Role of Platelets in Glutamate Mediated Excitotoxicity: An Overview

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Received date: July 07, 2015; Accepted date: August 10, 2015; Published date: August 17, 2015

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Abstract

Glutamate, the main excitatory neurotransmitter in central nervous system (CNS), binds primarily to two types of receptors on the post-synaptic membrane of neuronal cells, namely metabotropic receptors and ionotropic receptors. Metabotropic receptors are G-protein coupled receptors while ionotropic receptors are non-selective ligand-gated ion channels, which allow movement of cations like Na+, K+ and Ca2+. The three most studied ionotropic receptors are N-methyl-D-aspartate receptor (NMDAR), α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor (AMPA), and Kainate receptor.

Glutamate synapse plays a pivotal role in many neurodegenerative disorders like Huntington disease, trauma, epilepsy, Alzheimer and Amyotrophic lateral sclerosis, and cerebral ischemia. Understanding the mechanism of these glutamate ligand-gated ion channels (LGIC) can assist in the development of therapy against ischemic stroke, a leading cause of death and disability worldwide.

This review focuses on how aggregated platelet microthrombi cross blood-brain barrier, reach neural parenchyma and release glutamate. Accumulated glutamate hyperstimulates glutamate LGIC, thus leading to neurotoxicity and apoptosis of neuronal cells.

Introduction

Stroke is one of the major challenges faced by almost all the developing countries. Stroke has great impact worldwide as it is one of the capital causes of death and affliction. WHO endeavored to define stroke as "Rapidly developing signs of focal (or global) disturbance of cerebral function, lasting more than 24 hours or leading to death, with no apparent cause other than that of vascular origin"[1].

According to the recent report submitted to WHO, the rate of stroke is 41/100,000 population in Nigeria and 316/100,000 population in Dar-es-salaam. In Europe stroke incidence rate is 101.1-239.3/100,100 men population and 63.0-158.7/100,000 women population [2]. The prevalence of stroke in India in recent years has been around 84-262/100,000 in rural and 334-424/100,000 population per year in urban areas [3].

Ischemic stroke becomes a higher risk, when platelet aggregates (primary injury) release glutamate and elevate amino acid concentration in the body fluid. The released glutamate interacts with glutamate receptors on endothelial cells of blood-brain barrier leading to breakdown of the barrier. Glutamate excitotoxicity is the final pathway that results in either clinical deficits or complete loss of the functions of neuronal cells (secondary injury).

Parallelism between Platelet and Neuron

Sharing the common embryonic origin i.e. ectodermic, platelets and neurons show a strong resemblance among themselves. To study neuronal molecular mechanism, the most significant model is platelet because it has morphological and functional similarities with neuron. Serotonin transport mechanism is similar in platelet and neuron [4]. Other neurotransmitters like dopamine, gamma-aminobutyric acid (GABA), and glutamate are secreted by both the cells [5]. Glutamate receptors are present on neuronal as well as non-neuronal tissues like bone marrow, kidney, heart, pancreas and platelets [6].

Platelets have been used in understanding the mechanism of various neurological disorders like migraine, Parkinson, autism, schizophrenia and Alzheimer’s [7]. Alzheimer is caused due to senile plaque formation in the brain by amyloid β and 95% of circulating amyloid-precursor protein in blood is contributed by platelets. We have shown that when platelets are treated with Aβ25-35, small GTPase RhoA gets activated and in turn activate platelets by modulating actomyosin, leading to cytoskeleton reorganization [8].

Glutamate Receptor

Glutamate is the main excitatory neurotransmitter in vertebrate CNS and the most abundant amino acid in diet. It plays critical role in many neurophysiologic functions like memory, learning, cognition, mood alteration and executive functioning. Glutamate has the post-synaptic effect on two broadly classified glutamate receptors, i.e., ionotropic and metabotropic receptors. The former are G protein-coupled receptors, which recruit second messengers such as diacylglycerol and cAMP to modify neuronal excitability. Ionotropic glutamate receptors are non-selective cationic ligand gated ion channels having significant role in excitatory signaling. Mainly three classes of receptors fall in this category viz. NMDAR, AMPAR, Kainate receptors, named after responsiveness and affinity to preferred agonists, N-methyl –D-aspartate, a amino-3-hydroxy-5-methyl-4-...
isoaxoepropionic acid, kainic acid, respectively. NMDAR is a major subfamily of glutamate receptors that favors Ca2+ inflow but also allows the passage of monovalent cations like Na+ and K+. Different subunits of NMDAR are expressed that include GluNR1, GluNR2A, GluNR2B, GluNR2C, GluNR2D, and GluNR3A. Two subunits of GluNR1 integrate with either same or different GluNR2 or rarely with GluNR3A subunit to form tetrameric complex. NMDAR needs ligands like NMDA or glutamate binding to GluNR2 subunit and co-ligands like glycine or D-serine binding to GluNR1 or GluNR3 subunit for its opening [9].

Another significant receptor is AMPA receptor, a heterotetramer complex composed of GluR1-R4 subunits. Intact receptor complex is Ca2+ impermeable ion channel, whereas absence of GluR2 subunit from the complex increases permeability for Ca2+ and Zn2+ by 2 to 3 folds. One of the calcium-activated cytosolic enzymes is protein kinase c-α (PKCa). Once activated in presence of Ca2+, PKCa forms complex with Protein interacting with C kinase 1 (PICK1) and translocate to plasma membrane where it phosphorylates GluR2 subunit of AMPAR at serine 880 residue. Phosphorylation of GluR2 triggers its endocytosis, thus making AMPAR Ca2+ permeable [10].

Primary and Secondary Neuronal Injury

Platelet aggregation leads to blockage of arteries supplying to brain, thus depriving brain of oxygen and nutrients. It results in drop in energy charge in cells and neuronal cell death (primary injury) (Figure 1). Recently it has been demonstrated that during the course of primary injury platelet activation/aggregation amplification is regulated by NMDA receptors. Platelet aggregation is inhibited by antagonists of NMDA receptor like MK801 hydrogen maleate, 3, 5-dimethyl-1-adanantamine hydrochloride (memantene), D-2-amino 5-phosphopentanoic acid (AP5), while NMDAR agonists like L-glutamate, glycine, NMDA facilitate platelet activation [11].

It is hypothesized that platelets on aggregation release glutamate, which mediates excitotoxic brain injury and neuronal dysfunction. Glutamate is stored in platelet dense granules and is released when platelets form aggregates. Platelets express vesicular glutamate transporters (VGLUT), through which they release glutamate after formation of microthrombi [12].

Thrombin-evoked release of glutamate from platelets has been quantified. Platelets release glutamate within 60 seconds of thrombin treatment, raising glutamate concentration from 175 µM to a maximum of 425 µM. Blood brain barrier becomes permeable after subarachnoid hemorrhage and delayed cerebral ischemia. It might be possible that when stimulated platelets form thrombus, glutamate released from them binds to and stimulates NMDAR expressed on endothelial cells. It leads to disintegration of blood-brain barrier and decamping of platelets into neuronal parenchyma, thus mediating glutamate excitotoxicity by exposing neuronal cells to glutamate. Neurotoxicity or secondary injury is caused when released glutamate binds to its ionotropic receptors like NMDA and AMPA on neuronal cells, which leads to disturbance in ionic homeostasis. Secondary injury to neurons can either be due to clinical deficits or neuronal programmed cell death (Figure 1). There is a direct relationship between microthrombosis and changed neuronal physiological properties. Such changes can result in difficulties with memory, consolidation, confusion and executive functions [13,14].

Neurotoxicity Mediated by Excessive Glutamate Accumulation

Hyperstimulation of glutamate LGIC disrupts the ionic homeostasis. A transporter protein, Na+-K+-Cl- (NKCC1) co-transporter 1, of neuron is stimulated in presence of glutamate and disrupts ion homeostasis in ischemic condition. NKCC1 allows the entry of Na+ and Cl−, thus causing ionic imbalance. Due to increased concentration of ions, water accumulates inside cell and it eventually leads to cell death. NKCC1 activity is stimulated by activation of iotonic and metabotropic glutamate receptors. Thus excitotoxicity caused due to excessive Cl− and Na+ ions can be averted by either removing these ions from extracellular space or by blocking glutamate receptors [15].

Intracellular calcium is maintained at low concentrations (µM) as compared to extracellular calcium. Endoplasmic reticulum and mitochondria constitute important Ca2+ stores inside cell. Ca2+ homeostasis of cell is disturbed when glutamate receptor is hyperstimulated. Glutamate ionotropic receptors assist entry of cations like Na+ and Ca2+. Increased Na+ concentration inside cell depolarizes cell membrane resulting in opening of voltage-gated Ca2+ channels, which further increase intracellular calcium level [16]. Higher level of intracellular calcium activates various enzymes like protein kinase II, phospholipases, proteases, nitric oxide synthase and ornithine decarboxylase. These enzymes have different consequences on cell. Activated phospholipase A2 persuades the genesis of arachidonic acid and platelet-activating factor (PAF, 1-O-alkyl-2-acetyl-sn-3-phosphocholine). PAF induces glutamate release and arachidonic acid impairs reuptake of glutamate from synaptic cleft, resulting in hyperstimulation of glutamate receptor and neurotoxicity [17]. As a result of impairment in reuptake of glutamate, glutamate receptors are
further stimulated, enhancing more production of arachidonic acid. High level of arachidonic acid generates ROS, which promote protein degradation, DNA fragmentation, and lipid peroxidation of neuronal cells [18].

    Raised cytosolic Ca2+ also stimulates nitric oxide synthase, which induces nitric oxide (NO) synthesis. Being thermodynamically unstable NO reacts with other gaseous molecules, anions and ROS to form nitrates, nitrites and peroxynitrites. Nitric oxide and superoxide radicals combine to form peroxynitrite, which is a highly reactive oxidant and not easily eliminated by antioxidant system. Peroxynitrite has very short half-life of approximately 1-2 seconds and when it degrades it forms many neurotoxic products [19]. Peroxynitrite oxidizes nucleic acids, cytoplasmic proteins and also inhibits mitochondrial respiration by inducing oxidation of electron transport chain (ETC) complexes. Inhibition of ETC complexes leads to formation of ROS, which finally induces apoptosis [20]. NO also evokes apoptosis of neuronal cell by increasing Bax/bcl2 gene expression. The proapoptotic factors, Bax and bcl2, promote cytochrome c release from mitochondria into cytoplasm and activate caspase 3, stimulating the programmed cell death [21].

**Conclusion**

As the mechanism of signaling of glutamate receptor on neuron and platelet is not fully understood, its role in secondary injury after cerebral ischemia remains unclear. Glutamate excitotoxicity caused by platelet aggregates plays significant role in cerebral ischemia. Glutamate ionotropic receptors like NMDAR and AMPAR trigger secondary excitotoxicity by deregulating ionic homeostasis, inhibiting mitochondrial respiration, and generating reactive species of oxygen and nitrogen. Although drugs targeting NMDAR have been developed, they are associated with many deficits and side effects like hallucination, memory loss, ataxia, nightmares and catatonia [22]. Thus divulging the signaling mechanism of glutamate ionotropic receptor system can assist in drug intervention and restoring glutamate mediated excitotoxicity.

**References**


