Synaptic alterations as a common phase in neurological and neurodevelopmental diseases: JNK is a key mediator in synaptic changes

Brain synapses play a key role in neuronal communication; this “conversation” is at the basis of properly functioning synaptic dysfunctions that lead to brain disorders. We study the modulators of this crucial synaptic function and we here present the evidence supporting the c-Jun N-terminal kinase (JNK) pathway as a pivotal actor in this scenario.

The term “synapse” is derived from the Greek σύν ("together") and νέων ("tap", "connect") meaning “conjunction”. Synapses are highly specialized cellular connections that represent the basic structural units of neuronal communication. There are two different types of synapses: in electrical synapses, presynaptic and postsynaptic cell membranes are connected by special junctions called gap junctions, channels that allow ions to pass from one neuron to another. In chemical synapses, representing the majority of synapses in the mammalian nervous system, the electrical activity in the presynaptic neuron is converted into the release of a chemical neurotransmitter that binds to receptors located in the plasma membrane of the postsynaptic cell. Therefore, two separate cellular components build up the chemical synapse: the pre-synaptic element, which specialized in the release of the neurotransmitter, and the post-synaptic element, which binds the chemical compounds coming from it into an electrical signal. The pre- and postsynaptic elements are associated and mirror each other changes, so that changes in one element will correspond to changes in the second.

What is described above identifies the old “bi-partite” synapse, more recently updated with the addition of a third structural element: tiny astrocytic processes. These astrocytic processes respond to synaptic activity and, in turn, participate in the regulation of synaptic transmission. Nowadays we also consider a fourth cell type, the microglia that makes brief, repetitive contacts with synapses. These dynamic interactions take place in health but also in pathological conditions, like Alzheimer’s disease (AD), autism, and immune molecules participate in eliminating the dysfunctional synapses (Wilton et al., 2019).

Synapses, both excitatory and inhibitory, are fundamental in shaping brain function and undergo dynamic changes. The dynamic changes of synapses are mainly studied by the visualization of the post-synaptic structure called spines. In fact, by following spines in the living brain it was discovered that they were kept in a state of equilibrium guaranteed by their continuous formation and elimination, resulting in decreased or increased spine turnover, or temporarily shifting to facilitate their additions or eliminations (Holtmaat et al., 2005).

It is therefore very important to keep in mind that a single spine is in a constant evolution; it changes its shape according to the needs and activity of the pre- and postsynaptic neurons and of the neuronal network. This phenomenon is known as synaptic plasticity, a very important feature of both inhibitory and excitatory neurons, by which they can adapt themselves to environmental changes. Inhibitory synapses are difficult to visualize and are less studied, thus resulting in biology modifications poorly understood compared to the excitatory synapses. However, inhibitory synapses are fourfold more dynamic than their shaft counterparts (Chen et al., 2012).

The plasticity of the brain is fundamental to preserving its adaptive functionality at the basis of learning and memory, but also stress stimuli can induce structural changes at the synapse level. Growing evidence suggests that, in brain diseases, the first neurodegenerative mechanism takes place at the synapse. It is, therefore, crucial to understand the intracellular mechanisms underlying synaptic modulation to design powerful neuroprotective strategies.

With this in mind, we decided to examine synaptic dysfunction as a common alteration of neurobiological mechanisms from neurodevelopmental (Angelman and Rett syndromes) to chronic neurodegenerative diseases (Alzheimer and Tauopathy).

Plasticity is a central feature in the developing as well as in the adult brain. Several evidence show that neurodevelopmental disorders, especially Autism Spectrum Disorders (among them Angelman, Rett and Dravet syndromes, X fragile, etc.), are characterized by too many synaptic connections causing hyper connectivity of brain circuits. On the contrary, studies on aging and neurodegenerative brain diseases revealed an opposite trend with loss of synaptic connection (Penzes et al., 2011).

In this context, synaptic dysfunction, which usually precedes neuronal death, has been linked to many neurodegenerative diseases, such as Parkinson’s disease, Huntington’s disease, and AD, as well as neuropsychiatric diseases. Many mutations in the human synapse proteome (the “synaptome”) have been described to underlie psychiatric and neurological disorders.

Synaptic dysfunction is an emerging hypothesis explaining affective disorders and it was shown that loss of synaptic homeostasis in specific networks also contributes to chronic pain (Torres et al., 2017). In addition, an imbalance at the pre-synaptic terminal, it regulates the zipping of the vesicles and the release of neurotransmitters (Sclip et al., 2014; 2 -at the post-synaptic level, it controls the correct organization of receptors and scaffold proteins in the active zone of the post-synaptic density region (PSD) where the receptors of the neurotransmitter (Schip et al., 2014) take their place; 3 -in the soma, it governs apoptotic, necrotic and autophagic neuronal death, acting on different targets and intracellular organelles (Figure 1).

Why JNK is a pivotal key player in neurodegenerative mechanisms? Because JNK modulates 3 major actions in neurons that are highly polarized cells. Indeed, JNK plays diverse functions in different cellular compartments: 1- at the pre-synaptic terminal, it regulates the zipping of the vesicles and the release of neurotransmitters (Biggi et al., 2017); 2- at the post-synaptic level, it controls the correct organization of receptors and scaffold proteins in the active zone of the post-synaptic density region (PSD) where the receptors of the neurotransmitter (Schip et al., 2014) take their place; 3 -in the soma, it governs apoptotic, necrotic and autophagic neuronal death, acting on different targets and intracellular organelles (Figure 1).

Thus, the central focus of our research is to study the key proteins in synaptic dysfunction/dysmorphogenesis. The first synaptic changes represent a promising therapeutic strategy and a crucial temporal window for neuroprotective treatment. In fact, synapses dynamically change and adapt to stress stimuli passing through an initial reversible phase, during which synaptic function is impaired but can be rescued, to a second phase, if stress persists, where synaptic injury becomes irreversible, and progresses to synaptic death/lack, and eventually neuronal death.

Here we reported our studies on neurodevelopmental (Angelman and Rett syndromes) as well as neurodegenerative diseases (Alzheimer and Tauopathy) (Schip et al., 2014; Buccarello et al., 2018; Musi et al., 2020, 2021). Previous evidence from our work showed that JNK signaling pathway is involved in stress response as well as in other different physiological functions, and is a common actor in the synaptophagy of all these brain diseases.

Figure 1 Brain disorders are characterized by a shared feature called synaptic dysfunction (A) and JNKs localizations in neurons (B).
formation of the synaptic vesicles and modulating the zipping and release of the vesicles. On the other hand, in the post-synaptic terminal JNK interacts and phosphorylates the most abundant secretory traffic players (-43kDa) in the PSD region and also Shank3, these scaffolds regulate the N-methyl-D-aspartic acid and aminophosphonic acid receptors in the PSD region (Kunde et al., 2012). Furthermore, the same JNK3 interacts with transcriptional factors, regulating c-jun, activating transcription factors 2, and c-fos in the nucleus, as well as with mitochondrial (among the most important is JNK1 and Golgi (i.e., scaffolding and secretory trafficking) machinery. JNK has many somatic targets like caspases, Neurofilament H, SQSTM1 (p62), and many others. In schizophrenia, JNK3 also implicated in axonal transport (de Los Reyes Corrales et al., 2021), another important function in neurons. All these physiological mechanisms are vital in a healthy brain but can be compromised leading to brain disorders.

In this perspective, we examined JNK activation in Rett and Angelman Syndromes and in AD and tested whether its specific inhibition would provide neuronal protection. We selected two developmental and a neurodegenerative disease (AD), as we are searching for basic cellular and intracellular mechanisms and common key players that govern synaptic dysfunction.

More in detail, in the context of the Rett Syndrome, a neurodevelopmental disorder, characterized by pseudo-normal development, in which patients usually achieve normal neurodevelopmental milestones but start to regress between 8 and 36 months of age with loss of language and fine motor coordination, locomotive impairment and hand stereotypes, previous work of our group investigated the synaptic dysfunction in two different mouse models (Musi et al., 2021). We identified JNK as an important actor downstream of methyl CpG binding protein 2 (MECP2), the gene mutated in the MECP2-mutated iPSCs and mouse models, which causes molecular disorganization of the PSD region in the post-synaptic element. Its disorganization has been related to locomotor and cognitive impairments that are the hallmarks of this neurodevelopmental disease. We demonstrated that the specific inhibition of JNK by D-JNK1, a cell-permeable peptide, strongly ameliorates the symptoms and the disorganization of the PSD region in both mice models. To understand the molecular mechanisms involved in human disease, we used a model of patient induced pluripotent stem cells (iPSCs) from fibroblasts of Rett patients. We demonstrated JNK activation also in human neurons, differentiated from MECP2-mutated iPSCs, compared to the isogenic control excitatory iPSCs-neurons (Penzes et al., 2020). The JNK signal was activated in the MECP2-mutated iPSCs and not in control iPSCs, and D-JNK1 blocked the MECP2mut-induced neuronal death (Musi et al., 2021). Importantly, this is the first proof of concept that JNK is a key player in Human Rett syndrome.

In Angelman Syndrome, another genetic neurodevelopmental disease characterized by autistic feature, mental retardation, and locomotor disorder, we analyzed the synaptic dysfunction in the UBE3A−/− mouse model, being UBE3A the gene mutated in this syndrome (Musi et al., 2020). JNK was strongly activated also in the brain of these mice, suggesting its important role also in this neurodevelopmental disorder. In line, these mice displayed deregulation of the markers for the excitatory spines. D-JNK1 treatment improved these behavioral abnormalities and this correlated with the stabilization of the synaptic biomarkers (Musi et al., 2020).

In neurodegenerative diseases, we mainly studied AD, since JNK phosphorylates both amyloid precursor protein (APP) and its cleaved form, and two key players of AD, accelerating the formation of β-amyloid oligomers and the deposition of neurofibrillar tangles. JNK regulates the synaptic dysfunction in vivo in both AD and in vitro Alzheimer’s disease models. Indeed, in AD animal model JNK inhibition strongly improved the synaptic impairment also ameliorating the cognitive performances (Scipì et al., 2014).

To summarize, the data obtained in these studies strongly support the notion that JNK is a central actor in the degeneration mechanisms of the synaptic dysfunction in both neurodevelopmental and neurodegenerative diseases (Figure 1).

However, JNKs are a family (Jnk1, Jnk2, and Jnk3) of mitogen-activated protein-kinases in which, Jnk1 and 2 are ubiquitous, while Jnk3 is expressed mainly in the neuronal tissue and is the most highly responsive isoform to stress in the brain pathological context, thus representing a more promising and specific target. JNK3 displays an important role in a mouse model of AD (S×FAD mice) and, in line, increased levels and activation of JNK3 have been found in the post-mortem brain of AD patients but also in their cerebrospinal fluid (Gouraud et al., 2015). These indications corroborate the idea that JNK3 is a new therapeutical target to tackle AD and other brain diseases, strongly governed by synaptic dysfunction.

In the past, kinase inhibitors were considered dangerous, with non-specific activities and therefore very difficult to apply in clinical studies. However, thanks to the discovery of specific protein-protein interactions, JNK3 specifically inhibits the field is growing. Nowadays, in the field of drug discovery, kinases have become one of the most important targets in chronic and acute diseases. In fact, currently, there are 68 FDA-approved drugs targeting different protein kinases, six of which were approved over the last year.

It is of note that the specific JNK3 inhibition may be used to modulate synaptic changes and prevent synaptic dysfunction in many different brain diseases, as shown by our research and other group’s work. As brain damage, underlying synaptic dysfunction still represents the highest burden for society and we strongly believe that it is crucial to develop a strategy to protect synapses. We are now testing JNK3 inhibitors in new neurodegenerative models focusing on neuroinflammation to understand the potentiality of this specific inhibition to tackle central nervous system dysfunctions.

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