Novel coupling is painless

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The first report that several neurotransmitters, including γ-aminobutyric acid (GABA), decreased neuronal action potential duration in dorsal root ganglia (DRG) neurons appeared more than 35 years ago (Dunlap and Fischbach, 1978). Dunlap and Fischbach (1978) realized that the effects of GABA could not occur via the ionotropic GABA receptor—at that time the only known GABA receptor—and that the target ion channel was most likely a voltage-gated calcium (Ca\(^{2+}\)) channel, rather than a Na\(_{\text{v}}\) or K\(_{\text{v}}\) channel. They correctly concluded that another type of GABA receptor must exist, which we now know is the G protein–coupled GABA type B (GABA\(_{\text{B}}\)) receptor. The modulated channel was later identified as the chick homologue of the N-type Ca\(^{2+}\) channel Ca\(_{\text{V}}\), (α\(_{\text{IV}}\)) (Cox and Dunlap, 1992), one of three members of the Ca\(_{\text{V}}\) family. GABA\(_{\text{B}}\) receptors in human and rodent sensory neurons and in various expression systems were shown subsequently to inhibit native N-current and recombinant Ca\(_{\text{V}}\) current, respectively (Raingo et al., 2007; Callaghan et al., 2008; Adams and Berecki, 2013). Inhibition primarily occurs by a voltage-dependent mechanism common to various neurotransmitters whereby G\(\beta\γ\) binds to Ca\(_{\text{V}}\) slowing channel opening, whereas positive voltage steps relieve this inhibition (Marchetti et al., 1986). The closely related P/Q-type (α\(_{\text{P/Q}}\)) channel, Ca\(_{\text{V}}\), exhibits similar modulation by GABA (Mintz and Bean, 1993). The third member of the Ca\(_{\text{V}}\) family, Ca\(_{\text{V}}\) (α\(_{\text{E}}\)), is less susceptible to direct G\(\beta\γ\) modulation than the other two family members (Sheket et al., 1997). The revelation that mice with a deletion in either Ca\(_{\text{V}}\) or in Ca\(_{\text{V}}\) exhibited reduced neuropathic pain–like behavior, indicating that these channels participate in pain sensation signaling (Saegusa et al., 2000, 2001), sparked great interest in the regulation of Ca\(_{\text{V}}\) inhibition by GABA\(_{\text{B}}\) receptors in DRG neurons. Astonishingly, however, the precise mechanism of GABA\(_{\text{B}}\) receptor modulation of Ca\(_{\text{V}}\) channels has remained ill defined.

Given Ca\(_{\text{V}}\)'s functional importance in pain pathways, the uncertainty surrounding whether GABA\(_{\text{B}}\) receptors modulate Ca\(_{\text{V}}\) seems remarkable. Several observations may provide insights as to why this question still awaits an answer. In neurons, native Ca\(_{\text{V}}\) is referred to as R-type current, which is poorly defined as the current remaining after blocking the activity of T channels (Ca\(_{\text{V}}\)) with nickel, L channels (Ca\(_{\text{V}}\)) with dihydropyridines, and N, P, and Q channels with α-conotoxins. Because selective and complete pharmacological blockade of any target rarely occurs, doubt has surrounded the notion that native neuronal “R-type” current arises from a pure population of Ca\(_{\text{V}}\) channels (Wilson et al., 2000; Yang and Stephens, 2009); Ca\(_{\text{V}}\) activation occurs over a range of relatively negative voltages (approximately −40 to −50 mV), which has been used as an additional defining characteristic; however, this activation profile overlaps with that of other Ca\(_{\text{V}}\) currents (Williams et al., 1994). A further confounding issue is that R current often contributes a small percentage of the total whole cell Ca\(_{\text{V}}\) current, making it difficult to measure its modulation accurately. In any event, the GABA\(_{\text{B}}\) receptor’s ability to modulate R-type current has not been adequately examined in neurons, nor has its ability to modulate Ca\(_{\text{V}}\) been tested in a recombinant system.

In searching for novel treatments of neuropathic pain, in this issue Berecki et al. have now answered the long-languishing question of whether GABA\(_{\text{B}}\) receptors modulate Ca\(_{\text{V}}\) activity. In so doing, they have advanced three distinct research fields: (1) development of synthetic forms of naturally occurring toxins for treatment of neuropathic pain, (2) Ca\(_{\text{V}}\) current modulation, and (3) GABA\(_{\text{B}}\) receptor function. They demonstrate that cyclized Vc1.1 (cVc1.1), an orally active compound, derived by the cyclization of the synthetic α-conotoxin peptide Vc1.1 (Clark et al., 2010), selectively activates GABA\(_{\text{B}}\) receptors to inhibit recombinant Ca\(_{\text{V}}\) activity. Ca\(_{\text{V}}\) inhibition occurs by a voltage-independent, peristissus toxin (PTX)-sensitive mechanism that requires c-src kinase, a nonreceptor tyrosine kinase (see Okada, 2012), and channel phosphorylation in its proximal C terminus. The signaling pathway is similar to one that mediates voltage-independent inhibition of the Ca\(_{2.2}\) splice variant observed in a subpopulation of nociceptive DRG neurons after GABA\(_{\text{B}}\) receptor stimulation (Bell et al., 2004; Raingo et al., 2007). In characterizing
this mechanism, the authors identify cVc1.1’s inhibition of CaV2.3 activity via GABAB receptor activation as a potential therapeutic strategy for treating certain forms of neuropathic pain.

Pain sensation is complicated. Initially, pain is perceived by nociceptive sensory neurons whose cell bodies reside in DRG (see Costigan et al., 2009; Woolf, 2010). These neurons project to lamina I and II in the dorsal horn to trans-synaptically stimulate ascending spinal neurons. The signal then travels to the thalamus where sensory information is distributed to higher cortical areas. Nociceptive pain is characterized as a high threshold pain activated by immediate, intense stimuli such as noxious heat or a sharp prick. If an injury is sustained, inflammatory pain will occur, which is characterized by hypersensitivity or tenderness from an immune response in the area of an injury that may last for days but usually is reversible. Inflammatory pain, like nociceptive pain, is protective because it serves to discourage use and promote healing. In contrast, neuropathic pain, a disease state of the nervous system (Woolf, 2010), is maladaptive in that pain sensation remains despite the disappearance of the original insult. Often neurons sustain an injury such that the pain threshold decreases so that innocuous stimuli are perceived as painful and are characterized by increased intensity and duration. These changes may persist or become irreversible (Costigan et al., 2009).

Neuropathic pain is remarkably difficult to treat perhaps because of its varied quality, intensity, and source (Costigan et al., 2009). Not surprisingly, several ion channels have been implicated in mediating pain including NaV, CaV, HCN, TRPV1, and various K+ channels, NMDA receptors, and nicotinic receptors (Raouf et al., 2010). Therapeutically targeting any one of these channels to produce analgesia has undesirable side effects because of their widespread distribution both peripherally and centrally. Moreover, developing highly selective channel ligands is difficult, and their off-target binding contributes to further unwanted actions. Lastly, once a promising drug is identified, delivering it to its site of action in patients is problematic because of high biodegradability within and low cellular uptake through the gut, difficulty crossing the blood–brain barrier, and short half-life (Adams and Berecki, 2013). An example of the promises and difficulties of developing effective treatment for pain is the Conus snail peptide toxin MVIIA, a highly selective antagonist of CaV2.2 activity with FDA approval for use as a non-opiate analgesic and marketed under several names (SNX-111, Prialt, Ziconotide). By blocking CaV2.2 activity, MVIIA reduces transmitter release from nociceptive nerve endings (McGivern, 2006). However MVIIA must be injected intrathecally for treating neuropathic pain and has a narrow therapeutic range.

Despite limitations in using toxins to treat neuropathic pain, David Adams’ laboratory in collaboration with many colleagues has synthesized various native and modified Conus snail toxins and tested them for analgesic actions (Clark et al., 2010; Adams and Berecki, 2013). Vc1.1 is a 16-amino acid peptide, originally isolated and purified from Conus victoriae venom, but with the native posttranslational modifications of the toxin removed. It is classified as a cholinergic α-conotoxin based on sequence homology and its pattern of disulfide bridges (Sandall et al., 2003). Vc1.1 selectively antagonizes neuronal α9ε10 nicotinic ACh receptor functioning. In vivo, Vc1.1 acts as an effective and long-lasting analgesic in the chronic constriction injury and partial nerve ligation rat models of human neuropathic pain (Sandall et al., 2003; Satkunanathan et al., 2005). It also accelerates recovery from nerve damage in these two models. When Adams and colleagues tested Vc1.1, they found what seemed to be a second, more potent action of Vc1.1. Vc1.1 appeared to selectively inhibit high threshold (HVA) but not low threshold CaV currents in DRG neurons. At that time, only one other α-conotoxin, Rg1A, was known to inhibit HVA CaV current in DRG neurons; native, reduced, or alkylated forms of Vc1.1 as well as five other α-conotoxins exhibited no inhibitory action on CaV currents. When CaV2.2 activity was blocked with the selective α-conotoxin CVID, Vc1.1 lost its inhibitory action on DRG neuron HVA CaV currents, identifying this modulated HVA CaV current as N current (Callaghan et al., 2008).

Further characterization of Vc1.1’s actions on DRG neurons revealed that N-current inhibition by Vc1.1 occurred independently of rises in intracellular Ca2+, but instead depended on a G protein–dependent signaling cascade because current inhibition was lost in the presence of GDP-β-S, PTX, or pp60src tyrosine kinase inhibitory peptide (Callaghan et al., 2008). From this unusual profile, the group screened a variety of receptor agonists expressed in DRG neurons that might mediate Vc1.1’s inhibitory actions on CaV2.2 activity. Surprisingly, only GABAB receptor antagonists occluded N-current inhibition by Vc1.1 and Rg1A (Callaghan et al., 2008; Callaghan and Adams, 2010). Concomitantly, the GABAB receptor agonist baclofen and Vc1.1 were nonadditive in their inhibitory effects. Vc1.1 and Rg1A inhibited N current in mouse DRG neurons from α9 nicotinic receptor subunit knockout mice, confirming that Vc1.1’s actions occurred independently of nicotinic receptors because α10 is unable to form functional channels without the α9 subunit. A GABAB receptor antagonist blocked toxin-mediated relief from allodynic pain in animal studies (Klimis et al., 2011), consistent with Vc1.1 somehow acting via GABAB receptors and independently of nicotinic receptor subunits.

In most DRG neurons, GABAB receptors inhibit N current by a Gβγ-mediated, voltage-dependent mechanism as noted above. However, in a subpopulation of DRG neurons, baclofen modulates N current by an
revealed that the first (Y1761) but not the second (Y1765) tyrosine is required for voltage-independent current inhibition, presumably because its phosphorylation is required. In contrast, the critical tyrosine-containing amino acid motif found in the C termini of both Ca\textsubscript{V2.2}\textsubscript{e}[37a] and Cav2.3 is absent in Ca\textsubscript{V2.1}, which may explain this channel’s insensitivity to voltage-independent modulation. To test that possibility, a tyrosine residue was swapped in to either of the two homologous sites in the C terminus of Ca\textsubscript{V2.1}, and current was retested for sensitivity to cVc1.1. A small amount of current inhibition was now additional PTX-sensitive, G\textsubscript{B\gamma}-independent and voltage-independent mechanism involving c-src kinase (Diversé-Pierluissi et al., 1997; Raingo et al., 2007). The Lipscombe laboratory and their colleagues previously identified a Ca\textsubscript{V2.2} splice variant selectively enriched in capsaicin-responsive DRG nociceptive neurons that exhibits both voltage-dependent and independent inhibition of N current by baclofen (Bell et al., 2004; Raingo et al., 2007). In exon [37a] of Ca\textsubscript{V2.2}, a 14–amino acid insert in the proximal end of the carboxy-terminal tail contains two tyrosine residues (Y1743 and Y1747), whereas a phenylalanine has replaced the second tyrosine in the mutually exclusive exon [37b]. Voltage-independent inhibition of Ca\textsubscript{V2.2}\textsubscript{e}[37a] requires phosphorylation of the second but not the first tyrosine by c-src kinase. If mutated to phenylalanine (Y1747F) as is found in Ca\textsubscript{V2.2}\textsubscript{e}[37b], voltage-independent inhibition is lost and only voltage-dependent inhibition remains (Raingo et al., 2007). In contrast, complete relief of Ca\textsubscript{V2.2}\textsubscript{e}[37b] current inhibition by baclofen occurs with positive voltage, indicating that the 37b splice variant is insensitive to voltage-independent inhibition by c-src kinase. Collectively, the findings suggest that as with baclofen, Vc1.1 binds to GABA\textsubscript{B} receptors activating the same voltage-independent signaling pathway in DRG neurons to selectively inhibit Ca\textsubscript{V2.2}\textsubscript{e}[37a]. Whether other Ca\textsubscript{V2} family members could be modulated by this pathway remained an open question.

Here, Berecki et al. (2014) tested human recombinant Ca\textsubscript{V2.1} and Ca\textsubscript{V2.3} channels expressed in HEK 293T cells along with the GABA\textsubscript{B} receptor to answer the question of whether other Ca\textsubscript{V2} family members are sensitive to baclofen and/or Vc1.1 modulation by GABA\textsubscript{B} receptors (Fig. 1 A). Although the traditional GABA\textsubscript{B} receptor agonist baclofen inhibited currents from both channels, Vc1.1 and its cyclized form, cVc1.1, inhibited Ca\textsubscript{V2.3} but not Ca\textsubscript{V2.1} currents. The GABA\textsubscript{B} receptor antagonist CGP55845 had no effect on currents itself but antagonized the actions of cVc1.1, indicating that it too must somehow activate the GABA\textsubscript{B} receptor. The characteristics of Ca\textsubscript{V2.1} modulation by baclofen fit classical G\textsubscript{B\gamma}-mediated, voltage-dependent inhibition (Fig. 1 B), as prepulses relieved all the inhibition (Diversé-Pierluissi et al., 1997). In contrast, Ca\textsubscript{V2.3} current inhibition by either baclofen or cVc1.1 shows no voltage-dependent modulation; prepulses were unable to relieve any of the inhibition. As with Ca\textsubscript{V2.2}, inhibition requires c-src kinase because coexpression with a double mutant, inactive src kinase or synthetic pp60c-src kinase inhibitory peptide blocked Ca\textsubscript{V2.3} current inhibition, whereas the opposite treatment, overexpression of wild-type c-src kinase, enhanced inhibition (Fig. 1 C).

Sequence analysis revealed that Ca\textsubscript{V2.3} has two tyrosines (Y1761 and Y1765) in its C-terminal tail homologous to the two tyrosines in Ca\textsubscript{V2.2}\textsubscript{e}[37a]’s proximal C terminus (Raingo et al., 2007). Ca\textsubscript{V2.3} mutagenesis

**Figure 1.** GABA\textsubscript{B} receptor activation elicits both voltage-dependent and voltage-independent inhibition of Ca\textsubscript{V2} current. (A) After GABA\textsubscript{B} receptor activation, Ca\textsubscript{V2} family members exhibit distinct modulation by voltage-dependent (VD) and/or voltage-independent (VI) current inhibition. (B) Example current traces illustrate the two forms of modulation. (Left traces) VD inhibition exhibits slowed kinetics (red traces) compared with control (black) currents that is relieved with positive prepulses (PP). (Right traces) VI inhibition (VI) remains following a PP. (C) Schematic of the proposed mechanism for Ca\textsubscript{V2} inhibition by the two pathways. GABA\textsubscript{B1} and GABA\textsubscript{B2} subunits form the GABA\textsubscript{B} receptor. GABA or baclofen binds to the B1 subunit, activating the receptor. cVc1.1 (V) binds at an undefined site within the interface of the two subunits. Activated G\textsubscript{B\gamma} binds to multiple sites (red squares), whereas G\textsubscript{B} stimulates c-src kinase to phosphorylate tyrosine residues (blue circles) in the C-terminal tail of certain Ca\textsubscript{V2} family members. Black squares, Ca\textsubscript{V2} high affinity–binding sites.
observed with each point mutation, indicating that tyrosine phosphorylation in the proximal C-terminal tail is critical for c-src kinase modulation of channels.

These and previous findings from Adams and colleagues (Callaghan et al., 2008; Callaghan and Adams, 2010; Clark et al., 2010; Cuny et al., 2012) associate three toxins, Rg1A, Vc1.1, and cVc1.1, with GABA<sub>B</sub> receptor activation; however, whether they bind directly to the receptor or to another target protein remains controversial (McIntosh et al., 2009). One B1 and one B2 subunit heterodimerize to form a functional GABA<sub>B</sub> receptor (White and Cousins, 1998). B1 binds GABA in its large extracellular N-terminal domain. B2 is required for high affinity agonist binding to B1, contains a binding site for allosteric modulators, and recruits the G protein (Benke, 2013). Studies using siRNA to lower either B1 or B2 levels found that baclofen, Vc1.1, and Rg1A require both B1 and B2 subunits for N-current inhibition in DRG neurons, consistent with binding to GABA<sub>B</sub> heterodimers (Cuny et al., 2012). When Berecki et al. (2014) transfected HEK 293T cells with only the B1 subunit of the GABA<sub>B</sub> receptor, Ca<sup>V2.3</sup> inhibition by both baclofen and cVc1.1 was lost, supporting—but not proving—the notion that the toxin is somehow interacting with the receptor to elicit its activation. Potentially confounding this notion are findings that very high concentrations of Vc1.1 cannot not displace <sup>3</sup>H-CGP-5462 binding from human recombinant B1/B2 GABA<sub>B</sub> receptors (McIntosh et al., 2009) or from binding in DRG neurons (Adams and Berecki, 2013). These findings indicate that Vc1.1 does not compete with baclofen for a binding site but raise several questions surrounding Vc1.1’s site of action. In particular, if Vc1.1 does bind to the GABA<sub>B</sub> receptor, where is the binding site located, and how does toxin binding activate the receptor? Recently, molecular dynamic simulation studies of toxin binding to GABA<sub>B</sub> receptors identified a putative toxin-binding site at the internal B1/B2 heterodimerization interface (Adams and Berecki, 2013). Whether toxins do indeed bind an ectodomain to activate GABA<sub>B</sub> receptors awaits further binding, structure–function, and crystal structure studies. Once the toxin-binding site is confirmed, functional questions can be addressed, including whether other splice variants of Ca<sup>V2.3</sup>, which contain the critical tyrosine residue in the C terminus tail, are similarly modulated, and where in the pain pathway are Ca<sup>V2.3</sup> and GABA<sub>B</sub> receptors found within the same macromolecular signaling complex.

This insightful study by Berecki et al. (2014) demonstrates how the elaborate details of signaling by individual G protein–coupled receptors (GPCRs), in this case GABA<sub>B</sub> receptors, can give rise to a remarkable level of control over specific members of the Ca<sup>V2</sup> family. By characterizing the details of each branch of GABA<sub>B</sub> receptor signaling, potential therapeutic targets can be identified to strategically treat specific types of pain. Indeed, behavioral studies using siRNAs for Ca<sup>V2.2e[37a]</sup> indicate that this channel selectively participates in mediating normal thermal nociception as well as thermal hyperalgesia that accompanies nerve injury (Altier et al., 2007; Andrade et al., 2010). Splice variants of Ca<sup>V2</sup> channels exhibit different voltage and gating properties that will affect transmitter release and membrane excitability (Bell et al., 2004). Furthermore, whether a channel can be modulated will depend on its splicing and on the colocalization of critical signaling molecules such as c-src kinase within a sensory neuron. Lastly, expression of a particular GPCR within any given sensory neuron will determine whether modulation could be initiated by a transmitter such as GABA or a ligand such as cVc1.1. In this study, rather than using a toxin to directly inhibit ion channel activity, a toxin was used to activate a GPCR to inhibit the activity of a subset of Ca<sup>V2</sup> channels. Moreover, cVc1.1 is ~1,000 times more potent than baclofen in inhibiting Ca<sup>V2.3</sup> currents. Most exciting is previous findings that oral delivery of cVc1.1 produces significant pain relief in the chronic constriction injury rat model of neuropathic pain (Clark et al., 2010). Whether cVc1.1 will find its way into the pharmacological toolbox of treatments for clinical neuropathic pain awaits future studies. Who knew that such a simple question of whether GABA modulates Ca<sup>V2.3</sup> currents would yield a new target, ligand, and potential mechanism for treating neuropathic pain?

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