Microglial activation pathology via prolonged systemic inflammation and neurodegenerative cycling [19].

**List of Abbreviations**

- APC: Antigen presenting cells
- BBB: Blood brain barrier
- CCL: Chemokine (C-C motif) ligand
- CD: Cluster of differentiation
- CNS: Central nervous system
- CCR: Chemokine (C-C motif) receptor
- CD: Cluster of differentiation
- CNS: Central nervous system
- CCL: Chemokine (C-C motif) ligand
- JAK-STAT: Janus kinase-signal transducer and activator of transcription
- MAPK: Mitogen-activated protein kinase
- MCP: Monocyte chemotactic protein
- NFE2L2: Nuclear factor erythroid 2- like protein 2
- NF-κB: Nuclear factor-kappa B
- NLRs: NOD-like receptors
- NMDA: N-Methyl-D-aspartate
- NOD: Nucleotide-binding and oligomerization domain
- ORM: Orosomucoid molecule
- RNS: Reactive nitrogen species
- ROS: Reactive oxygen species
- TGF: Transforming growth factor
- TLRs: Toll-like receptors
- TNF-α: Tumor necrosis factor alpha

**Neuro-inflammation and Microglia**

Microglial cells are located in the spinal cord and brain but specifically in the hippocampus and substantia nigra [15]. These cells are about 5 to 20 percent of the whole populace of glial cells in the CNS and are known to represent the immune system in the CNS, due to their ability to secrete cytokotoxic factors, act as antigen-presenting cells (APC), and exert phagocytic effect [16]. They are derivative of macrophages formed in the naive yolk sac via hematopoiesis, then translocate to the emerging neural tube where microglia are developed [17]. In normal conditions, microglia cells guard the brain surroundings by inducing a rapid reaction to alterations and successfully moderate inflammation. Several signs that jeopardize homeostasis of the central nervous system for example residues and/or structures from fungi, viruses, and bacteria; complement factors, cytokines, chemokines, antibodies, and abnormal endogenous ones, as well as endogenous ones, as well as...
proteins among others, are detected by the microglial cells and thus initiate their activation [15]. Therefore, there are 2 main functional parts of the microglia-maintenance of the central nervous system and immune defense [17].

Microglial cells possess two functional states of polarization—firstly is the phenotypical polarization to develop a typical pro-inflammatory or a different phenotype that is anti-inflammatory and pro-healing [18]. Hence, various anti- and pro-inflammatory cytokines, and other stimuli, can polarize microglia near distinct functional phenotypes.

Inflammatory phenotype of activated cells is described by the up-regulation of tumor necrosis factor (TNF)-α, CD16 Fc receptors, CD86, CD64, CD32, IL-1β, -6,-12,-23, chemokine and inducible nitric oxide synthase (iNOS), however microglia possessing anti-inflammatory phenotype exhibit upregulation of the insulin-like growth factor (IGF)-1, mannose receptor (CD206), arginase (Arg)-1,activating receptor expressed on chitinase 3-like 3 (Ym-1), myeloid cells 2 (RECM2), among others [18]. These factors are embodied in the stimulation of microglial cells leading to the additional generation of some inflammatory mediators and cytokines, capable of contributing to apoptosis of the neurons in manifold neurodegenerative diseases. With these features, microglial cells are considered as the local immune cells in the brain, and also engenous to minor changes in central nervous system equilibrium and then readily activated during most neuropathological conditions.

Among other things, a significant point of note is the starting of the destruction process microglial cells tend to act. In the early stage of Alzheimer's disease, there is an increase in the activation of microglial; indicating that microglia initiates the removal of dangerous elements contributing to the disease like amyloid-β plaques. Hence, an acute neuro-inflammatory response is believed to have a beneficial effect on the central nervous system, with the ability to reduce damage and repair damaged tissue. Besides, microglia have the potential of removing glutamate, which is a recognized neuro-toxic substance that exerts its actions on NMDA (N-Methyl-D-aspartate) receptors from the neurons and then results in the death of the neurons. In Alzheimer's patients, the significance of glutamic acid and related microglia cell role was evidenced by the therapeutic actions of memantine drug (NMDA receptors antagonists) that enhances the cognitive potential and daily functions [20]. Meanwhile, it is imperative to note that microglial cells can be triggered by endogenous proteins or with environmental toxins, resulting in over-activation of cells and generation of ROS and TNF neuronal toxicity [21].

**Neuro-inflammation and Astrocytes**

Astrocytes, the fewest abounding and diverse kind of glial cells in the CNS can change their morphology based on their localization, subtype, and developmental stage [22]. For instance, the protoplasmic ones are the gray matter astrocytes, displaying short branches, while the white matter fibrous astrocytes show long unbranched methods [23]. The astrocytes support the components of the neurons in the neural tissue and microglial cell and also respond to insults of all forms in the central nervous system through the reactive astrogliosis; this procedure is sensitive and dependable biomarker of diseased tissues. These cells are accountable for multifaceted and vital functions in a healthy central nervous system, such as their involvement in the primary functions in the processing of information by neural circuits and synaptic transmission [12] and are also involved in synaptogenesis and dynamically control signal transmission, information processing, synaptic and neural plasticity, and also give metabolic support for neurons [24]. As a result, astrocytes have been proven to partake in some essential processes for example regulating environmental homeostasis of ions, pH, the flow of blood, reducing oxidative stress [25], and also accountable for a huge sum of equilibrium tasks in the central nervous system [26]. With these abilities, microglia cells and astrocytes, behave like the core effectors of the neuro-inflammatory feedback. After trauma or sense of destruction signal, astrocytic cells together with microglia speedily respond to pathology and undertake significant changes in their functioning and morphology [24]. Therefore, the feedback objective is to regulate and eliminate the insult on the brain, but this feedback could have lethal outcomes. Reactive gliosis is a self-sustaining activity that eventually aggravates injury and also signifies a non-physiologic condition in which the astrocytes may forgo their supportive effects [25]. The process that leads to the stimulation of the cells remains uncertain, but several factors tangled in brain damages can stimulate their feedback. For instance, the presence of amyloid in Alzheimer's disease has been confirmed to activate astrocytes. Similar to microglia, astrocytes can also phagocytose and destroy amyloid-β, and to bring this ability, microglia and astrocytes are triggered via TLRs (Toll-Like Receptors) and RAGE (receptor for advanced glycation end-products), therefore initiating local inflammation [27]. Once the astrocytes response is triggered, there is a change in their morphology and significantly intensifies the expression of the glial fibrillary acidic protein (GFAP), which is a known indicator of astrocytes reactivity [28]. The changes result in a disorder in the astrocytes normal activities, which are important for the normal function of the neurons. Internally, astrocytes activation includes stimulation of transcription factor- NF-kB, regulating the release of adhesion molecules and chemokine, and hence favors outer lymphocyte penetration and upregulating inflammatory response resulting in neurodegeneration [27]. Besides, the transcriptional activity of NF-kB inhibition in astrocytes can broadly decrease inflammation, therefore signifying NF-kB inhibition in the astrocytes as a possible therapy for diseases like Alzheimer’s disease [30].

With this evidence, it has been confirmed that activated astrocytes have the potential to initiate neurodegeneration; furthermore, activated astrocytes display inflammation-related factors, for example, S100β peptide, which represent a key factor for neuro-inflammation. The S100β protein, a neurtrophin produced by astrocytes, is responsible for the development, function of neurons, and survival under physiological conditions. The peptide S100β is over-expressed and is identified with the pathology progression of subjects with severe brain trauma and neurodegenerative diseases [27].

**Crosstalk between Astrocyte and Microglia**

Microglia, the major immune cells in the CNS, partake in many neuro-pathological conditions and, in synergy with astrocytes, aid the recovery of the CNS from injury and stress [31]. Due to their bi-directional conversation and autocrine feedback, during the injury modulation in the CNS, the cross-talk amid astrocytes and microglia has come into the vanguard of glial research and electromagnetic force therapy utilized in inducing neuro-restoration. At the epicenter of this mutual adjustment is the control of microglial cells of the innate immune roles of astrocytes, influencing their role, either neuro-toxic or neuroprotective [31]. Microglia, activated before astrocytes, release NADPH oxidase derived hydrogen peroxide (H2O2) [32], complement component 1q (C1q), tumor necrosis factor (TNF)-α, and interleukin (IL)-1 β (Fig. 3) [33] to control astrocyte stimulation and promote A1 reactive astrocytosis which is the major destructive mechanism of astrocytes activity.

![Figure 3: Schematic diagram showing Neuro-inflammation-mediated neurodegeneration in the brain](https://doi.org/10.54117/sjmams.v1i1.1)

This enables the microglial cell the primary target for electromagnetic force (EMF) therapy of the central nervous system. Though astrocitic
cells are known as neuronal supportive cells, helping with the regulation of the CNS homeostasis, several studies describe astrocytes' function in the microglial functions and cytokine regulation, and the inborn immunity response in the CNS [35]. Microglia cross the CNS for removal of pathogens, repair the tissue, or scar formation like (TLRs), complement, mannose, NOD2, and Toll-like (TLRs), astrocytes take a close cross-talk with the neighboring microglia in the CNS for removal of pathogens, repair the tissue, or scar formation induction [37]. Likewise, recent studies have established the astrocyte-microglia cross-talk is an important step in the innate reaction to injury in inflammation in the CNS, demonstrating that the astrocyte's response to toll-like receptor (TLR)-2, 3, and 4 is significantly boosted by or directly associated to, the existence of microglial cells within nearby tissue [38]. This shows an important interaction between the microglia and astrocytes in neuro-repair and neuro-restoration. Simply through cautiously controlled interactions of astrocytes and macrophages that react to inflammation can be resolved and controlled. A1 astrocytic cells are pro-inflammatory, showing gene upregulation possibly damaging to synapses, and are initiated by microglia release of interleukin (IL)-α, TNFα, and complement component (C1q). A2 reactive astrocytes release proteins that stimulate synaptogenesis of the CNS [18]. Also, as are triggered under ischemic states: A2 astrocytes have neurorestorative and -protective effects and display the phenotypes required to be induced for electromagnetic force (EMF) therapy to success leads to neuro-regeneration [38]. Working synergistically with astrocytic cells, the highly abounding cells in the CNS, microglia –typically function as chaperones to the astrocytes, controlling astrocyte's innate immune effects at pathological states by secreting molecules having an impact on the intracellular signal transduction via MAPK and JAK-STAT pathways. To further increase the inflammatory response, microglial cells upregulate the nuclear factor kappa B (NF-kB) signaling pathway, and upon triggered, astrocytes augment pro-inflammatory gene expression, and promote the formation of pro-inflammatory growth factors, chemokines, and cytokines. Then, astrocytes control microglial cell phenotypes and functions via the astrocyte-derived factors, complemented proteins, cytokines, and chemokines [15].

Conclusion
The origin and various causes of CNS microglia and astrocytes activation, amplification, and termination are vital in unraveling treatment for a well-modulated and maintained systemic immune system response. Seeing it as a possible therapeutic step, inhibiting the inflammatory process is a potential therapeutic strategy to combat inflammation-induced neurodegeneration.

References


