

Passing the Wave - How Does the Action Potential Pulse Pass Between Axons and Dendrites?

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Received: December 04, 2024; Published: December 24, 2024

Abstract

The action potential is an ensemble of three inseparable concurrent states, one of which is a soliton known as the action potential pulse (APPulse). Here we consider how the pulse may pass from one neuron to another. On the basis of the evidence we present here our working hypothesis is that the APPulse is passed mechanically from presynaptic terminals to postsynaptic dendrites via surrounding astrocytes. These cells are strongly connected to both pre- and postsynaptic neurons by synaptic adhesion molecules: cadherins and nectins.

Keywords: Action Potential; Action Potential Pulse; Hodgkin and Huxley; Computational Action Potential; Puncta Adherentia Junctions

Abbreviations

AP: Action Potential; APPulse: Action Potential Pulse; HH: Hodgkin and Huxley; CAP: Computational Action Potential; PAJs: Puncta Adherentia Junctions

Introduction

During the last few years much evidence has accrued to show that the action potential as originally described by Hodgkin and Huxley [1] (HH) is accompanied by a synchronized, coupled soliton pressure pulse in the membrane known as the action the action potential pulse (APPulse) [2-6]. Furthermore, action potentials are an ensemble three separate states (Figure 1): the HH physiological action potential (AP); a non-binary computational element, the computational action potential pulse (APPulse). The speed of action potential conduction is defined by the exact properties of the membrane through which the pulse travels using the concomitant entropy (released energy) from the physiological action potential. The conduction velocity, and therefore the latency of the AP, is therefore fixed to the properties of the membrane. Collisions between converging APs cancel according to the properties described by HH and the pulse structure. This adaptation and combination of both the HH and Heimburg\Tasaki [11,12] theories is essential to explain the higher and previously unconsidered functioning of neurons such as computation, especially within a brain neural network.

From a physiological point of view, it is clear that synaptic transmission allows the HH action potential to pass from one neuron to another, but the question is, how does the APPulse pass from one cell to the next or is it generated *de novo* by each cell as it passes through a chain of neurons in the central nervous system. We think the latter suggestion to be unlikely as it would introduce delays in passing

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Figure 1: The nerve impulse is an ensemble of three inseparable, concurrent states of the action potential. What an observer will perceive depends on their investigational perspective. The physiological action potential is the orthodox action potential described in detail by Hodgkin and Huxley [1]. The action potential pulse is the mechanical pressure wave for which substantial evidence has been presented elsewhere [2,3] and the computational action potential, first described by Johnson and Winlow [6]. From Bioelectricity, 3,161-170, with permission.

information from one cell to another. Another solution would be a physical connection between pre- and postsynaptic neurons and the most likely candidates to provide this are, in our opinion, the astrocytes found at central synapses in the nervous system.

Synapses

Synapses function at a slower rate than action potentials, with latencies typically in milliseconds or up to100ms where inhibitory synapses are involved [13]. However, they remain stable over the computational limits of the APPulse, i.e. in microseconds. Synapses are inherently plastic, varying latencies with time and activity, making any form of computation error prone if timing is of the same order and there is no feasible opportunity to redact error; in neural networks, quantal frequency synchronisation permits computation within the limits of plasticity and redacts error: this must take place during timing when plasticity is fixed in microseconds or less.

The probable role of astrocytes in physically connecting pre and postsynaptic neurons

Some glial cells function primarily as a scaffold for the physical support for neurons. Astrocytes are the most abundant types of neuroglia and neuroglia in general have been described as "living nerve glue" by Kettenmann and Verkhratsky [14]. While neurons have been extensively studied over many decades, the glial cells less so, both types of cells co-evolved [14,15]. Astrocytes have been shown to form tripartite synapses [16] throughout the CNS [15] and are known to interact with neurons by releasing neurotransmitters [16] often modulating neuronal intracellular calcium release and thus transmitter release from presynaptic terminals [17].

Here we consider the astrocytes found at central synapses where they form specific connections between presynaptic axon terminals and postsynaptic dendrites via synaptic adhesion molecules (SAMS), thus effectively gluing them together and preventing stress damage. Thus, they may also have a previously undescribed, additional role to play in the passage of the APPulse at the tripartite synapses, where they physically connect the pre- and post-synaptic neurons (Figure 2) via cadherins and nectins. Any physical movement in the presynaptic cell will be physically transmitted to the postsynaptic cell via the astrocyte. If this is the case, it would allow physical transmission of the APPulse from the presynaptic nerve terminal to postsynaptic dendrites.

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Cadherins and nectins

Cadherins were first described by Takeichi [18] and are transmembrane proteins mediating cell-cell adhesion. At synapses they are termed synaptic adhesion molecules and contact pre- and post-synaptic cells at *puncta adherentia* junctions (PAJs), thus providing an extracellular skeleton holding the cells tightly together and preventing separation due to physical stress (Figure 2). Nectins also have similar functions at PAJs [19].



Figure 2: A simplified diagram of the mechanical connections between astrocytes and cell adhesion molecules (cadherins) at a central tripartite synapse. The connections are made at PAJs. The presynaptic cell and the astrocytes release neurotransmitters, but this not shown here - for further detail see Togashi., et al. [15] and Araque., et al [16].

Discussion

Synapses are an essential part of the process of perception and motor control, in that they have finite control of latencies over the 0.3 ms to 100 ms slower than parallel processing which in the retina occurs at 10⁻⁶s [7]. Synapses separate cellularly diverse structures where each neuron, its shape morphology and position histologically has a functional basis. Synapses provide changing latency permitting memory loops to align and form abstract patterns. For example, object representation in the retina and LGN from large object forms a pattern in the visual cortex L1 and L2 changing to detail L5 which are assimilated into the pyramidal neurons which synchronise from L2 to L5 and output accurately final objects in L6 where synchronisation occurs with L1 from parallel output from the LGN. This redistribution of circulating synchronised memory is essential in learning when considering the cerebral cortex and is predetermined by the plasticity inherent in the synapses increases the span of latency timings and changes the neural networks by raising the concentrations and effectiveness of neurotransmitters. The close-joints of cadherins etc and protein structure may also facilitate a direct transmission of impulses by mechanical impulses described by the APPulse, reducing the latencies, similar to gap junctions, and increasing uniform stable transmission. This concurs with our theory that synapses act as changers of latency in an inherently plastic environment. Thus, a brain neural network is by no means fixed in functional form but adjusts according to activity.

We have shown that propagation must occur through mechanical means and the only available candidate is the well-documented mechanical pulse that accompanies the HH AP. This pulse is ubiquitous whether the neuron is spiking or is spikeless, as have been described in the eye and elsewhere [7]. That the pulse is propagated mechanically is evident from the timing constraints of the demonstrated

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computation in the eye [7] and must take place mathematically within 10⁻⁶s: without this timing synchronisation and thus computation, as demonstrated [7] cannot be reliable and accurate. This is the predicted precision of timing between the start of propagation from the beginning of the soliton to the opening of the ionic gates.

Conclusion

The action potential pulse may be mechanically transmitted between neurons by astrocytes connected to pre- and postsynaptic neurons by cadherins and nectins.

Conflict of Interest

The authors received no external support for this work and have no conflicts of interest.

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