Astrocyte syncytium: a functional reticular system in the brain

In contrary to the discrete neuronal circuits, astrocytes share their cytoplasm through gap junctional coupling into a syncytium. This distinctive difference in neuronal and astrocytic anatomy recalls the historical debate between Camillo Golgi and Ramón y Cajal on the wiring principle of the nervous system over a century ago (Cimino, 1999). At that neuron-centric era, the Cajal advocated the cell theory that considered the nervous system to be made up of discrete individual cells. Golgi on the other hand favored the reticular theory that viewed the nervous system as a singular continuing network system. Nevertheless, regardless of the difference in their viewpoints, both theories were intended to postulate an anatomical blueprint of neural circuitry, with no attention given to neuroglia. It was until the 1950s, the distinct membrane borders between synapses were revealed by electron microscopy, which means that neurons are indeed discrete individual cells. This declared Ramón y Cajal’s cell theory the victor in the debate, while Golgi’s reticular theory was disregarded. However, the existence of a ‘low-resistance pathway’ between neuroglia was soon identified in the optic nerve in the 1960s (Kuffler et al., 1966), which we now know to be gap junction coupling that connects the cytoplasm of astrocytes into a syncytial network. Ever since, decades of studies clearly demonstrated that syncytial coupling into the network is a most prominent feature of astrocytes and these glial networks are intimately interwoven with the neuronal circuits running across the entire central nervous system (Giaume et al., 2010). Surprisingly, the question of whether a reticular system exists in parallel with the neuronal circuits has received a little research attention to this day. Interestingly, the wiring pattern of astrocyte syncytium is reminiscent of the reticular theory postulated by Golgi. In this perspective, the “revived” use of “reticular theory” is solely dedicated to the brain reticular system established by astrocytes.

Astrocytes vary substantially in cytoarchitecture and spatial organization: Heterogeneity of astrocytes was initially recognized by cellular morphology, which led to the classification of protoplasmic astrocytes in grey matter and fibrous astrocytes in white matter. Now this notion continuously expands to astrocytes’ embryonic origins, gene expression, and physiological functions (Khakh and Sofroniew, 2015). Aided by the latest tissue clearing technology, a novel scale survey favors the notion that syncytial isopotentiality is a general feature among protoplasmic astrocytes in different cortical regions, in the barrel fields of layer IV somatosensory cortex, and velate astrocytes in the cerebellum. Interestingly, syncytial isopotentiality exists in the syncytium established by Bergmann glia. Although fibrous astrocytes are morphologically distinct from protoplasmic astrocytes and reside in an environment devoid of synapses, syncytial isopotentiality also appears in corpus callosum white matter. Syncytial isopotentiality has also been revealed in grey matter spinal cord astrocytes (Huang et al., 2018). Our results from a large-scale survey favor the notion that syncytial isopotentiality is a general feature of astrocyte networks.

Astrocyte syncytium: a reticular system safeguarding neuronal circuits: A general role of astrocytes is to safeguard neuronal circuits. For example, a K+ channel-mediated homeostatic mechanism, the K+ spatial buffering hypothesis, was formulated over 50 years ago by Kuffler and associates. Should this mechanism be truly operational, one of the speculated requirement is that astrocytes need to be strongly coupled into an isopotential network. Now this crucial mechanism, syncytial isopotentiality, has finally been unearthed. As a demonstrated case, this network mechanism facilitates K uptake with a far greater efficacy. Syncytial isopotentiality should also provide a high efficiency for spatial redistribution of K+ ions inside an astrocyte syncytium and with the extracellular space surrounding the very same astrocyte network (Ma et al., 2016).

How does disease conditions affect syncytial isopotentiality? Guided by the discovery of syncytial isopotentiality and the new method developed in our study, a major stroke-induced mechanism underlying astrocyte membrane depolarization was identified. This mechanism was determined to be caused by an energy failure-induced dissipation of K+ content in the entire astrocyte network (Du et al., 2018). In various kind of neurological injuries and diseases, it has been increasingly recognized that astrocytes become reactive (Sofroniew, 2014). Changes in astrocyte morphology, gene expression, and the
organization patterning could altogether occur under disease conditions. This brings into question of how such multifaceted changes could alter the strength of network coupling and function. This is a new research area that is wide open for the future. It remains completely unknown the extent to which an impaired syncytial isopotentiality contributes either as a cause or an intermediate step in neurological diseases. For instance, the syncytial architecture as a whole can be altered in certain disease conditions, such as epilepsy. Also, the Cx43 expression is known to be altered in disease conditions (Giaume et al., 2010). Therefore, a potential loss or gain of syncytial network connectivity may both occur in a context-dependent manner (Sofroniew, 2014). However, what happens to the astrocyte network communication remains to be determined. Now the availability of a new and powerful methodology should facilitate the future establishment of a causal relationship between an impaired syncytial isopotentiality and a specific type of neurological disorder.

**Future research perspectives:** Evidence from others and our studies indicate that establishment of syncytial coupling is a ubiquitous feature of astrocytes across the brain. Now, syncytial isopotentiality has been revealed as an operational mechanism of the astrocyte network and a methodology has also been established for further exploration of this mysterious reticular system in the brain. Nowadays, the “connectome” is used to refer to the structural and functional mapping of neural connectivity in the brain. Although astrocytes are wired together into a syncytium, much of the current effort is concentrated on creating wiring diagrams of neurons in different brain areas. It is time to put equal amounts of effort to establish the architectural wiring principle and function of this reticular system.

It should also be noted that the astrocyte syncytium is an electrically low-resistance reticular system. Although a better understanding of this “low-resistance pathway” has been gained at the functional level (Ma et al., 2016), the ultrastructural underpinning and biophysical rationale remain to be established. The low-resistance reticular system also means a low membrane resistance that is mostly caused by a high permeability to K+ ions (Du et al., 2015).

This front, what remains elusive is the identity of the entire repertoire of the channels that create the astrocyte membrane to be so “leaky”. Although such a low-resistance system was first brought to light by Stephen Kuffler over 50 years (Kuffler et al., 1966), the role of this reticular system in normal brain function and the diseased brain continues to be a mystery. Interestingly, newborn astrocytes are solitary individual cells (Zhong et al., 2016). Curiously, how newly generated astrocytes converge into a shared astrocyte network and when a functional astrocyte network emerges in the developing brain is basic neurobiological questions to be determined. Additionally, a functional readout for defining an astrocyte syncytium that is developmentally mature is yet to be established. This functional readout would be highly useful to determine whether a delay or reversal of this developmental process could be causative to a disease under investigation.

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