Neurons have high densities of voltage-gated Na\(^+\) channels that are restricted to axon initial segments and nodes of Ranvier, where they are responsible for initiating and propagating action potentials. New findings (Bréchet et al., 2008) reveal that phosphorylation of several key serine residues by the protein kinase CK2 regulates Na\(^+\) channel interactions with ankyrin G. The presence of CK2 at the axon initial segment and nodes of Ranvier provides a mechanism to regulate the specific accumulation and retention of Na\(^+\) channels within these important domains.

Neurons process and encode information through synaptic activity that results in the generation and propagation of action potentials (APs). Voltage-gated Na\(^+\) (Na\(^+\)) and K\(^+\) (K\(^+\)) channels, highly enriched at the axon initial segment (AIS) and nodes of Ranvier (Fig. 1 A), are central to this activity. Electrophysiological measurements suggest that AIS Na\(^+\), channel density is \(\sim 50\) times that of proximal dendrites (Kole et al., 2008), and in myelinated axons, nodal Na\(^+\) channel densities are at least 25 times those of the internode (Shragger, 1987). How do neurons restrict Na\(^+\) channels to the AIS and nodes? Experiments indicate that the cytoskeletal scaffold ankyrin G (ankG) is responsible for clustering Na\(^+\) and K\(^+\) channels. For example, loss of ankG by RNA interference or in mutant mice blocks AIS and nodal clustering of ion channels (Zhou et al., 1998; Dzhavashvili et al., 2007; Hedstrom et al., 2007), and efforts to identify AIS localization determinants in Na\(^+\) channels revealed a highly conserved ankG-binding AIS-targeting motif in the II-III linker domain (Garvid et al., 2003; Lemaitre et al., 2003). Intriguingly, KCNQ2/3 K\(^+\) channels, also enriched at the AIS, appear to have independently evolved an AIS-targeting motif nearly identical in sequence to that found in Na\(^+\) channels (Pan et al., 2006).

Besides ankG, axons also have high levels of ankyrin B (ankB). These similar cytoskeletal scaffolds occupy complementary regions of the axon (Fig. 1 B), such that in myelinated axons, ankB is restricted to paranodes, whereas ankG is only found at nodes (Fig. 1 A, inset; Ogawa et al., 2006); in hippocampal neurons, ankG is found only at the AIS, whereas ankB is found throughout the distal axon (Boiko et al., 2007). Because both ankB and ankG share a highly conserved membrane-binding domain (MBD) that interacts with the AIS-targeting motif, it has been difficult to understand how Na\(^+\) channels are restricted only to regions with ankG. In the current issue, Bréchet et al. (see p. 1101) help to resolve this conundrum by demonstrating that spatially regulated phosphorylation of several key serine residues in the AIS-targeting motif strongly enhances its affinity for the MBD.

Phosphorylation (and dephosphorylation) of AIS and nodal proteins has been shown previously to play important roles in regulating AIS assembly. For example, phosphorylated p65/RelA and IκBα, an inhibitor of the nuclear factor κB (NFκB) transcription factor, are enriched at the AIS and nodes of Ranvier (Schulz et al., 2006), and inhibition of IκBα phosphorylation disrupts AIS formation (Sanchez-Ponce et al., 2008). In contrast, rather than promote binding, phosphorylation of a tyrosine residue conserved in L1 family cell adhesion molecules enriched at the AIS (e.g., neurofascin-186 and NrCAM) blocks their interaction with ankG (Garver et al., 1997). It may be that phosphorylation is a common mechanism for regulating interactions among AIS proteins.

To investigate this possibility and to further define the mechanism of Na\(^+\), channel clustering and interaction with ankG, Bréchet et al. (2008) performed a structure function analysis of the AIS-targeting motif using a K\(^+\)/Na\(^+\) chimera that included only the AIS-targeting motif of Na\(^+\) 1.2. They found that by combining a glutamate mutation (Na\(^+\) 1.2 E1111A) with mutation of single or multiple serines in the AIS-targeting motif (Na\(^+\) 1.2 S1112A, S11123-24A, or S11126A; Fig. 1 C), AIS recruitment of the mutant chimera was completely abolished. Subsequent sequence analysis suggested that these serines were potential phosphorylation sites for the protein kinase CK2, and in vitro phosphorylation assays confirmed this prediction. Using surface plasmon resonance, they then showed that CK2 modulated the association of the AIS-targeting motif with the ankyrin MBD. In fact, CK2-mediated phosphorylation increased the affinity of the AIS-targeting motif for the ankyrin MBD \(\sim 1,000\)-fold! The relevance of the phosphorylation for Na\(^+\), channel clustering was further demonstrated by showing that the K\(^+\)/Na\(^+\) 1.2

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It is tempting to speculate that the enrichment of Na\textsubscript{v} channels only in ankG-rich regions might be further ensured by an ankB-interacting phosphatase. This would provide an additional layer of regulation, ensuring that Na\textsubscript{v} channels are spatially restricted and only interact with ankG. Because Na\textsubscript{v} channels can also bind to ankB-MBD, inappropriate phosphorylation of the serines within the AIS-targeting motif could result in ectopic clustering of channels along the axon, as well as hyperexcitability. Alternatively, failure to appropriately phosphorylate channels (or increased activity of phosphatases) could lead to a neuron in a hypoexcitable state and conduction block. It will be interesting to determine if diseases or injuries that lead to altered cellular excitability might not result from inappropriate Na\textsubscript{v} channel phosphorylation or dephosphorylation (Fig. 1 C). Although ankG and ankB are very similar in their MBDs, ankB contains a unique insertion that facilitates intramolecular binding between ankB’s N-terminal membrane-binding and C-terminal domains.

Protein kinase CK2’s role in regulating Na\textsubscript{v} channel clustering was further confirmed by demonstrating that CK2 is highly enriched at the AIS of hippocampal neurons both in vitro and in vivo, and at central nervous system and peripheral nervous system nodes, where it colocalizes with Na\textsubscript{v} channels and ankG. Thus, CK2 is spatially restricted to sites where it can promote the binding of Na\textsubscript{v} channels to ankG. Inhibition of CK2 in cultured hippocampal neurons caused a decrease in the density of Na\textsubscript{v} channels at the AIS. Surprisingly, CK2 inhibition also caused a reduction in the amount of ankG at the AIS, which suggests either that CK2 directly regulates ankG stability (possibly through βIV spectrin) or that Na\textsubscript{v} channels can also influence the stability of ankG. This latter possibility is consistent with the observation that silencing of Na\textsubscript{v} channel expression in motor neurons by RNA interference reduces AIS ankG clustering (Xu and Shrager, 2005).

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Rather than dephosphorylation of the Na+ channel, ankG specificity may be a consequence of ankB’s MBD being occupied by its own C-terminal domain (Abdi et al., 2006).

Although the results of Bréchet et al. (2008) provide important insights into the mechanisms of AIS assembly, as any good paper should, they also raise additional questions. For example, how is CK2 restricted to the AIS? Does its localization depend on ankG? Because KCNQ2/3 K+ channels share a common AIS-targeting motif, does CK2 also regulate the localization of these K+ channels? Further, it is also important to note that mutation of the serines alone (whether single or multiple) did not affect the localization of K+2.1-Na+1.2 chimeras unless these mutations were combined with the glutamate (Na+1.2 E111A) mutation. The significance of E111A remains obscure but suggests that phosphorylation is not the only mechanism regulating Na+ channel clustering. It is also intriguing to note that additional levels of subcellular specialization exist among Na+ channels at the AIS. For example, although ankG extends throughout the AIS in retinal ganglion cells, Na+1.1 occupies a microdomain in the proximal AIS (close to the cell body), whereas Na+1.6 channels occupy the complementary distal AIS (Van Wart et al., 2007). The reasons for this additional degree of specialization are unknown, and it will be interesting to determine if phosphorylation contributes to formation of these microdomains.

Finally, perhaps the most intriguing implication of this work is the potential for dynamic regulation of Na+ channel density. Because action potential initiation depends on the density of Na+, (and K+) channels, modulation of AIS channel numbers by regulating CK2 activity or through competing phosphatases (Fig. 1 C) could be an important way to dynamically regulate the biophysical properties of the spike-generating machinery. If true, besides synaptic plasticity, AIS plasticity may be another way in which neurons can strengthen or modify neural circuits.