Kv2 channels create endoplasmic reticulum / plasma membrane junctions: a brief history of Kv2 channel subcellular localization

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ABSTRACT
The potassium channels Kv2.1 and Kv2.2 are widely expressed throughout the mammalian brain. Kv2.1 provides the majority of delayed rectifying current in rat hippocampus while both channels are differentially expressed in cortex. Particularly unusual is their neuronal surface localization pattern: while half the channel population is freely-diffusive on the plasma membrane as expected from the generalized Singer & Nicolson fluid mosaic model, the other half localizes into micron-sized clusters on the soma, dendrites, and axon initial segment. These clusters contain hundreds of channels, which for Kv2.1, are largely non-conducting. Competing theories of the mechanism underlying Kv2.1 clustering have included static tethering to being corralled by an actin fence. Now, recent work has demonstrated channel clustering is due to formation of endoplasmic reticulum/plasma membrane (ER/PM) junctions through interaction with ER-resident VAMP-associated proteins (VAPs). Interaction between surface Kv2 channels and ER VAPs groups channels together in clusters. ER/PM junctions play important roles in inter-organelle communication: they regulate ion flux, are involved in lipid transfer, and are sites of endo- and exocytosis. Kv2-induced ER/PM junctions are regulated through phosphorylation of the channel C-terminus which in turn regulates VAP binding, providing a rapid means to create or dismantle these microdomains. In addition, insults such as hypoxia or ischemia disrupt this interaction resulting in ER/PM junction disassembly. Kv2 channels are the only known plasma membrane protein to form regulated, injury sensitive junctions in this manner. Furthermore, it is likely that concentrated VAPs at these microdomains sequester additional interactors whose functions are not yet fully understood.

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My introduction
Kv2.1 was first discovered by cDNA expression cloning from rat brain in Rolf Joho’s lab in 1989 [2]. This channel, originally designated drk1, encoded a classic delayed rectifier K\textsuperscript{+} channel with a high activation threshold of −15 mV. These investigators postulated that the unusually large cytoplasmic C-terminus may be involved in both channel regulation and subcellular localization. As discussed below, these ideas were correct. Three years later in the Snyder lab, Kv2.2, first termed cdrk1, was cloned from rat taste buds using drk1 sequence to screen a cDNA library [3]. Kv2.1 and Kv2.2 are conserved in the channel forming core, containing the six transmembrane domains and ion conducting pore, and within a limited region of the proximal C-terminus, but otherwise show little sequence identity [3]. mRNA expression analysis indicated areas of overlapping expression in the CNS but suggested that cells predominantly use either Kv2.1 or Kv2.2 [4,5]. Studies of Kv2.1 have dominated the Kv2 literature, partially due to a Kv2.2 cloning artifact which wasn’t realized for some time [6], and Kv2.1 is emphasized in the discussion presented below. However, Kv2.2 shares many properties with Kv2.1, as well as some intriguing differences, which are noted where appropriate.

RNA expression studies and antibody-based protein detection indicate that Kv2 channels are perhaps the most widely expressed voltage-gated potassium channels in the brain.
K⁺ channel genes in terms of tissue distribution. They are present in cortical [7], hippocampal [8], and α-motoneurons [9] but also in retinal bipolar cells [10], cardiac myocytes [11], vascular and gastrointestinal smooth muscle [12,13], and pancreatic beta cells [14,15]. Kv2.1 protein expression is high not only in neurons but even in cell types such as vascular smooth muscle where the outward currents average 100–200 pA [12]. From a functional standpoint, Kv2.1-derived K⁺ currents regulate the action potential waveform in pyramidal neurons undergoing high frequency stimulation [16], cardiac action potential duration [17], vascular smooth muscle membrane potential and contraction [12], beta cell potential and insulin release [14,15,18], and cortical neuron apoptosis [19–22]. Mutations altering Kv2.1 conductance are linked to epilepsy in humans [23,24] and Kv2.1 knockout mice are epileptic, hyperactive, and display defects in spatial learning [25]. Additionally, these mice have elevated serum insulin levels, a prolonged glucose-induced beta cell action potential duration and a diminished firing frequency [18]. However, this review will not focus on the electrical activity of Kv2 channels but will emphasize the non-conducting and structural roles of these channels to induce stable, micron-sized endoplasmic reticulum/plasma membrane junctions that likely function as sites for specialized Ca²⁺ homeostasis and membrane trafficking.

Kv2 channels “cluster” on the membrane

Two years after the cloning of Kv2.1 cDNA the Trimmer group used immunolocalization in rat cerebral cortex to explore the distribution of Kv2.1 [26]. Immunoreactivity was described as neuronal specific, with staining present on the soma and both the apical and proximal dendrites. Importantly, Kv2.1 immuno-staining was described as having a “punctate, membrane-associated nature” as opposed to a fully diffuse localization across somato-dendritic compartments. This was the first evidence that Kv2.1 adopts a unique and enigmatic subcellular localization pattern on the neuronal plasma membrane. Two years later these findings were both supported and expanded upon by the Snyder group that first identified Kv2.2 [4,5]. Whereas the Trimmer group initially focused on pyramidal cells of the rat cortex, Hwang et al. also found α-Kv2.1 staining in Purkinje and granule cells of the cerebellum and granule cells of the olfactory bulb. Furthermore, the Snyder group reported molecular layer staining consistent with stellate and basket cells. Again, Kv2.1 immunoreactivity was described as “punctate” in nature. Given the additional finding that Kv2.1 migrates heterogeneously during SDS gel electrophoresis, the Snyder lab postulated that differences in cellular localization could be due to either Kv2.1 posttranslational modification or “may result from some intrinsic targeting information contained within the amino acid sequences of the two K⁺ channels that directs them to the different subcellular compartments where they are then post-translationally modified” [5]. These hypotheses would prove especially apt in the years to come.

In 1996, evidence suggesting a specific domain of Kv2.1 is important for its localization, as well as the putative role of posttranslational modification, began to solidify [27]. The Trimmer group generated two Kv2.1 C-terminal truncation mutants, ΔC187 and ΔC318, and found that while wildtype Kv2.1 and the ΔC187 mutant both formed 0.5–1.0 μm “clusters” on the cell membrane of MDCK cells, removal of additional amino acids from the C-terminus abolished clustering, as the ΔC318 construct was uniformly distributed. It was also revealed that while wildtype Kv2.1 would cluster in MDCK cells, the clustered phenotype was reported absent in COS-1 cells. Concomitantly, Scannevin et al. noted differences in the gel mobility of Kv2.1 in the two separate cell types, i.e. Western analysis revealed a higher molecular weight in the clustering MDCK cells compared to a lower molecular weight in the non-clustering COS-1 cells, consistent with a link between channel posttranslational modification and clustering. Given earlier findings using in vivo ³²P-labeling [28], it was deemed likely a portion of these posttranslational modifications represented phosphorylation events.

At this time, the mechanism by which Kv2.1 localized into clusters was poorly understood, for all that was known was that a portion of the C-terminus was critical and that clustering may be phosphorylation-dependent. Then in 2000, the Trimmer lab used a series of truncation and
internal deletion mutations to identify a 26 amino acid region within the C-terminus of Kv2.1 that was responsible for clustering [29]. They called this 26 amino acid region the proximal restriction and clustering (PRC) domain. The addition of this domain to the C-terminus of Kv1.5 endowed a clustering phenotype to its otherwise fully diffuse membrane localization while its removal completely abolished clustering in Kv2.1. In addition, several point mutations within the PRC domain also abrogated the clustering phenotype.

Kv2.2 also contains a PRC-like domain and clusters on the membranes of neurons, however it would take 18 years from the time it was first cloned for the field to recognize this fact due to an unfortunate single-nucleotide deletion that resulted in a diffuse truncated protein [6].

As it was previously established that Kv2.1 channels are capable of shaping dendritic \([\text{Ca}^{2+}]\) transients [8], Antonucci et al. in 2001 [30], examined whether Kv2.1 colocalized with \([\text{Ca}^{2+}]\) signaling proteins. They reported colocalization between Kv2.1 clusters and both calsequestrin and ryanodine receptors in pyramidal neurons. In addition, overexpression of a GFP-Kv2.1 construct via adenovirus altered the localization of both calsequestrin and ryanodine receptors, causing them to more closely resemble the exact distribution of Kv2.1 macroclusters. This effect of Kv2.1 overexpression suggested a direct or indirect association between Kv2.1 and these two \([\text{Ca}^{2+}]\) signaling proteins.

### Kv2 clustering is regulated by neuronal activity and insult

In 2004, Misonou et al. [31], proposed a link between neuronal electrical activity, Kv2.1 phosphorylation, and somatic clustering. First, they observed that the induction of class 5 motor seizures by kainate injection declustered Kv2.1 in vivo. Next, 10 \(\mu\)M glutamate treatment of cultured hippocampal neurons to increase spontaneous bursting activity caused a similar declustering with no effect on the total surface level of Kv2.1 protein. These two findings effectively coupled the electrical activity of the cell with the subcellular localization of Kv2.1. The glutamate-induced declustering was also associated with a coincident decrease in the phosphorylation state of the channel; both dephosphorylation and declustering were dosage- and time-dependent and reversible with glutamate washout. Pre-incubation with 100 \(\mu\)M AP-5 (D-2-amino-5-phosphonopentanoate) and 10 \(\mu\)M CNQX (6-cyano-7-nitroquinoxaline-2,3-dione) prior to glutamate treatment blocked dephosphorylation and declustering, suggesting that glutamate was signaling these downstream events through NMDA and AMPA-kainate glutamate receptors. In addition, removal of \([\text{Ca}^{2+}]\) from the media or \([\text{Ca}^{2+}]\) channel block with \(\text{CdCl}_2\) blunted glutamate-induced dephosphorylation and declustering.

Misonou et al. next identified the protein phosphatases responsible for Kv2.1 dephosphorylation. Treatment with okadaic acid, an inhibitor of protein phosphatase 1 and protein phosphatase 2A increased phosphorylation in unstimulated cultures, but failed to prevent glutamate-induced dephosphorylation of the channel. Cyclosporin A, on the other hand, an inhibitor of calcineurin, had no effect on phosphorylation in unstimulated cultures, but did abolish glutamate-induced dephosphorylation and declustering. This led to the conclusion that “glutamate-induced dephosphorylation of Kv2.1 and lateral translocation of Kv2.1 from a clustered to uniform localization are tightly coupled and occur by means of ionotropic glutamate receptor stimulation, leading to \([\text{Ca}^{2+}]\)-dependent activation of calcineurin.” In two 2005 papers [32,33], Misonou and coworkers discovered that Kv2.1 cluster dispersal also occurred after hypoxic or ischemic insult. They postulated that “Kv2.1 clusters form specialized ‘micro-signaling domains’ to regulate somatodendritic \([\text{Ca}^{2+}]\) signaling at these sites” [32]. The glutamate-induced declustering of Kv2.1 is illustrated in the top row of Figure 1.

Contrary to Kv2.1, Kv2.2 is largely resistant to insult-induced declustering. In both mouse brain sections and HEK293 cells, Kv2.2 clusters largely remain intact after hypoxia and elevated cytoplasmic \([\text{Ca}^{2+}]\) [7]. This resistance to declustering is paired with less phosphorylation at rest, as well as less dephosphorylation during insult [7]. The significance of this difference in sensitivity to declustering stimuli, if one exists, has yet to be examined.
Relationship between Kv2.1 clustering and channel activity

Throughout early studies of Kv2.1 localization there were few solid ideas with respect to the physiological significance of Kv2.1 clustering. However, glutamate treatment affected more than just the phosphorylation state and subcellular localization of Kv2.1. In the final finding of the Misonou et al., 2004 paper, they concluded that glutamate-induced declustering resulted in a > 20 mV hyperpolarizing shift in the $V_{1/2}$ of neuronal $I_K$, from +15.5 ± 0.5 mV to −8.4 ± 0.5 mV. This shift was reversible after a 2 h glutamate washout and was blocked by calcineurin inhibitors. This last conclusion shaped the field’s thinking in regards to the physiological significance of Kv2.1 clustering, effectively linking the channel’s voltage-sensitivity to its subcellular localization. The next year Misonou et al. went further, hypothesizing that clusters on the soma and proximal dendrites acted as “on/off” switches that regulated the intrinsic excitability of the cell [33]. It was thought at the time that clusters represented reservoirs of high-threshold Kv2.1 channels, ready to be hyper-activated upon cluster release when needed.

Throughout the early 2000s, the overall hypothesis was that Kv2.1 was tethered to cytoskeletal elements adjacent to the ER and near ionotropic receptors, kinases, and phosphatases that served to either maintain or disrupt Kv2.1 clustering and thus regulate channel voltage-dependence [34]. As is often the case, a more complex picture arose as we learned more.

Working under the assumption that Kv2.1 clustering regulated the voltage-dependence of channel activation, our group, the Tamkun laboratory, began cell biological studies of Kv2.1 behavior on the membrane surface to better understand the mechanism and dynamics underlying Kv2.1 localization and the regulation of channel function. Surprisingly, both traditional FRAP approaches and newly developed single molecule, quantum dot-based, imaging indicated that Kv2.1 channels were not statically tethered within clusters at the cell surface as originally proposed [34], but rather were corralled within a cytoskeletal fence [35]. Here the clustered channels had a lateral mobility similar to Kv2.1 outside the clusters, which agreed with that expected for freely diffusing membrane proteins. Thus, it seemed likely that phosphorylation of the C-terminus allowed Kv2.1 to interact with an additional unknown protein and this additional mass prohibited the channel from diffusing through the cluster’s perimeter boundary. Non-clustered channels did not have this posttranslational modification, could not interact with this unknown accessory protein, and therefore were too small to be hindered by the cytoskeletal architecture and were free to diffuse into
and out of clusters. Supporting this hypothesis were the dual findings that in HEK293 cells Kv2.1 clusters tend to be entirely bordered by actin filaments and that disruption of actin with swinholide A dissolved Kv2.1 clusters [36]. A mechanism by which Kv2.1 was corralled in a phosphorylation-dependent manner by a cytoskeletal perimeter fence neatly fit all available data at the time.

Since these data cast doubt onto the prevailing model for Kv2.1 clusters, i.e. where static tethering regulates the voltage-dependence of activation, we decided to directly examine the function of clustered Kv2.1 using HEK293 cells expressing GFP-Kv2.1 and a cell-attached patch clamp approach [37]. While cell-attached patches on membrane regions devoid of Kv2.1 clusters, but populated with freely diffusing Kv2.1 channels, exhibited macroscopic delayed rectifier currents, little to no channel activity was detected when the patch clamp pipet was placed directly onto Kv2.1 clusters, as if the clustered channels were non-conducting even at +60 mV. When comparing the whole cell gating current magnitude to ionic currents in transfected HEK293 cells a striking mismatch was found, suggesting that all the Kv2.1 channels responded to a depolarizing stimulus with a moving S4 domain but only a small percentage of these transitions resulted in pore opening. This finding agreed well with early work from the Pongs lab [38] that indicated a discrepancy between Kv2.1 gating and ionic current in *Xenopus laevis* oocytes, where less than 1% of the gating channels actually opened. To test whether Kv2.1 clusters acted as reservoirs of non-conducting channels that were activated upon release, we next measured whole cell currents before and after inducing Kv2.1 declustering via either actin depolymerization to dissolve the hypothesized diffusion-limiting fence, or alkaline phosphatase in the patch clamp pipet to dephosphorylate the clustered channel [37]. Both treatments resulted in declustering, however while the alkaline phosphatase treatment resulted in the expected shift of voltage dependence, declustering via actin depolymerization did not. Neither treatment increased current density, which would be expected if non-conducting channels suddenly became conducting once declustered. These findings were contrary to the prevailing theories about the channel, as they demonstrated that clustering per se has little impact on channel function. While phosphorylation seems to both govern some aspects of channel electrical activity as well as clustering, location and conductance were not inextricably linked.

Following studies would confirm these findings. Baver and O’Connell [39] showed that the NMDA receptor-based regulation of Kv2.1 activity occurs in the absence of Kv2.1 clustering. In addition, our group would later find that the non-conducting state was regulated by surface channel density and not location on the cell surface [40]. The non-conducting state existed in C-terminal truncation mutants that lack the PRC domain and cannot cluster and the percentage of non-conducting channels increased as a function of surface channel number [40]. Further supporting a separation between localization and conductance, in 2015 the Trimmer lab found that the cell cycle-dependent regulation of Kv2.1 clustering in COS-1 cells, which is due to changes in Kv2.1 phosphorylation, does not affect Kv2.1 currents [41]. While we now know that uncoupling of S4 movement from pore opening is regulated by channel density, the exact mechanism underlying this disconnect remains a mystery.

### Non-conducting functions of Kv2.1 clusters

If the clustered channels are not, and do not become, conducting upon declustering, what is their purpose, especially considering the gating current data that indicates non-conducting Kv2.1 channels still sense changes in membrane potential? The high levels of Kv2.1 protein in multiple cell types suggest a structural role and these high levels would also mandate the non-conducting state, for without this, neurons would be electrically silenced.

Non-conducting Kv2.1 had already been linked to exocytosis, for the Lotan group found that Kv2.1 facilitates dense core vesicle release from neuroendocrine cells independently of potassium flux via Kv2.1 interaction with syntaxin [42,43]. Unfortunately, since this work did not employ imaging, no relationship was drawn between these results and Kv2.1 localization. Motivated by
this Lotan work, our lab next asked whether the Kv2.1 clusters acted as insertion platforms for membrane protein delivery to the plasma membrane [44]. Approximately 85% of both Kv2.1 and Kv1.4 channel plasma membrane insertion events occurred at the Kv2.1 cluster perimeter. As Kv1.4 is freely diffuse, this localized delivery is not specific to cluster-resident proteins. In addition, since endocytosis was also observed at the perimeter of Kv2.1 clusters, these microdomains were postulated to act as membrane trafficking hubs [44,45]. Very recent work from the MacDonald and Gaisano labs [14,15] further demonstrates that Kv2.1 clusters regulate insulin exocytosis in pancreatic beta cells.

Du and colleagues [8], using a combination of immunohistochemical and electron microscopy approaches, had previously found that Kv2.1 clusters were often localized on neuronal cell membranes directly apposed to both ER/PM junctions and astrocyte membranes. These junctions, or discs of flattened cortical ER located 5–8 nm from the intracellular side of the cell membrane, were continuous with either the smooth or rough ER. This association between Kv2.1 clusters and membrane junctions had also been reported in rat α-motoneurons where the Kv2.1 clusters are apposed to cholinergic C-type synaptic terminals [9]. As discussed above, the Trimmer group had reported association between Kv2.1 clusters and ER proteins such as ryanodine receptors. Thus, still searching for non-conducting functions associated with Kv2.1 clusters, our lab returned to the idea that Kv2.1 clusters existed at sites of ER/PM junctions as first seen by Du et al. in 1998. Given that the clustered, but non-conducting, Kv2.1 channels still likely responded to voltage, and being influenced by the work of Kurt Beam [46], we wondered whether they could act in a fashion similar to L-type Ca\(^{2+}\) channels at the skeletal muscle triad junction where Ca\(^{2+}\) channels couple membrane depolarization to Ca\(^{2+}\) release from internal stores via mechanical activation of ryanodine receptors. Therefore, we further examined the relationship between Kv2.1 and the ER.

In untransfected HEK293 cells, endogenous ER/PM junctions were small in size and transient in nature as observed using TIRF imaging of fluorescent ER markers as well as electron microscopy [47]. In contrast, following Kv2.1 expression, cortical ER adjacent to the Kv2.1 clusters was dramatically remodeled. These results suggested that Kv2.1 binds to the ER surface and tethers the ER to the PM. This tethering generates the observed Kv2.1 cluster phenotype and explains the high Kv2.1 mobility within the clusters, assuming the ER binding partner, whether it be protein or lipid, is mobile within the ER membrane. Thus, Kv2.1 channel bound to the ER is corralled by the edge of the ER membrane at the point where this organelle turns inward towards the more internal cytoplasm. Further confirming a non-conducting role of Kv2.1 is to induce ER/PM junctions, glutamate-induced Kv2.1 declustering in cultured hippocampal neurons resulted in concomitant retraction of ER localized under Kv2.1 clusters after cluster dispersal as illustrated in [47]. The next obvious step required to understand the Kv2.1-ER relationship was for us to identify the Kv2.1 binding partner on the ER cytoplasmic face, be that protein or lipid. As both Kv2.1 and Kv2.2 form clusters, our lab privately hypothesized that a common mechanism behind this clustering existed and that Kv2.2 also likely interacted with an unknown ER resident to form ER/PM junctions. Indeed, recent efforts have confirmed that cortical ER remodeling is a function of both Kv2.1 and Kv2.2 [48,49].

**Kv2 channels and the VAPs**

With the hope that Kv2.1 interacts with an ER protein, our group used APEX-based proximity biotinylation and identified a 33 kD protein that likely represented the Kv2.1 binding partner [48]. With critical input from Tim Levine of University College, London, and using a combination of redistribution assays, FRET, and siRNA knockdown experiments, we confirmed that this 33 kD protein was the ER VAMP-associated protein, or VAP. We also found, as expected, that both Kv2.1 and Kv2.2 interact with VAPs. It is this Kv2-VAP interaction which tethers the ER to the PM and is thus responsible for Kv2 clustering. For additional information, see Johnson et al. [48].

VAP was first discovered in *Aplysia californica* and was originally referred to as VAP-33 as the
polypeptide migrated at 33 kD during SDS gel electrophoresis [50]. Four years later, three mammalian homologues were uncovered: VAPA, VAPB, and VAPC [51]. While VAPA was almost identical to the Aplysia VAP-33, VAPB and VAPC were novel homologues with the latter, VAPC, being a VAPB splice variant that lacks the transmembrane domain found at the C-terminus of VAP A and B [51,52]. All VAP proteins contain an immunoglobulin-like β sheet on their N-terminus followed by a coiled-coil domain [53]. A VAP consensus sequence, FK(V/I)KTT(VA)P(K/R)(K/R)YCV(K/R)P, within their N-terminal major sperm protein (MSP) domain may contribute to VAP oligomerization [54] in addition to the transmembrane domain [51,55]. A missense mutation within VAPB (P56S; highlighted in the above sequence) causes intracellular aggregation of the VAPB mutant and is associated with late-onset spinal muscular atrophy and amyotrophic lateral sclerosis [56]. This mutation recruits both wildtype VAPB and VAPA to these immobile aggregates and results in cell death in culture systems [57]. In addition, the P56S mutation interferes with VAPBs ability to interact with other proteins via its MSP domain [58]. Deletion of the Drosophila homolog, DVAP-33A, results in a decrease in the number of synaptic boutons while overexpression leads to an increase in bouton number and decrease in average size [59]. There is also a rich literature connecting the VAPs to viral replication which raises the question of whether Kv2-induced ER/PM junctions are in some way co-opted for viral transmission [60–62].

The interaction between the VAPs and their partners occurs through the binding of the MSP domain within VAP to an FFAT (two phenalalanines in an acidic tract) motif located on the binding partner. The consensus sequence for FFAT motifs is EFFDAxE, though this sequence can tolerate a high degree of variability [63]. A negative upstream flanker region aids in the interaction between VAP and the FFAT motif although serine phosphorylation can replace acidic residues in this region, thus allowing activation by phosphorylation [64–66]. While Kv2 channels do not possess a stereotypical FFAT sequence, we identified a noncanonical FFAT motif located within the PRC domain of both Kv2 channels using a series of protein chimeras. In Kv2.1 this motif is SFISCAT while in Kv2.2 it is SFTSCAT. Phosphorylated serine residues both within and upstream of this motif may generate the negative charge necessary for VAP interaction, which could explain the Kv2 phosphorylation requirement to form ER/PM junctions. This Kv2 FFAT motif-VAP MSP domain interaction at the ER/PM interface is illustrated in Figure 2.

Figure 2. Mechanism of ER/PM junction formation and Kv2 clustering. The binding of a noncanonical FFAT motif (SFISCAT in Kv2.1, pictured here) within the Kv2 C-terminus to VAPs embedded within the cortical ER membrane generates both membrane contact site formation and the Kv2 clustered phenotype.
that the VAP-dependent Kv2.1 clustering also generates a domain potentially involved in cell-to-cell communication since the Kv2.1 beta subunit, AMIGO, is an adhesion molecule [67–69]. The bottom row of Figure 1 illustrates the effect of glutamate-induced Kv2.1 dephosphorylation that frees the channel C-terminus from the ER VAPs, thus resulting in Kv2.1 declustering, VAP dispersal, and cortical ER remodeling.

Based on discussions with our laboratory, the Trimmer group was able to independently confirm the Kv2-VAP interaction. Importantly, the Trimmer lab generated anti-VAP antibodies and used immuno-staining to confirm that endogenous VAPs were concentrated at Kv2 clusters in both rat brain and cultured rat hippocampal neurons [70]. This concentration of endogenous VAPs at native Kv2.1 clusters complements our redistribution experiments in HEK293 cells where Kv2 induction of ER/PM contacts concentrated ER VAPs at this microdomain [48]. Our FRET data, which suggests VAPs may homo-oligomerize within Kv2.1-induced ER/PM junctions independently of Kv2.1 binding, provides a possible mechanism for this robust redistribution. In addition, VAP homo-oligomers could simultaneously bind Kv2 channels to form junctions and bind other interactors, thus concentrating additional proteins to these membrane contact sites (see Figure 2). VAPs have a large interactome, including kinases, vesicle trafficking proteins, kinesins, AKAPs and lipid transfer proteins [65,71], all of which may play important roles at Kv2-induced ER/PM junctions.

**Current model for Kv2.1-induced ER/PM contact sites**

The model illustrated in Figure 3 agrees with data obtained in transfected HEK293 cells, cultured hippocampal neurons, rat brain slices and intact rat brain [48,70]. Here serine phosphorylation adjacent to the Kv2.1 C-terminal FFAT motif allows channel binding to VAPs present on the cytoplasmic surface of ER. This binding not only tethers the ER membrane to within 10 nm of the PM but it also concentrates VAPs to this microdomain and generates the Kv2 clustered phenotype. Given that VAP homo-oligomerization may occur at these sites to form a VAP scaffold, Kv2 channels effectively generate a site where additional VAP interactors can concentrate. This could provide these sites with the capability to perform further functions, from lipid transfer between ER and PM, to aiding ER calcium store refilling.

**Kv2-based regulation of ER/PM junctions and sensitivity to cellular insult**

Individual Kv2.1 channels are capable of switching between being freely diffusive or clustered under homeostatic conditions [36], suggesting constant dynamic control over these microdomains that can

![Figure 3. Model of the Kv2.1-induced ER/PM junction. Calcineurin-dependent dephosphorylation of the Kv2.1 C-terminus results in release from the ER VAPs, Kv2.1 declustering, loss of the ER/PM junction, VAP redistribution, and cortical ER retraction. The clustered Kv2.1 channels are proposed to be non-conducting under resting conditions [37]. Both the Kv2.1-containing and Kv2.1-free ER/PM junctions are trafficking hubs where dense core vesicle exocytosis [14,15] and membrane protein delivery is concentrated [44], perhaps in part due to these membrane contact sites residing in, or forming, “holes” in the cortical actin [36]. However, Kv2.1-containing membrane junctions are proposed to also concentrate SNARE proteins, for Kv2.1 contains VAMP-2, SNAP25 and syntaxin binding domains [72–74].](image-url)
be pushed either towards enlarging microdomains or their disassembly. Given the correct stimuli, Kv2 dephosphorylation breaks the Kv2-VAP link. The ER then retracts into the cytoplasm and VAPs redistribute throughout the ER. It is likely that the functions specifically associated with the Kv2-induced ER/PM contacts are then lost. Inversely, other stimuli can lead to the formation of Kv2-induced ER/PM contacts or enlargement of preexisting junctions, thus creating conditions under which functions can now be performed or performed to a greater degree.

There are several known conditions under which Kv2-induced ER/PM junctions are impacted, some of which this review has already covered. Recent electron microscopy work in cultured hippocampal neurons performed by Dr. Tao-Cheng [75,76] found a significant decrease in both size and number of ER/PM junctions after depolarization with high K⁺ or NMDA treatment. These findings fit neatly with the known effect neuronal activity has on Kv2.1 channel localization [31]. While mass dephosphorylation of Kv2.1 can both cause declustering as well as a hyperpolarizing shift in voltage dependence, we now know that these two events are not inextricably linked as once believed [37]. For instance, the cell cycle dependent ebb and flow of clustering behavior in COS-1 cells is not associated with changes in channel electrical activity [41]. Also, while Kv2.1 clusters predominantly contain nonconducting channels, the regulation of the nonconducting state occurs independently of channel localization [37,40]. Clearly, more work will need to be accomplished on the non-conducting state.

Kv2-induced ER/PM junctions are also impacted by cellular insult and disease. As previously mentioned, Kv2.1 clusters disperse in response to hypoxic or ischemic conditions [32,33]. Kv2.2 appears to be less sensitive to this insult-induced declustering [7] and this sensitivity or resistance to ER/PM junction rearrangement may be one reason cells have opted to use one channel over the other. Kv2.1 clusters are enlarged in some rat models of Alzheimer’s disease with a coupled decrease in K⁺ current density that may be due to increased oxidative stress during development [77]. Human mutations predicted to disrupt Kv2.1-VAP interaction without impacting channel electrical activity are associated with developmental delay [78]. There is also a great body of work performed by the Fyffe lab examining Kv2.1 in rat motoneurons and C-bouton synapses [9,79–81]. In 2014 this group discovered that Kv2.1 clusters dispersed in response to peripheral nerve injury in both ipsilateral and contralateral limbs [81]. Curiously, without reinnervation Kv2.1 clusters reformed over time, but with reinnervation they did not.

Determining the functions associated with Kv2-induced ER/PM junctions and how this microdomain can be targeted to enhance neuroprotection is an area of new importance. Recently, work by the Aizenman and Shah labs revealed that Kv2.1-induced ER/PM junctions are necessary for apoptosis [82]. Interfering with the formation of Kv2.1-induced ER/PM junctions, without affecting channel electrical activity, disrupted the apoptotic stimulus-induced trafficking of Kv2.1 to the surface, thus interfering with the increase in K⁺ efflux necessary for apoptosis [19,20]. This inhibition of Kv2.1 delivery improved neuroprotection during oxidative stress [82]. These results fit well with our earlier work demonstrating that Kv2.1 ER/PM junctions serve as trafficking hubs for the delivery of Kv2.1 and other ion channels [44]. Future studies on this topic are certainly warranted.

ER/PM junction composition and function in the intact brain

ER/PM junctions were first described in muscle cells by Porter & Palade in 1957 [83]. Their existence in neurons would be described just five years later [84]. Recent focused ion beam-scanning electron microscopy (FIB-SEM) based reconstructions of ER/PM junctions in rodent brain by the De Camilli group, indicate these membrane contact sites represent up to 12% of the neuronal soma [85]. These membrane contacts also exist in dendrites along the dendritic shaft and in axons within the varicosities which are the predominant sites for neurotransmitter release. Whether Kv2.1 is involved in these non-somatic contacts remains an open question.

In the intact brain these membrane junctions are predicted to contain multiple components at any one time, (e.g. Kv2 channels, junctophilins,
STIM proteins, voltage-gated Ca\textsuperscript{2+} channels, Trp channels, extended synaptotagmins, etc.), and the specific composition of ER/PM junctions will depend on cell type. Junction composition will also be affected by cellular state, for example, the STIM-Orai complex will reside in junctions during periods of ER calcium store depletion, but not during other times, and we now know that Kv2-VAP complexes are regulated by neuronal activity and cellular insult in a phosphorylation-dependent manner [41,47,48,70,86–88]. In addition, ER/PM junctions within different compartments of a single cell may display different properties. For example Kv2.1-induced junctions on the AIS are more resistant to CO\textsubscript{2} and kainate-induced declustering than junctions on the soma [89].

Accumulating evidence suggests ER/PM junctions are not homogeneous, but that within these micro-domains there exists nanodomain heterogeneity. For example, different proteins capable of forming ER/PM junctions require different spacing between the PM and ER membrane. This requirement, as has been demonstrated by the Balla group, can cause exclusion of certain proteins from specific contact site regions based on size [91]. It is possible that within intact brain tissue a singular ER/PM junction is subdivided into various nanodomains which perform different functions. This may account for the “donut-like” localization pattern of some ER/PM junction resident proteins; the proteins which require less distance between membranes are forced to the center of the junction and those that require more distance between membranes are forced to the periphery [91]. Endogenous Kv2.1 clusters in neurons are sometimes reported to have such a “donut-like” pattern with proteins such as neuregulin-2 occupying the center space [91].

Since multiple proteins may be contributing to ER/PM junction structure in the intact brain, events which cause Kv2-VAP unbinding may not disrupt the junction, but rather may only lead to a loss of Kv2, VAP, and potential VAP interactors from this location, thus providing perhaps a more nuanced response to certain stimuli. The interplay between these various nanodomains and the inherent dynamics within the entire junction is an obvious area for future investigation as is the functionality provided by the Kv2-VAP interaction and how this functionality is disturbed during cellular insult.

**Future directions**

Now that we have begun to understand the molecular composition of Kv2-induced ER/PM junctions, future studies can confront the question so far largely missing from the history reviewed above: why use a non-conducting ion channel to form ER/PM junctions in the first place, i.e. what functionality does the cell gain from this arrangement over any other protein? Perhaps the voltage-sensing S4 domains of Kv2.1 communicate membrane potential to downstream effectors as occurs at the skeletal muscle PM/sarcoplasmic reticulum junction. Another potential advantage of using a K\textsuperscript{+} channel to form ER/PM contacts relates to ER Ca\textsuperscript{2+} store homeostasis. Perhaps the clustered Kv2 channels become conducting during ER Ca\textsuperscript{2+} store depletion, thus providing a localized K\textsuperscript{+} counter current to enhance any localized Ca\textsuperscript{2+} influx. More work is necessary to determine the exact role of the Kv2.1 syntaxin binding domain in exocytosis, especially considering recent findings suggesting that this domain regulates dense core vesicle exocytosis at Kv2.1 clusters and can rescue insulin release in diabetic pancreatic beta cells [14,15]. The Kv2.1 beta subunit, AMIGO, which is a cell adhesion molecule [68], could serve to position ER/PM junctions adjacent to cell-cell contact sites. Indeed, intercellular AMIGO interactions could be responsible for the symmetrical ER/PM contacts observed by the De Camilli group at some somatic contact sites [85]. And why would such activity- and insult-sensitive structures exist if not for some form of intercellular communication? As this story of Kv2.1 clustering continues to move forward we anticipate both the resolution of these questions and new surprises.

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