Signaling complexes of voltage-gated calcium channels

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Voltage-gated calcium channels are key mediators of depolarization induced calcium entry into electrically excitable cells. There is increasing evidence that voltage-gated calcium channels, like many other types of ionic channels, do not operate in isolation, but instead form complexes with signaling molecules, G protein coupled receptors, and other types of ion channels. Furthermore, there appears to be bidirectional signaling within these protein complexes, thus allowing not only for efficient translation of calcium signals into cellular responses, but also for tight control of calcium entry per se. In this review, we will focus predominantly on signaling complexes between G protein-coupled receptors and high voltage activated calcium channels, and on complexes of voltage-gated calcium channels and members of the potassium channel superfamily.

Introduction

Calcium ions serve as important mediators of cell signaling in both excitable and non-excitable tissues. Elevation in intracellular calcium levels triggers physiological responses that include cardiac muscle contraction, hormone secretion, neurotransmitter release, activation of calcium-dependent enzymes and calcium dependent gene transcription.1–5 In electrically excitable cells such as neurons and muscle cells, action potential evoked calcium entry is primarily due to the activation of both low and high voltage-gated calcium channels.6 Low voltage activated (LVA),7 and high voltage activated (HVA) calcium channels can be distinguished based on their threshold of activation, with LVA channels activating close to resting membrane potentials.8 In contrast, the family of HVA calcium channels can be distinguished from one another by their functional properties and pharmacological profiles.9–13 They also support distinct physiological functions. For example, T-type calcium channels are involved in regulating cellular excitability,14–16 N and P/Q-type channels mediate fast evoked neurotransmitter release,1 and L-type calcium channels mediate functions such as excitation coupling in the heart and muscle, insulin secretion and calcium dependent gene transcription.17–21 Along these lines, different calcium channels isoforms are targeted by clinically active drugs such as analgesics, general anesthetics, antiepileptics and cardioactive drugs,22–24 and mutations in various calcium channel isoforms have been associated with conditions such as familial migraine, deafness, epilepsy, cardiac arrhythmias and ataxia (reviewed in refs. 25 and 26).

Common to all types of voltage-gated calcium channels is a pore forming Ca_\(\alpha_1\) subunit that contains four homologous membrane domains that are flanked by cytoplasmic N- and C-termini and connected by cytoplasmic linker regions.2 While T-type calcium channels are thought to contain only a Ca_\(\alpha_1\) subunit, members of the HVA calcium channel family also contain a cytoplasmic Ca_\(\beta\) subunit, as well as a larger extracellular membrane anchored Ca_\(\alpha_2\delta\) subunit, and both of these subunits are represented by four different genes (reviewed in refs. 27 and 28). This coassembly with ancillary subunits is necessary in order to promote export of the channels from the endoplasmic reticulum,29 and also results in alteration of the functional properties of the channels.30 In addition, all of the known HVA channels co-assemble with calmodulin, a calcium binding protein that bestows calcium feedback regulation onto the channels in the form of calcium-dependent inactivation and facilitation.31–33 The incorporation of a calcium signaling protein into the channel complex is indicative of the notion that calcium channels do not operate in isolation, but tend to form larger signaling complexes that are not only designed to enhance coupling efficiency between calcium entry and downstream signaling events, but also provide for a means of regulating calcium channel activity per se.

There are numerous examples of cytoplasmic protein interactions with different types of voltage-gated calcium channels. For example, N and P/Q-type channels tightly associate with proteins of the synaptic vesicle release machinery, such as syntaxin 1 and SNAP-25, thus effectively coupling calcium entry to fast vessel release, but also to confer feedback regulation onto channel activity.34–36 Small GTPases such as Rem, Gem and Rem2 associate with HVA channels to regulate channel surface expression (reviewed in ref. 39). A-Kinase anchoring proteins, protein kinase A, and phosphatase 2A associate with certain L-type channel isoforms to tightly control the regulation of channel activity by phosphorylation.40–43 However, it has only emerged recently that various types of calcium channels interact not only with cytoplasmic signaling molecules, but also with other membrane proteins.
such as receptors and ion channels to form physical signaling complexes that either allow for more efficient regulation of calcium channel activity or facilitate the specificity of calcium signaling. Here we will focus on two such examples: We will first discuss how protein interactions with G protein coupled receptors enhance calcium channel modulation and trafficking. We will then review how interactions between calcium channels and potassium channels bestow calcium regulation onto potassium channel function and thus modify neuronal output.

G Protein Coupled Receptor: Calcium Channel Complexes

Most types of voltage-gated calcium channels are modulated following the activation of various types of G protein coupled receptors (GPCRs) (reviewed in refs. 45 and 46). Upon receptor activation, GDP bound constitutively to the Goα subunit is exchanged for GTP, resulting in a conformational change that either leads to a breakup of the Goβγ trimer, or at least in a conformational change within the trimer that results in the generation of two separate signaling entities—activated Goα and Gβγ dimers. Both Goα and Gβγ act on various effector molecules. For example, Goα may stimulate the activity of adenylly cyclase, leading to the production of cyclic AMP and the activation of protein kinase A. In the context of voltage-gated calcium channels, a classic example of such a soluble second messenger pathway is the regulation of L-type calcium channels by β-adrenergic receptors (βARs), which results in protein kinase A dependent phosphorylation of the channels in cardiac cells and a massive upregulation in channel activity that ultimately increases heart rate. While Gβγ subunits act to regulate various cell signaling pathways, they also can directly modulate certain types of ion channels via a membrane-delimited pathway. This was first described almost three decades ago for N-type calcium channels in sensory neurons. Activation of noradrenergic receptors was found to produce a rapid depression of N-type channel activity that was later found to be strongly voltage dependent, and to be mediated by direct binding of Gβγ to the N-type calcium channel α1 subunit. This binding interaction and the associated functional regulation appears to be destabilized by protein kinase C (PKC) dependent phosphorylation of the channel and there is evidence that PKC is in fact associated with the channel via an enigma homolog adaptor protein, suggesting the existence of a channel-kinase complex. Given that the Gβγ subunit is large and anchored to the inner leaflet of the plasma membrane and thus unlikely to diffuse quickly across large distances, one might expect that channels and receptors would have to be localized in close proximity to each other to account for the relatively rapid onset of G protein inhibition. This then raises the possibility that G protein coupled receptors and voltage-gated calcium channels may be located in close proximity either in the same lipid rafts, of perhaps even be physically associated.

L-Type Channels

The first description of a calcium channel-GPCR complex involved Ca1.2 L-type calcium channels and βARs. There are three known βAR subtypes expressed in the mammalian heart (1AR, β2AR, β3AR), with the 1AR and β2AR subtypes able to activate PKA and stimulate calcium entry via L-type calcium channels into cardiac myocytes. It has been shown that β2ARs form macromolecular signaling complexes with Ca1.2 calcium channels. These complexes also contain other elements of the signaling machinery, such as G proteins and caveolin. The functional significance of these assemblies has not yet been demonstrated in cardiac myocytes, nor have the molecular determinants that underlie the interactions between L-type calcium channels and receptors been identified. The investigation into coupling between these receptors and the channels has been hampered by the fact that it has proven to be a challenge to reconstitute βAR signaling to L-type channels in heterologous systems—a problem that has only been solved recently. Nonetheless, the existence of receptor-Ca1.2 channel complexes suggests that they may help optimize the coupling between the receptor and the channels.

P/Q-Type Channels

Like N-type calcium channels, P/Q-type channels can be regulated by a membrane delimited Gβγ mediated pathway in response to activation of a number of different receptor subtypes. These channels are expressed at high levels in cerebellar Purkinje neurons where they are regulated by activation of metabotropic glutamate receptors. Coexpression of these receptors with Ca2.1 (P/Q-type) channels in heterologous expression systems was shown to reduce current amplitude in an agonist independent manner, suggesting a direct regulation of channel activity by the receptor itself. Furthermore, it was shown that receptors and channels colocalized to the same dendritic compartments in Purkinje neurons, and that they could be co-immunoprecipitated both from transfected cells and native tissue. This close spatial arrangement may serve to optimize receptor signaling to the P/Q-type channels and thus to fine tune synaptic transmission. Furthermore, the agonist independent modulation of the channel by these receptors could potentially provide a mechanism by which P/Q-type channel function may be regulated by changes in receptor density.

N-Type Channels

The interaction between N-type channels and GPCRs has been extensively investigated. Like P/Q-type channels, N-type channels are heavily clustered at synaptic sites where they contribute to the release of neurotransmitters. Hence, any inhibition of N-type channels has the potential to inhibit neurotransmission. The significance of such regulation is underscored by the clinical use of morphine, a potent activator of μ-opioid receptors. Activation of these receptors results in the inhibition of N-type calcium channels at dorsal horn synapses, thus preventing the transmission of pain signals from primary afferent fibers.

Numerous other types of GPCRs have been found to inhibit N-type channels via a Gβγ mediated, voltage dependent pathway, and in some cases a voltage-independent inhibition has been observed (but will not be considered further here). One
At the same time, the coexpression of the N-type channel with ORL1 resulted in increased cell surface density of the channels. It is possible that the ORL1 receptor, when bound to the channel, occludes an ER retention motif, thus facilitating ER export and surface expression. This would fit with the observation that the ORL1 receptor binds to the channel’s C-terminal region—a locus that has been implicated in the ER retention of the channel.

Unlike the ORL1 receptor, μ-opioid receptors do not appear to regulate N-type channel trafficking. Intriguingly, when coexpressed with the ORL1 receptor, the μ-opioid receptor subtype becomes capable of internalizing N-type channels. This is presumably due to the fact that opioid receptors can form heterodimers with ORL1, thus giving rise to a trimeric signaling complex with unique trafficking properties (Fig. 1, inset).

Two other types of GPCRs have since been shown to regulate N-type channel trafficking. Both D1 and D2 dopamine receptors interact with the N-type calcium channel α1 subunit, albeit at distinct sites. Nonetheless, in both cases, receptor coexpression increases the cell surface density of the channels, whereas receptor activation mediates N-type channel internalization. In the case of D1 receptors, it was shown that the receptor targets the channels to dendritic sites in prefrontal cortex neurons, thus

![Figure 1. Physical interactions between certain types of GPCRs and voltage-gated calcium channels bring the receptors close to the channels, so that receptor/G protein activation results in effective coupling to the channels. In addition, N-type calcium channel interactions with either ORL1 receptors or dopamine receptors have been shown to support agonist dependent internalization of channel receptor complexes, and to facilitate insertion of complexes into the membrane after synthesis. Inset: ORL1 receptors mediate N-type channel internalization upon prolonged treatment with the agonist nociceptin. In contrast, the μ-opioid receptor agonist DAMGO results in internalization of these receptors, but not of the channels unless ORL1 receptors are also present (i.e., the ORL1 receptor acts as a bridge between the N-type channel and the μ-opioid receptor).](image-url)
effectively defining the subcellular localization of the channels.\textsuperscript{76} Hence, a GPCR is able to divert the N-type calcium channels from their normal presynaptic locus to mediate calcium signaling in dendrites.

At this point, it is not clear if other types of GPCRS are able to associate with N-type calcium channels to regulate their function and trafficking. However, efficient channel receptor coupling may necessitate a close spatial proximity between receptors and channels that may be best achieved through either direct physical interactions, or linkage via an intermediate protein.

In the above sections, we have discussed how the formation of calcium channel-GPCR complexes optimizes receptor-mediated control over calcium channel activity. Below, we will illustrate how the formation of protein complexes with potassium channels allows for better calcium channel mediated control of potassium channel function, and thus neuronal firing.

**Calcium-Potassium Channel Complexes**

Calcium channels have long been known to associate with calcium-dependent potassium (K\textsubscript{Ca}) channels to control membrane excitability through activation of potassium efflux. K\textsubscript{Ca} channels expressed in central neurons correspond either to big conductance (BK, mslo) or small conductance (SK1-3) channels.\textsuperscript{78,79} Past work has established the ubiquitous occurrence of calcium-K\textsubscript{Ca} channel interactions across cells of the nervous system, but increasingly, it has become clear that coupling can be highly specific to particular channel isoforms in different cells. These interactions were typically first assessed at the level of a functional coupling between calcium channel subtypes and BK or SK potassium channels in relation to specific processes (i.e., spike discharge vs. subthreshold synaptic events) (reviewed in ref. 80). Although this interaction can reflect a close proximity between calcium and potassium channels, it may or may not reflect interactions at the molecular level within an ion channel complex. Recent work, however, has identified an association between different calcium and K\textsubscript{Ca} channels that provides for highly localized and specific coupling between calcium influx and activation of a potassium channel within a protein complex. The first coupling between a calcium channel and a voltage-gated potassium channel was also reported in the form of a Ca\textsubscript{3}-K\textsubscript{4} complex,\textsuperscript{81,82} significantly increasing the range of neuronal functions that will benefit from calcium-dependent regulation of potassium channel function.

Here we review examples of calcium-K\textsubscript{Ca} channel coupling and the advantages conferred upon regulation of cell excitability in central neurons. We consider cases where potassium channel activation reflects a functional coupling as well as the existence of an actual ion channel complex between calcium and K\textsubscript{Ca} channels, or to the K\textsubscript{4}/KChIP complex. For further information readers are referred to several excellent reviews on details of K\textsubscript{Ca} and K\textsubscript{4} channel activation.\textsuperscript{79,80,83-87}

**Control of Cell Excitability by Calcium-K\textsubscript{Ca} Coupling**

BK channels are activated through a complex and synergistic interplay between membrane voltage and the internal concentration of calcium.\textsuperscript{83} The channels are comprised of an extracellular N-terminus, seven transmembrane segments (S0–S6) and an internal C-terminus. Calcium sensitivity is mediated through a C-terminal “calcium bowl” and RCK1 domain, with response in the 1–10 \( \mu \)M range of internal calcium concentration.\textsuperscript{83,84} SK\textsubscript{Ca} subunits are encoded by four genes (SK1–4), with central neurons differentially expressing SK1–3 channels. SK channels have no intrinsic voltage-sensitivity and are thus only sensitive to changes in the concentration of internal calcium. Calcium dependence is achieved by interacting with calmodulin at a domain on the C-terminus, with the complex exhibiting greater sensitivity to internal calcium concentration than BK channels, with a half maximal activation at ~0.3 \( \mu \)M calcium.\textsuperscript{80}

The different means by which BK and SK channels achieve calcium-dependent activation with or without sensitivity to membrane voltage fluctuations enable these channels to influence widely different forms of neuronal activity. In general, BK channels are recognized for their role in repolarizing the fast falling phase of sodium spike discharge and a subsequent fast AHP of \(<10\) msec.\textsuperscript{89-92} SK channels exhibit a time constant for activation of ~5 msec but much slower deactivation that allows for the generation of a medium AHP (mAHP) that lasts in the order of a 100 msec.\textsuperscript{83,93,94} As such, SK-mediated AHPs can respond to longer time frames of cell activity than BK channels by sensing the levels of calcium accumulated during repetitive spike discharge. SK channels can thus alter firing rate gain\textsuperscript{89,94} or control the duration of calcium-dependent depolarizing or burst responses.\textsuperscript{95,96} The activation of BK or SK channels by spike discharge can further establish a baseline level of membrane excitability according to the rate of tonic background firing.\textsuperscript{97} Associating specific calcium channel subtypes with either BK or SK channels thus provides control over select aspects of neuronal activity.

The existence of toxins or drugs specific to different calcium channels and to K\textsubscript{Ca} channels has helped to identify the source of calcium influx responsible for controlling K\textsubscript{Ca}-mediated responses. The number of these studies conducted to date are too numerous to review here, but serve to indicate the existence of at least a functional coupling between different HVA or LVA calcium channels and BK or SK channels that are often specific to different cell types. Thus, N-type calcium channels were found to functionally couple to the activation of BK channels in CA1 pyramidal cells\textsuperscript{98} and SK channels in cells of deep cerebellar nuclei,\textsuperscript{97,98} thalamic reticular nucleus,\textsuperscript{95} and nucleus basalis.\textsuperscript{100} P-type calcium channels couple to the activation of BK channels\textsuperscript{101,102} or SK channel\textsuperscript{102} in cerebellar Purkinje cells. L-type calcium channels have been reported to activate BK channels in neocortical pyramidal cells\textsuperscript{103} and CA1 hippocampal pyramidal cells,\textsuperscript{98} as well as to SK channels in CA1 pyramidal cells.\textsuperscript{98} Finally, T-type calcium channels are at least functionally coupled to BK channel activation in medial vestibular neurons,\textsuperscript{104} and to activation of SK channel isoforms in dopaminergic neurons,\textsuperscript{105} thalamic reticular cells\textsuperscript{106} and nucleus basalis neurons.\textsuperscript{100}

In some cases, calcium entry through more than one calcium channel isoform can activate the same K\textsubscript{Ca} current in a given cell, as found for BK channel activation by both N- and L-type channels in neocortical pyramidal cells.\textsuperscript{103} Conversely, calcium entry
also exhibit the Ca v-BK complex. At this time no reports have surfaced on the potential for T-type calcium channels to form nanodomain interactions with either BK or SK channels.

The mechanism by which a nanodomain interaction can take place between HVA calcium and BK channels was only recently resolved. In particular, all of Ca v1.2 (L-type), Ca v2.1 (P/Q-type) and Cav2.2 (N-type) (but not R-type) calcium channels could be coimmunoprecipitated with BK channel and β subunit proteins from rat brain or from lysates of expression systems. Expression of each of the HVA calcium channels with BK channels (and β subunits) revealed that calcium-dependent activation of BK current was insensitive to internally perfused EGTA, but blocked by BAPTA, consistent with a nanodomain interaction. This interaction appeared to involve only the α-subunits of calcium and BK channels given the ability to coimmunoprecipitate these proteins in the absence of auxiliary subunits (Fig. 2A).

The physiological relevance of an HVA-K Ca interaction is several fold. By closely associating specific calcium channel subtypes to BK or SK channels, the activation of potassium current can be restricted to a given voltage response, or tailored to different activity patterns of a cell. Thus, spike frequency can also exhibit the Ca v-BK complex. At this time no reports have surfaced on the potential for T-type calcium channels to form nanodomain interactions with either BK or SK channels.

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be modulated according to the rate of spike repolarization and generation of an AHP, a process that would benefit from the nanodomain interaction between HVA calcium and BK channels. By comparison, microdomain interactions between T-type calcium and SK channels would be sufficient to control lower frequency oscillatory responses. The cell-specific interactions apparent between calcium and K channels also provide control over the voltage for potassium channel activation according to the associated calcium channel isoform. Thus, a comparison between the I-V plots for BK channels coexpressed with either L-type or P/Q-type calcium channels in Xenopus oocytes revealed distinctly different voltage ranges for potassium current activation, as conferred upon the potassium channel by the associated calcium channel. These distinctions were further shown to provide very different profiles of potassium current in response to spike-like voltage clamp commands of different half-widths. These differences would be expected to have significant influence on potassium current associated with spike discharge between different cells, and between soma and axon terminals, with consequent effects on transmitter release.

The current data thus reveal that interactions between calcium channels and either SK or BK channels can be highly specific to different cells, and tailored through the formation of protein complexes and differential calcium requirements to modify a wide range of neuronal outputs. While altered neuronal output will indirectly affect the activities of voltage-gated calcium channels (i.e., via changes in membrane potential), it remains to be determined if the physical association with these potassium channel family members may directly alter calcium channel function.

Ca-K Interactions

Most recently, the range of possible interactions between calcium and potassium channels was greatly extended by reports of a new interaction between LVA Ca 3 T-type calcium channels and K4 A-type potassium channels. This interaction is unique in representing the first identification of an association between a calcium channel and a purely voltage-gated potassium channel. T-type calcium and A-type potassium currents share the properties of fast activation and inactivation, near complete inactivation at rest, and recovery from inactivation during membrane hyperpolarizations. This extensive overlap in the voltage-dependent and kinetic properties of Ca 3 calcium and K4 potassium channels thus appears ideally suited to associating these channel types as part of a single complex. Periods of inhibition in cells expressing these channels will then increase the availability of both A-type and T-type currents, allowing for their transient activation upon return to resting potential. The consequence of this combination of inward and outward current activation was illustrated in cerebellar stellate cells, where an A-type mediated increase in first spike latency (FSL) was transformed to a decrease in FSL over the slightly shifted voltage range for recovery of T-type channels from inactivation. Further investigation of this interaction revealed that Ca 3 and K4.2 channels coimmunoprecipitate from brain lysate, with pull-down experiments indicating an association at the molecular level between K4.2 channels and Ca 3 C-termini (Fig. 2B).

The existence of this association was instrumental in resolving a long-standing issue of the potential function for a previously defined association between K4 channels and K+ Channel Interacting Proteins (KChIPs). First discovered in yeast-two hybrid screens searching for K4 associated proteins, KChIPs bind K4 channels at two sites: a hydrophobic N-terminal segment and at the T1 assembly domain loop. By way of binding, KChIPs dramatically change the voltage-dependent and kinetic properties of K4 channels. Specifically, KChIP binding promotes the trafficking of K4 channels to the plasma membrane, hastens the rate of Ia recovery from inactivation, slows the rate of Ii inactivation, and causes shifts in the voltage-dependence of activation and inactivation. Importantly, KChIPs are calcium binding proteins that belong to the neuronal calcium sensor (NCS) family. Similar to other NCS family members, KChIPs contain four EF-hand binding motifs, two of which (EF-3 and EF-4) bind calcium. Initial mutagenesis studies revealed that KChIP1 molecules that were incapable of binding calcium could still associate with K4 channels but lost the ability to modulate K4 properties, suggesting a role for calcium in K4 function. Indeed, a study investigating the effects of intracellular calcium on K4.3-KChIP2d (a minimal KChIP isoform which only contains one EF-hand) demonstrated that chelation of intracellular calcium abolished the established KChIP effects on gating. Conversely, recent structural studies of the K4.3-KChIP complex have revealed that some KChIP EF-hand mutants lose the ability to bind K4.2 channels, highlighting the important role calcium binding plays in structural maintenance of KChIP proteins.

While isolated studies had shown that neuronal Ia could be modulated by fluctuations in intracellular calcium, identifying a calcium source capable of regulating K4 properties had remained elusive. However, it was recently determined that Ca3-mediated calcium influx in cerebellar stellate cells serves to modulate the voltage-dependence of K4 inactivation (Vh) at the nanodomain level (Fig. 2B). Heterologous expression studies further showed that KChIP3 (but not KChIP1, 2 or 4) was the calcium sensor involved in the modulation of K4 Vh. Similarly, internal perfusion of specific antibodies directed at KChIP proteins in stellate cells revealed that KChIP3 was the critical link between Ca3-mediated calcium influx and the calcium-dependent modulation of K4 function. Moreover, without the actions of the Ca3-K4 complex, the voltage range for K4 channel inactivation would be shifted far enough leftward as to essentially fall outside of the physiological range of usual membrane potential fluctuations. The Ca3-K4 interaction thus appears to be instrumental to the contribution of A-type K4 potassium channels to neuronal activity. Although the number of physiological processes that could be affected by the Ca3-K4 complex remain to be fully explored, the current data set reveal that the Ca3-K4 complex is key in controlling stellate cell excitability by modifying the gain of firing frequency. Given the prominent expression of these channel types and KChIP3 proteins, it is predicted that calcium-dependent modulation of K4 availability and thereby neuronal excitability.
will be a widespread phenomenon throughout central and peripheral neurons. As in the case of calcium activated potassium channels, it remains to be seen if KCa1Ps or K_4 channels can directly alter calcium channel function/membrane expression levels.

**Concluding Remarks**

Here, we have highlighted two general examples of how cell signaling can be optimized through the formation of specific signaling complexes between membrane proteins. While receptor heterodimerization has been a well-established phenomenon, the emergence of singular complexes between receptors and ion channels, or between two different types of ion channels is a more recent concept. The notion that channels and receptors may exist as macromolecular signaling complexes makes intuitive sense, as this not only allows maximizing the efficiency of coupling and crosstalk between the individual players, but may also allow for unique signaling properties that are specific to a particular subcellular locus. Although the examples of physically signaling complexes involving different types of channels and receptors are still relatively limited, it is likely that more such examples will emerge in the near future.

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