HMGB1-TLR Signaling in Rasmussen’s Encephalitis

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Abstract

Rasmussen’s encephalitis is neurological disorder of childhood characterized by uni-hemispheric inflammation, intractable focal epilepsy and progressive cognitive and neurological deficits. Currently, the pathogenesis of Rasmussen’s encephalitis is still enigmatic and hemispherectomy is the only effective method to control the seizures associated with Rasmussen’s encephalitis. Recently data indicated that intrinsic activation of endogenous pro-inflammation high-mobility group box-1 (HMGB1) and Toll-like receptor (TLR) is involved in the development of Rasmussen’s encephalitis. Activation of HMGB1-TLR signaling plays a critical role in brain inflammation, development of epilepsy and cognitive dysfunction. Targeted therapy on HMGB1-TLR signaling might be a novel strategy with anti-inflammation, anti-epilepsy as well as improving cognitive dysfunction associated with epilepsy in Rasmussen’s encephalitis.

Keywords: Rasmussen’s encephalitis; Epilepsy; Inflammation; Cognition; HMGB1

Introduction

Rasmussen’s encephalitis is a very rare chronic progressive inflammatory neurological disorder of uncertain etiology affecting mostly children and associated with hemispheric atrophy, focal epilepsy (epilepsia partialis continua), cognitive deterioration and progressive neurological deficits, resulting from progressive loss of function subserved by the involved cerebral hemisphere [1-3]. The etiology and pathogenesis of Rasmussen’s encephalitis, in particular, the factors responsible for the characteristic of asymmetry are still elusive. Currently, hemispherectomy is the only effective method to control the seizures associated with Rasmussen’s encephalitis. Increasing experimental and clinical evidence supports a link among inflammation, epilepsy (both in terms of epileptogenesis and the long-term consequences of seizures) and cognitive dysfunction associated with epilepsy, which indicates that activation of inflammatory processes in the brain is a common feature of various epileptic disorders [4]. Recently study demonstrates that intrinsic activation of endogenous pro-inflammation high-mobility group box-1 (HMGB1) and Toll-like receptor (TLR) are involved in the development of Rasmussen’s encephalitis [5]. Activation of HMGB1-TLR plays a critical role in brain inflammation, development of epilepsy and cognitive dysfunction, and inhibition of HMGB1-TLR signaling demonstrates anti-inflammation, anti-epilepsy as well as improving cognitive dysfunction associated with epilepsy [4].

HMGB1-TLR Signaling and Brain Inflammation

HMGB1-TLR may represent a novel pro-inflammatory axis following sterile brain injury [6]. Activation of proinflammatory cytokines especially HMGB1-TLR signaling involves in immune response to inflammation associated with many neurodegenerative diseases, including Alzheimer’s disease, Parkinson’s disease, amyotrophic lateral sclerosis, multiple sclerosis and epilepsy [4,7], and subserve neuromodulatory functions implicated in brain physiology and may contribute to acute and chronic neurodegeneration. HMGB-1, characterized as a key cytokine [8], is a 216 amino acids (29 kDa) DNA-binding protein with a highly conserved structure in several species [9]. HMGB-1 participates in nucleosome formation and regulation of gene transcription [10,11], including proinflammatory genes [12]. HMGB1 acts as a “danger signal” and alerts the immune system to damaged or dying cells. In response to inflammatory stimuli, HMGB-1 is secreted by activated macrophages [13], natural killer cells [14], myeloid dendritic cells [8] and astrocytes [15] binding to RAGE and TLR, including TLR2 and TLR4. The hyperacetylated form of HMGB1 regulates transcription of various pro-inflammatory cytokines, including IL-1β, through binding to TLR2, TLR4 and also to RAGE [12,16]. In human brain cortex, HMGB1 immunoreactivity expressed in nuclei of neurons and glia cells as reported by Zurolo et al. in 2011 and Luan et al. in 2016. When under immune/inflammatory challenges or injurious conditions, HMGB-1, normally residing in nuclei, translocates to the cytoplasm and/or extracellular space [12,17]. Extracellular HMGB1 acts as a ‘danger signal’ to orchestrate a homeostatic defensive response in challenged tissues; however, under circumstances that remain poorly understood, the resolution of inflammation is compromised, its pro-inflammatory properties, acquired upon its cellular release and consequent TLR/RAGE stimulation, appear to contribute to the pathogenesis of various inflammatory and CNS diseases [18].

TLR4 and TLR2 are preferentially expressed on cells that play central roles in innate immune responses, including macrophages and microglia, but it is present at very low or undetectable levels on neurons and astrocytes [5,19,20]. As the major initial sensors of danger or stranger signals recognized by TLR4, Microglia secretes...
HMGB1/TLR Signaling and Epilepsy

Extensive experimental and clinical evidence supports a link between inflammation and epilepsy, both in terms of epileptogenesis and the long-term consequences of seizures, which indicates that activation of inflammatory processes in the brain is a common feature of various epileptic disorders [4,16,23,24]. Therefore, brain inflammation is regarded as a diagnostic, prognostic or therapeutic biomarker for epilepsy [25]. Recent studies demonstrate that activation of HMGB1/TLR signaling pathways plays a crucial role in the development of epilepsy. Activation of HMGB1–TLR signaling has been recently demonstrated in epileptogenic brain tissue from drug-resistant epilepsy patients with Rasmussen’s encephalitis [5], the prototype of inflammatory and immune-mediated epileptic encephalopathy, but also with temporal lobe epilepsy [15,16] and focal malformations of cortical development [19] without infectious or immune-mediated etiology. In particular, antagonists of HMGB1 and TLR4 retard seizure precipitation and decrease acute and chronic seizure recurrence, which highly suggest that activation of HMGB1–TLR-RAGE pathway possibly represents a common mechanism of epileptogenesis [26].

Rasmussen’s encephalitis is a rare chronic progressive inflammatory neurological disorder with pharmacoresistant focal epilepsy (epilepsia partialis continua) [1,2,27]. There might be a linking between activation of the innate immune system and consequent inflammation to epilepsy of Rasmussen’s encephalitis. The chronic inflammation is—at least partly—the consequence of a cycle in which inflammation induced cell injury leads to the release of endogenous danger-associated molecular pattern that drives the inflammatory response, causing further damage and inherent epileptogenicity of brain lesions, pathogenic or otherwise [6]. The intralesional overexpression and cellular distribution of HMGB1 and its cognate receptors TLR2, TLR4 and RAGE in specimens from Rasmussen’s encephalitis patients with medically intractable epilepsy has been demonstrated (5): (i) Cytoplasmic translocation of HMGB1 was detected in the neurons, reactive astrocytes and reactive microglial cells within the lesions; (ii) Increases in the immunoreactivity of HMGB1 in reactive astrocytes (predominant in cytoplasm) were observed; (iii) Intralesional expression of RAGE, TLR4 and TLR2 was also elevated in both neurons, reactive astrocytes and reactive microglial cells, particularly in reactive astrocytes. Although HMGB1 release and signaling may be a general feature of all epilepsies, expression of HMGB1, TLR2, TLR4 and RAGE was more markedly increased in perivascular areas, and

endothelial cells in walls of blood vessels within the lesions cortex displayed immunoreactivity in Rasmussen’s encephalitis patients. These findings are concomitant with the reactive astrogliosis, neuron loss and inflammation (i.e. CD8-positive, CD3-positive T lymphocyte). The evidence further supported the role of HMGB1–TLR pathway in activation of immune and endothelial cells in the pathogenesis of Rasmussen’s encephalitis [5].

Transcriptional activation of genes can be triggered by HMGB1 via nuclear factor-κB and/or activator protein–1-dependent transcription of inflammatory genes, which may perpetuate brain inflammation and contribute to long-term molecular plasticity involved in epileptogenesis and neuropsychiatric comorbidities (Figure 1). It has been shown that repressor element 1-silencing transcription factor (REST) may control the fundamental transcription patterns that drive circuit excitability, seizures and epilepsy via recruitment of histone methyltransferase G9a [28]. As G9a is essential for the establishment and maintenance of H3K9me2 [29], aberrant distribution of heterochromatin marker and alteration of chromatin structure may directly leads to the initiation and deterioration of seizures and epilepsy. Interestingly, as histone modifier G9a also contributes to the maintenance of genomic imprinting [30], the REST-G9a-imprinted genes axis may explain the genetic tendency in the family suffered seizures and epilepsy. Therefore, understanding the epigenetic mechanisms involved in seizures and epilepsy may shed light for a better treatment of these disorders.

Figure 1: HMGB1–TLR signaling pathway in epilepsy. HMGB1 can activate specific receptors TLR expressed by glia and neurons, inducing rapid non-transcriptional effects on voltage-gated and receptor-gated ion channels, neurotransmitter release and glutamate receptors leading to increased neuronal excitability. Transcriptional activation of genes can also be triggered by HMGB1 which may perpetuate brain inflammation and contribute to long-term molecular plasticity involved in the development of epilepsy and neuropsychiatric comorbidities related with epilepsy. Abbreviations: HMGB1, high-mobility group box 1; TLR, Toll-like receptor; NFκB, nuclear factor kappa B; AP1, activator protein 1.
HMGB1/TLR Signaling and Cognition

Brain inflammation has been implicated in the pathophysiology of several neuropsychiatric conditions. Therefore, inflammatory processes which are triggered in the brain by an epileptogenic insult may, concurrently with seizures, lead to the development of neuropsychiatric abnormalities [4]. As a crucial regulator mediating epilepsy development and brain inflammation, HMGB1/TLR signaling also has been implicated in the pathophysiology of cognitive and psychiatric phenotypes. RAGE-mediated inflammatory pathway plays an important negative regulator of learning and memory. In particular, the increased RAGE signaling has been implicated in mechanisms of memory impairments in Alzheimer’s disease, and RAGE is regarded as a potential therapeutic target to ameliorate cellular dysfunction and activation of immune and endothelial cells in the pathogenesis of processes which are triggered in the brain by an epileptogenic insult [31]. Recent evidence indicated that HMGB1 and its corresponding receptors are upregulated in the inflammation pathway in epileptic tissue and may play a crucial role in the development and perpetuation of seizures and cognitive dysfunction associated with epilepsy [4,15]. To elevate the brain levels of HMGB1 via intracerebroventricular administration has been proven to precipitate seizures and disrupted object memory mediated by either TLR4, or RAGE [32]. Refractory focal seizures and cognitive deterioration are the clinical features of Rasmussen’s encephalitis, a rare progressive inflammatory disorder of uncertain etiology [1,2]. Upregulation of HMGB1, TLR4 and RAGE in reactive microglia, reactive astrocytes, neurons and vascular endothelial cells has been demonstrated in the brain lesion area of Rasmussen’s encephalitis. Therefore, HMGB1-induced overstimulation of RAGE or TLR4 might contribute to memory deficits in Rasmussen’s encephalitis.

Currently, we still have no clear answer for the etiology and pathogenesis of this destructive inflammatory response in Rasmussen’s encephalitis; in particular, the factors responsible for the characteristic of asymmetry are still elusive. We are uncertain of the possible initial antigens to trigger the inflammatory response and why the process is unilateral. So far, hemispherectomy is the only effective method to control the seizures associated with Rasmussen’s encephalitis. To date anti-inflammatory therapy has not produced lasting responses in Rasmussen’s encephalitis, such as corticosteroids pulses and intravenous immunoglobulins. Therefore, imperfectly effective immunological interventions might unacceptable postpone hemispherectomy. Evidence demonstrated HMGB1-TLR pathway in activation of immune and endothelial cells in the pathogenesis of Rasmussen’s encephalitis. Therefore, perhaps more specifically targeted therapy such as targeting on HMGB1-TLR might produce lasting and promising clinical responses.

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