Regulation of neuronal excitation–transcription coupling by Kv2.1-induced clustering of somatic L-type Ca\(^{2+}\) channels at ER-PM junctions

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In mammalian brain neurons, membrane depolarization leads to voltage-gated Ca\(^{2+}\) channel-mediated Ca\(^{2+}\) influx that triggers diverse cellular responses, including gene expression, in a process termed excitation–transcription coupling. Neuronal L-type Ca\(^{2+}\) channels, which have prominent populations on the soma and distal dendrites of hippocampal neurons, play a privileged role in excitation–transcription coupling. The voltage-gated \(K^+\) channel Kv2.1 organizes signaling complexes containing the L-type Ca\(^{2+}\) channel Cav1.2 at somatic endoplasmic reticulum–plasma membrane junctions. This leads to enhanced clustering of Cav1.2 channels, increasing their activity. However, the downstream consequences of the Kv2.1-mediated regulation of Cav1.2 localization and function on excitation–transcription coupling are not known. Here, we have identified a region between residues 478 to 486 of Kv2.1’s C terminus that mediates the Kv2.1-dependent clustering of Cav1.2. By disrupting this Ca\(^{2+}\) channel association domain with or with a cell-penetrating interfering peptide, we blocked the Kv2.1-mediated clustering of Cav1.2 at endoplasmic reticulum–plasma membrane junctions and the subsequent enhancement of its channel activity and somatic Ca\(^{2+}\) signals without affecting the clustering of Kv2.1. These interventions abolished the depolarization-induced and L-type Ca\(^{2+}\) channel-dependent phosphorylation of the transcription factor CREB and the subsequent expression of c-Fos in hippocampal neurons. Our findings support a model whereby the Kv2.1-Ca\(^{2+}\) channel association domain-mediated clustering of Cav1.2 channels imparts a mechanism to control somatic Ca\(^{2+}\) signals that couple neuronal excitation to gene expression.

Significance

In hippocampal neurons, gene expression is triggered by electrical activity and Ca\(^{2+}\) entry via L-type Cav1.2 channels in a process called excitation–transcription coupling. We identified a domain on the voltage-gated \(K^+\) channel Kv2.1 that promotes the clustering of L-type Cav1.2 channels at endoplasmic reticulum–plasma membrane junctions in the soma of neurons. Importantly, we discovered by disrupting this domain that the Kv2.1-mediated clustering of Cav1.2 at this somatic microdomain is critical for depolarization-induced excitation–transcription coupling.

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The authors declare no competing interest.

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channels also increases Cav1.2 open probability (35, 37). However, neither the molecular determinants of LTCC targeting to somatic Kv2-associated ER-PM junctions nor the physiological consequences of Ca<sup>2+</sup> signals produced at these sites are known. Here, we identified a Ca<sup>2+</sup> channel association domain (CCAD) within Kv2.1’s cytoplasmic C terminus that is necessary for Cav1.2 localization to ER-PM junctions. The disruption of the Kv2.1 CCAD selectively impairs the ability of Kv2.1 to recruit Cav1.2 to ER-PM junctions but does not impact Kv2.1 clustering at these sites. A cell-penetrating interfering peptide based on the CCAD sequence caused the spatial and functional decoupling of Cav1.2 from Kv2.1-associated ER-PM junctions and significantly reduced Cav1.2 channel activity and LTCC-dependent gene expression in neurons. Our data provide additional evidence that Cav1.2 channels on neuronal somata play a prominent role in excitation–transcription coupling and that the Kv2.1-dependent regulation of the clustering, activity, and localization of this population of Cav1.2 channels shapes their physiological function.

**Results**

**Identification of a Ca<sup>2+</sup> Channel Association Domain in the Cytoplasmic C Terminus of Kv2.1.** Cav1.2 channels colocalize with Kv2.1 channels in the somata of hippocampal neurons (35), as illustrated by confocal images obtained from a mouse brain section immunolabeled with antibodies against Kv2.1 and Cav1.2 (Fig. 1A). When Cav1.2 channels are expressed in human embryonic kidney (HEK) 293T cells, they also cluster with Kv2.1 at ER-PM junctions (Fig. 1B). To define the molecular requirements of Kv2.1-mediated rearrangement of Cav1.2 channels, we coexpressed Cav1.2 tagged with monomeric RFP [Cav1.2-RFP (38)], with a series of previously described Kv2.1 deletion mutants and chimeras (39, 40) with altered Kv2.1 C-terminal domains (Fig. 1C) in HEK293T cells. We used total internal reflection fluorescence (TIRF) microscopy to image the PM and near-PM populations of the immunolabeled and fluorescent proteins. We determined the degree of colocalization between the Kv2.1 isoforms and Cav1.2 by measuring Pearson’s correlation coefficient (PCC) values of the respective fluorescence signals (Fig. 1D). As a quantitative index of Cav1.2 channel clustering, we used the coefficient of variation of Cav1.2-RFP fluorescence intensity (CV, obtained by dividing the SD of the fluorescence signal by the mean fluorescence signal). A larger CV value indicates a higher degree of channel clustering (27).

Similar to previous results (35), we found that full-length Kv2.1 colocalized with Cav1.2-RFP channels, as indicated by high PCC values (Fig. 1D). Coexpression with Kv2.1 also resulted in the enhanced clustering of Cav1.2 channels, as indicated by the increased Cav1.2 CV values upon Kv2.1 coexpression relative to control values (Fig. 1E). We next investigated

**Fig. 1.** Identification of a domain within the C terminus of Kv2.1 required for coclustering with Cav1.2. (A) Confocal optical section of the dentate gyrus granule cell layer acquired from an adult mouse brain section immunolabeled for Cav1.2 (red) and Kv2.1 (green). Hoescht staining (blue) is shown in the merge panel (Scale bar, 2 μm in both the main panel and the magnified inset whose location in the main panels are outlined.) (B) TIRF images of HEK293T cells transfected with Cav1.2 (red) and WT and chimeric Kv2.1 constructs (green) (Scale bar, 10 μm and holds across all panels). (C) Diagram of Kv2.1 chimeras, and truncation, and internal deletion mutants used to identify the region in Kv2.1 required for the reorganization of Cav1.2 channels. In the left column, coclustering of WT and chimeric Kv2.1 constructs with Cav1.2 is indicated by Y (yes) or N (no), as determined by displaying a significantly lower PCC value as compared to Kv2.1 WT. Within the region encoding the Kv2.1 C terminus, the putative CCAD is highlighted in red, and the PCC domain in green. (D) PCC values of the fluorescence signals of WT and chimeric Kv2 constructs and Cav1.2 channel fluorescence signals expressed in HEK293T cells. Each point represents a single cell; bars are mean ± SEM (****P < 0.0001 and ***P = 0.0009; one-way ANOVA followed by Dunnett’s post hoc test versus Kv2.1 WT). (E) CV values of Cav1.2 channel fluorescence when coexpressed with Kv2 channels in HEK293T cells. Each point represents a single cell; bars are mean ± SEM (****P < 0.0001 and ***P < 0.0006; one-way ANOVA followed by Dunnett’s post hoc test versus Cav1.2 alone).
whether the domain underlying the Kv2.1-mediated recruitment of Cav1.2 to ER-PM junctions was located on the cytoplasmic C terminus of Kv2.1 by evaluating a chimeric protein containing the entire Kv2.1 C terminus (amino acid residues 411 to 853 of rat Kv2.1) appended onto the unrelated Kv1.5 channel (Kv1.5N-Kv2.1C [411 to 853]) chimera, which localizes to ER-PM junctions (29), coclustered with Cav1.2 channels and increased Cav1.2 CV values relative to Cav1.2 alone (Fig. 1 B, D, and E). This result demonstrated that the cytoplasmic C terminus of Kv2.1 contained determinants sufficient to both mediate its clustering at ER-PM junctions and to stabilize clustered Cav1.2 channels at these sites.

To further refine which region(s) within the Kv2.1 C terminus underlies the reorganization of Cav1.2 channels, we next used a chimeric protein in which a much smaller portion of the Kv2.1 C terminus (amino acid residues 536 to 600, which contain the PRC domain) was appended to Kv1.5. Although the Kv1.5N-Kv2.1C (536 to 600) chimera still exhibits Kv2.1-like clustering at ER-PM junctions (29) due to the presence of an intact PRC domain (Fig. 1 B and C), it was incapable of recruiting Cav1.2 channels. The evaluation of a set of Kv2.1 C-terminal internal deletion and truncation mutants supported that the PRC-dependent formation of organized ER-MP junctions was necessary but not sufficient for the Kv2.1-mediated reorganization of Cav1.2 channels and defined a CCAD between residues 411 and 520 of the Kv2.1 C terminus required for Kv2.1 to reorganize Cav1.2 channels (Fig. 1C).

To gain insights into this putative CCAD-containing region of Kv2.1, we determined the frequency of nonsynonymous polymorphisms in the coding region of the human KCNB1 (Kv2.1) gene in genomic and exome sequences from over 141,000 individuals (https://gnomad.broadinstitute.org/). The lack of genetic variation within a region is indicative of functional importance (41). There is little variation in regions that encode domains critical to the function of Kv2.1 as a voltage-sensing and K⁺-conducting channel (Fig. 2A). Conversely, the C terminus of Kv2.1 displays much more genetic variation, with the exception of the invariant PRC domain (residues 584 to 597) and a charged domain within the proximal C terminus region important for the cell-surface expression of Kv2.1 [within residues 444 to 477 (42)]. A closer inspection of the region-encoding residues 477 to 520 of the Kv2.1 C terminus revealed an additional invariant domain corresponding to residues 475 to 493 of human Kv2.1 (Fig. 2A). This same region of Kv2.1 has previously been identified to mediate interaction with the vesicle fusion SNARE protein syntaxin-1A (Stx-1A) in vitro and in neuroendocrine and pancreatic beta cells (43–45). However, whether this region of Kv2.1 is also involved in regulating the subcellular distribution and function of Cav1.2 had not been experimentally addressed.

**Point Mutants and Interfering Fragments of the Kv2.1 CCAD Disrupt Cav1.2 Channel Clustering with Kv2.1.** A previous investigation of the Stx-1A binding domain of Kv2.1 revealed a critical role for tryptophan residues at positions 484 and 486 in mediating the ability of a Kv2.1-derived synthetic peptide to bind Stx-1A in vitro (46). Given the high-sequence conservation of W484 and W486 across vertebrates, the lack of genetic variation at these sites in Kv2.1 in the human population, and the results of our Kv2.1 chimera and deletion mutant experiments, we tested whether mutating these same residues would also impact the ability of Kv2.1 to regulate the subcellular organization of Cav1.2. We introduced either the W484A or W486A point mutations in a DsRed-tagged, non-K⁺-conducting point mutant of Kv2.1 (DsRed-Kv2.1 P404W). We chose to use this nonconducting mutant background as the ability of Kv2.1 to organize ER-PM junctions (31) and to increase Cav1.2 localization at these sites is independent of its ability to conduct K⁺ (35), and using this mutant avoids any additional and indirect contributions of the K⁺-conducting function of Kv2.1, for example on membrane potential, to its regulation of Cav1.2. We will hereafter refer to this nonconducting point mutant as “Kv2.1 WT,” in reference to its WT cytoplasmic C terminus.

When expressed in HEK293T cells alone, Kv2.1 WT and the W484A- and W486A point mutants showed similar overall expression and PM clustering (SI Appendix, Fig. S1). We used TIRF imaging to evaluate the ability of these mutants to colocalize with and cluster coexpressed Cav1.2-GFP (Fig. 2 B–H). Similar to our previous observations (35), Kv2.1 WT colocalized with Cav1.2-GFP and increased its PM clustering (Fig. 2B) relative to Cav1.2-GFP expressed with DsRed alone (Fig. 2C). Neither increased colocalization nor clustering of Cav1.2 was seen upon coexpression with a Kv2.1 PRC domain point mutant (S586A) (Fig. 2D) that itself does not cluster at ER-PM junctions (29). Kv2.1 constructs containing the W484A or W486A mutations still clustered like Kv2.1 WT (Fig. 2 E and F), which was expected as they retain an intact PRC domain. However, when compared to Kv2.1 WT, these mutants were deficient in their ability to recruit and reorganize Cav1.2-GFP channels, as indicated by the reduced Cav1.2 PCC and CV values relative to Kv2.1 WT (Fig. 2 I–K). Together, these results demonstrate that W484 and W486 are necessary for the Kv2.1-dependent reorganization of Cav1.2 channels.

We next tested how the introduction of a cell-penetrating peptide, containing amino acids (a.a.) 478 to 486 of the putative Kv2.1 CCAD, impacted the ability of Kv2.1 to recruit Cav1.2 channels in HEK293T cells. This peptide [TAT-CCAD, also called TAT-C1a(b) (44)] contains these 9 a.a. of Kv2.1 fused to a TAT motif that directs its efficient uptake into cells following its addition to culture medium (47). The overnight treatment of HEK293T cells coexpressing Kv2.1 WT and Cav1.2-GFP with TAT-CCAD reduced coclustering of Kv2.1 and Cav1.2 without disrupting Kv2.1 clustering itself (Fig. 2 G–K). A version of this peptide in which the Kv2.1 sequence was scrambled (TAT-Scr) had no impact on the ability of Kv2.1 WT to reorganize Cav1.2 (Fig. 2 G–K). These results support that the CCAD found within a.a. 478 to 486 in Kv2.1 is both necessary and sufficient for the ability of Kv2.1 to increase the clustering of Cav1.2.

**The Kv2.1 CCAD Is Required for Kv2.1-Mediated Increases in Cav1.2 Clustering and Function.** We next applied a bimolecular fluorescence complementation approach to test the hypothesis that as the TAT-CCAD peptide disrupts Kv2.1-Cav1.2 association, it would interfere with the previously demonstrated ability of Kv2.1 to increase Cav1.2 coclustering and function (35, 37). For these experiments, we used a system employing Cav1.2 channels fused to the N- or C terminus of the split-Venus fluorescent protein, consisting of Cav1.2-VN155 ([I152L]) and Cav1.2-C155, respectively, VN155 ([I152L]) and Cav1.2-VN155 do not fluoresce in isolation, but they reconstitute the fluorescent Venus protein when brought into close proximity by interaction of the separately tagged Cav1.2 subunits (38). We have used these reporters to show that depolarization-induced Ca²⁺ influx through Cav1.2 channels increases spontaneous Cav1.2-Cav1.2 interactions (38, 48), as shown in Fig. 3d, and that coexpression with Kv2.1 further increases the probability of Cav1.2-Cav1.2 channel interactions (37). Here, we found that the application of TAT-CCAD reduced the Kv2.1-dependent increases in Cav1.2-Venus fluorescence intensity without affecting the intrinsic fluorescence of DsRed-Kv2.1 itself (Fig. 3B). In contrast, cells treated with the control TAT-Scr peptide exhibited typical Kv2.1-mediated increases in Cav1.2 split-Venus fluorescence (Fig. 3B).

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These results indicated that the TAT-CCAD peptide selectively disrupted the ability of Kv2.1 to increase Cav1.2-Cav1.2 interactions. Based on our observation that the Kv2.1 W484A and W486A mutants were impaired in their ability to reorganize Cav1.2 channels, we hypothesized that the TAT-CCAD cell-penetrating peptide competes with the CCAD on Kv2.1 for binding to Cav1.2 or to an intermediary protein that promotes Cav1.2 localization at ER-PM junctions. Because the Kv2.1 W484A and W486A mutants would already be deficient for this interaction, we predicted that we would observe no additional effect of the TAT-CCAD peptide on Cav1.2-Cav1.2 interactions in cells expressing Kv2.1 CCAD mutants. For these experiments, we used the Kv2.1 W486A mutant, which displayed a larger reduction in its ability to reorganize Cav1.2 than did Kv2.1 W484A (Fig. 2 I and J). Cells expressing the Kv2.1 W486A point mutant did not display depolarization-induced elevations in split-Venus fluorescence, as seen for Kv2.1 WT (Fig. 3B). Moreover, neither the TAT-CCAD nor the TAT-Scr peptide had any additional effect on Cav1.2 Venus fluorescence in Kv2.1 W486A-expressing cells. These results support the model that an intact Kv2.1 CCAD domain is necessary for the Kv2.1-mediated increase in Cav1.2-Cav1.2 interactions, and that the TAT-CCAD cell-penetrating peptide acts as a competitive inhibitor of the direct or indirect association of Kv2.1 with Cav1.2 via this domain.

While collecting reconstituted fluorescence data in the split-Venus experiments, we also determined how disrupting the Kv2.1 CCAD impacted Cav1.2-Venus channel Ca\textsuperscript{2+} currents. Cav1.2 currents increase in parallel with stimulus-induced increases in Cav1.2-Cav1.2 interactions (38, 48) (Fig. 3C) and are further potentiated by Kv2.1-mediated clustering (37) (Fig. 3D). We found that the treatment of cells with the TAT-CCAD peptide, but not the TAT-Scr peptide, prevented these stimulus-induced increases in Cav1.2 currents (Fig. 3 D and E). Cells expressing the Kv2.1 W486A variant also exhibited a smaller increase in Ca\textsuperscript{2+} currents relative to those observed in postdepolarization conditioning in the presence of Kv2.1 WT (Fig. 3 F and G). Treatment with either of the TAT peptides produced...
no further effects in the Kv2.1 W486A-expressing cells (Fig. 3 F and G). Overall, the impact of these manipulations on Cav1.2 Ca2\(^{+}\) currents paralleled their effects on Cav1.2-Cav1.2 interactions as reported by split-Venus fluorescence.

Consistent with its role in enhancing Cav1.2 channel activity, coexpression with Kv2.1 also enhances depolarization-induced Ca2\(^{+}\) influx in Cav1.2-expressing HEK293T cells, as measured with the Ca2\(^{+}\)-sensitive dye Fluo-4 (35). To test whether the TAT-CCAD peptide would have any direct effects on Cav1.2 function in the absence of Kv2.1, we compared the effects of TAT-CCAD treatment on depolarization-induced Ca2\(^{+}\) influx in Cav1.2-expressing HEK293T cells in the presence and absence of Kv2.1 coexpression. We found that the TAT-CCAD peptide impaired the Kv2.1-dependent increases in depolarization-evoked Ca2\(^{+}\) influx in HEK293T cells relative to cells with no treatment or treated with the TAT-Scr peptide (Fig. 3 H–J). However, neither the TAT-CCAD nor TAT-Scr peptides had any effect on Ca2\(^{+}\) influx in cells expressing Cav1.2 alone, showing that these peptides impacted Cav1.2 function in a Kv2.1-dependent manner. As expected from the results of the Cav1.2-Venus experiments, Kv2.1 W486A did not increase Cav1.2-mediated Ca2\(^{+}\) influx as did Kv2.1 WT, and TAT-CCAD treatment did not impact Ca2\(^{+}\) influx in cells expressing Kv2.1 W486A. These observations show that the impact of TAT-CCAD on Cav1.2 function requires the coexpression of Kv2.1 WT. Taken together, these data further support that Kv2.1 channels promote Cav1.2 clustering and activity and that an intact Kv2.1 CCAD is required for this effect.

Disruption of the Kv2.1 CCAD Reduces Endogenous Cav1.2 Channel Activity and Its Clustering with Kv2.1 at ER-PM Junctions. We next tested how these CCAD mutations and the TAT-CCAD cell-penetrating peptide impacted the function and localization of endogenous Cav1.2. We first employed an optical approach to measure single-Cav1.2 channel activity in the form of Ca2\(^{+}\) sparklets, which are local elevations in intracellular Ca2\(^{+}\) levels.
produced by the opening of a single or small cluster of Cav1.2 channels (49). We measured Ca\(^{2+}\) sparks in INS-1 cells, a rat cell line derived from pancreatic β-cells with well-characterized Cav1.2-mediated Ca\(^{2+}\) currents (50). The compact morphology of these cells permitted effective control of the membrane potential to measure spontaneous openings of endogenous Cav1.2 channels at polarized membrane potentials. Similar to our previous observations with Cav1.2 channels expressed in HEK293T cells (35), we found that the expression of Kv2.1 WT (in the non-K\(^{+}\)-conducting P404W background) in INS-1 cells increased sparklet activity (NP) and the number of sparklet sites present in the TIRF footprint relative to control cells (Fig. 4 A–C). The fitting of multipeak Gaussians to the all-points histograms of the calibrated Ca\(^{2+}\) sparklet records (Fig. 4 D and E) indicated a quantal amplitude of 35.4 ± 2.3 nM Ca\(^{2+}\), similar to LTCC-mediated Ca\(^{2+}\) sparks measured under similar conditions in other cell types (48, 51, 52). Cumulative frequency distribution plots revealed a greater frequency of multiquantal Ca\(^{2+}\) sparklets, corresponding to multiple simultaneous LTCC openings, in the DsRed-Kv2.1 group compared to the control DsRed or DsRed-Kv2.1 W486A groups (Fig. 4D). Moreover, the treatment of cells with the LTCC agonist Bay K8644 increased the Ca\(^{2+}\) sparklet activity of control DsRed and DsRed-Kv2.1 W486A groups but did not further enhance the activity of the DsRed-Kv2.1 group (Fig. 4 B, C, and E). These observations suggest that exogenously expressing Kv2.1 WT brought the endogenous Cav1.2 channels close to their maximum open probability, similar to results obtained for exogenous Cav1.2 channels coexpressed with Kv2.1 in HEK293T cells (35). In contrast, Ca\(^{2+}\) sparks in cells expressing Kv2.1 W486A resembled those in cells lacking exogenous Kv2.1 expression, supporting the hypothesis that an intact CCAD is required for the Kv2.1-mediated potentiation of endogenous Cav1.2 channels.

We next examined how the TAT peptides impacted somatic LTCC-dependent Ca\(^{2+}\) signals in cultured rat hippocampal neurons transfected with GCaMP-Kv2.1. We previously determined that Ca\(^{2+}\) sparks reported by GCaMP-Kv2.1 are generated by the spontaneous opening of LTCCs juxtaposed to ER-localized RyRs at Kv2.1-associated ER-PM junctions (35). We found that treatment with TAT-CCAD occluded spontaneous Ca\(^{2+}\) sparks and reduced somatic Ca\(^{2+}\) sparks triggered by the treatment of cells with the LTCC activator Bay K8644 (Fig. 4 F and G). In contrast, more than half of cells treated with the control TAT-Scr peptide displayed spontaneous Ca\(^{2+}\) sparks. Together, these data suggest that the CCAD-mediated association of Cav1.2 with Kv2.1 enables the generation of LTCC- and RyR-mediated Ca\(^{2+}\) signals at somatic ER-PM junctions, and that the TAT-CCAD peptide interferes with the clustering of Cav1.2 at these sites.

We tested this possibility by assessing how the treatment of neurons with TAT peptides impacted the spatial coupling of somatic Cav1.2 channels with RyRs (35). We found that the TAT-CCAD treatment of hippocampal neurons decreased the colocalization of Cav1.2 with RyRs (Fig. 5 A, B, and E). In contrast, TAT-CCAD treatment did not impact the spatial coupling of Kv2.1 with RyRs (Fig. 5 A, B, E, and F). While treatment with the TAT-CCAD peptide reduced the association of Kv2.1 with Cav1.2 in the soma, it did not alter the localization of Cav1.2 in distal dendrites, as evidenced by the lack of an effect on the extent of colocalization of Cav1.2 with the postsynaptic protein PSD-95 (Fig. 5H). These results suggest that interfering with the function of the Kv2.1 CCAD selectively disrupts the Kv2.1-mediated localization of Cav1.2 channels at somatic ER-PM junctions. The findings that the TAT-CCAD peptide suppressed the Kv2.1-mediated increases in somatic Cav1.2 clustering and function, but had no impact on Cav1.2 alone or on its localization on distal dendrites that lack Kv2.1, suggest that it can be used as a selective modulator of somatic but not distal dendritic Cav1.2 channel function.

We next determined how treatment with the TAT-CCAD peptide influenced the spatial proximity of endogenous Kv2.1 and

![Fig. 4. Disruption of the CCAD abolishes Kv2.1-dependent increases in endogenous LTCC activity. (A) [Ca\(^{2+}\)]i records from representative Ca\(^{2+}\) sparklet sites recorded in INS-1 cells transfected with the indicated constructs before (upper traces) and after (lower traces) treatment with 500 nM Bay K8644. (B and C) Bar graphs of mean ± SEM. NP, (B) and sparklet density (C) before and after treatment with Bay K8644; (B): *p = 0.0297 and **p = 0.051; C: *p = 0.0201 and **p = 0.0077; one-way ANOVA followed by Dunnett’s post hoc test versus DsRed-C1 vehicle). Each point represents a single cell. (D and E) All-points histograms and cumulative frequency distributions (solid lines) of Ca\(^{2+}\) sparklet data recorded from control cells (D) or cells treated with 500 nM Bay K8644 (E). Insets show unobscured histograms for multiquantal events (black: DsRed-C1; red: Kv2.1 WT, and blue: Kv2.1 W486A). The data were fit with a multicomponent Gaussian function (dashed lines). (F) Confocal image of a hippocampal neuron transfected with GCaMP3-Kv2.1. Arrows indicate Kv2.1 clusters displaying spontaneous Ca\(^{2+}\) sparks whose fluorescence intensity profiles are plotted in adjacent panel (Scale bar, 10 μm). (G) Quantification of Ca\(^{2+}\) sparks sites observed in neurons treated with TAT-CCAD or TAT-Scr peptides. Each point represents a single cell; bars are the average of mean experimental values ± SEM (**p = 0.0027; Student’s t test).](https://doi.org/10.1073/pnas.2110094118)
Cav1.2 channels to one another using a proximity ligation assay [PLA (53)] to detect Kv2.1 and Cav1.2 channels that are within 40 nm of each other (37). The overnight treatment of neurons with TAT-CCAD reduced the Kv2.1-Cav1.2 PLA signal, an effect not seen with TAT-Scr (Fig. 6 A–C), supporting that this Kv2.1-derived cell-penetrating peptide interferes with the association of endogenous Kv2.1 and Cav1.2 channels. Consistent with the model that the Kv2.1-mediated clustering of Cav1.2 increases the association of Cav1.2 channels with one another (37) and our split-Venus results in HEK293T cells (Fig. 3), TAT-CCAD treatment also reduced the Cav1.2-Cav1.2 PLA signal (Fig. 6 A and B). The Kv2.1-Kv2.1 PLA signal was not impacted by TAT-CCAD treatment (Fig. 6 A and B), consistent with imaging results in HEK293T cells (Figs. 1 and 2), showing that while the CCAD is necessary and sufficient for the association of Kv2.1 with Cav1.2, it is not involved in the clustering of Kv2.1 itself.

**Disruption of the Kv2.1 CCAD Suppresses Excitation–Transcription Coupling.** Neuronal LTCCs including Cav1.2 uniquely contribute to membrane depolarization-dependent changes in transcription factor activation by mediating Ca\(^{2+}\) influx within specialized and compartmentalized signaling complexes (3, 17, 54–57). Excitation-transcription coupling can be initiated at distal synaptic sites (58–60), although somatic depolarization in itself is sufficient (61) and certain forms require activation of somatic LTCCs (56). While much is known of the protein complexes involved in mediating Cav1.2-dependent forms of synaptic plasticity and excitation-transcription coupling triggered at distal synaptic sites (12, 62), the nature of signaling complexes underlying somatic LTCC-mediated excitation-transcription coupling has not been defined. We previously established that Kv2.1 organizes Cav1.2 in close proximity to somatic ER-PM junctions in hippocampal neurons (35). The results presented here indicate that disruption of the Kv2.1 CCAD permits selective decoupling of Cav1.2 from these sites. Therefore, we tested how selective disruption of Cav1.2 recruitment to Kv2-organized ER-PM junctions would impact depolarization-induced Cav1.2 Ca\(^{2+}\) influx-dependent changes in transcription factor activation and gene expression. Phosphorylation of the transcription factor CREB at S133 is triggered by Ca\(^{2+}\) influx through LTCCs and stimulates expression of immediate early genes encoding multiple proteins, including the transcription factor c-Fos, that serve important roles in learning and memory (17, 55, 63–65). We treated cultured hippocampal neurons with TAT peptides overnight and investigated how membrane potential depolarization induced by elevated extracellular K\(^{+}\) affected nuclear CREB phosphorylation and induction of c-Fos expression. We first silenced the synaptic and intrinsic activity of neurons for two hours using a buffer solution containing CNQX and AP5 to block glutamate receptors, and tetrodotoxin (TTX) to block voltage-gated Na\(^{+}\) channels (17); TAT-CCAD or TAT-Scr peptides were also included in this preincubation medium. Neurons were then depolarized for 90 seconds with 45 mM K\(^{+}\) buffer, also supplemented with CNQX, AP5, and TTX, predicted to depolarize the membrane potential to > –19 mV (17), to induce cell autonomous LTCC-dependent transcription factor activation (66) (Fig. 7A). We found that TAT-CCAD peptide treatment reduced the depolarization-triggered increases in the levels of nuclear pCREB (Fig. 7 B and C) and c-Fos expression (Fig. 7 D and E), supporting a critical role for LTCCs clustered at Kv2-associated ER-PM junctions in mediating LTCC excitation-transcription coupling. Treatment with TAT-Scr peptide was without effect. The impact of TAT-CCAD treatment was mimicked by treatment with the LTCC blocker nimodipine, which as previously reported also blocked induction of pCREB phosphorylation and c-Fos expression (66) (Fig. 7 C and E). Importantly, nimodipine occluded any further effects of the TAT-CCAD treatment on both pCREB
and c-Fos induction (Fig. 7C and E), showing that the effects of this cell penetrating peptide were mediated through its impact on LTCCs. Given their impact in increasing Cav1.2 clustering and activity, we next evaluated whether exogenous expression of non-conducting Kv2.1 channels in hippocampal neurons could impact LTCC-dependent changes in pCREB or c-Fos. We performed the same high K+-induced depolarization assay using neurons transfected with nonconducting (i.e., in the P404W point mutant background) Kv2.1 WT. We found that expression of Kv2.1 WT in cultured neurons increased K\textsuperscript{+}-induced CREB phosphorylation (Fig. 7F) and c-Fos (Fig. 7G) expression relative to neurons expressing DsRed. However, hippocampal neurons expressing the Kv2.1 CCAD mutant W486A had levels of nuclear pCREB (Fig. 7F) and c-Fos (Fig. 7G) comparable to neurons lacking exogenous Kv2.1 expression. Together, these results support that the somatic population of LTCCs associated with Kv2.1 plays a prominent role in excitation-transcription coupling, and that Kv2.1-mediated increases in LTCC clustering and activity, and/or recruitment to ER-PM junctions regulate LTCC-dependent excitation-transcription coupling (Fig. 7H).

Discussion

In this study, we made three fundamental observations regarding the organization of somatic Kv2.1, Cav1.2, and ER-PM junctions and their role in regulating excitation-transcription coupling. First, we discovered that Kv2.1-mediated ER-PM junction formation and the recruitment of Cav1.2 channels are separable functions of Kv2.1 mediated by distinct domains (the PRC and CCAD, respectively) within its C terminus. Second, the targeted disruption of the Kv2.1 CCAD impaired Kv2.1-dependent increases in Cav1.2 clustering and activity, and the association of Cav1.2 with ER-PM junctions, but unlike mutations in the PRC domain, did not impact the clustering of Kv2.1 at these sites. Third, the selective uncoupling of somatic LTCCs from Kv2.1-mediated ER-PM junctions in hippocampal neurons using CCAD mutants or a cell-penetrating interfering peptide suppressed LTCC-dependent excitation-transcription coupling in the form of depolarization-induced nuclear CREB phosphorylation and c-Fos expression. These findings suggest that the subset of Cav1.2 channels localized in specific microdomains at somatic Kv2.1-mediated ER-PM junctions play a key role in generating the Ca\textsuperscript{2+} signals that trigger depolarization-induced gene expression. Based on these results, we propose a model in which the CCAD links Kv2.1 to somatic Cav1.2 channels, promoting their clustering at ER-PM junctions, modulating their activity, and impacting their role in excitation-transcription coupling (Fig. 7H).

Our results show that LTCC-dependent excitation-transcription coupling was impaired by the selective, CCAD-dependent uncoupling of somatic Cav1.2 channels from Kv2.1-mediated ER-PM junctions. It has long been recognized that LTCCs play a privileged role in the Ca\textsuperscript{2+} influx mediating important forms of excitation-transcription coupling in brain neurons (63, 67, 68). However, for the most part, these studies were not able to distinguish the roles of the distinct populations of LTCCs on the neuronal soma and on distal dendrites near dendritic spines. Given its proximity to the nucleus, Ca\textsuperscript{2+} influx via somatic LTCCs is expected to be especially important for triggering gene expression (56, 61, 69, 70). Recently, additional experimental support for the concept of “soma to nucleus signaling” (61) was provided by studies employing neurons grown in microfluidic chambers that allowed for the separate interrogation of the contribution of the somatic and distal dendritic populations of LTCCs (56). These studies revealed a specific role for somatic LTCCs in mediating the Ca\textsuperscript{2+} influx underlying the calcineurin-dependent activation of the NFAT transcription factor (56). As Kv2.1 expression is restricted to the soma and proximal dendrites, the impact of the Kv2.1-mediated regulation of LTCCs would presumably be restricted to these same neuronal compartments and would not impact LTCCs on distal dendrites at/near dendritic spines. Our findings show that treatments with either the TAT-CCAD cell-penetrating peptide or the LTCC blocker nimodipine inhibit excitation–transcription coupling to a similar extent and that the treatments occlude one another. Taken together with the other results presented here, this provides additional evidence that LTCCs on the soma and proximal dendrites, and specifically those localized in specific microdomains at Kv2.1-containing ER-PM junctions, play a prominent role in mediating the Ca\textsuperscript{2+} increases that trigger excitation–transcription coupling. The Kv2.1-mediated increase in the localization of Cav1.2 with ER-PM junctions results in their increased spatial and functional association with ER-localized RyRs and more robust Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+} release.
Nuclear pCREB levels in hippocampal neurons transfected with the indicated constructs and treated with 4.7 mM KCl. (A) Schematic of experimental protocols for 45 mM K+ depolarization-induced phosphorylation of CREB at Ser133 (pCREB) or the expression of c-Fos (NC medium: neuronal culture medium). (B) Representative images of nuclei and pCREB immunofluorescence in neurons treated with TAT-Scr or TAT-CCAD under each condition (4.7K: 4.7 mM K+; 45K: 45 mM K+; and 45K + nim: 45 mM K+ + 10 μM nimodipine) (Scale bar, 10 μm). (C) Nuclear pCREB levels in hippocampal neurons treated as indicated. Each point represents a single cell (solid circles) or the mean value from each of three independent experiments (open circles); bars are the mean ± SEM (***P < 0.0001; one-way ANOVA followed by Dunnett’s post hoc test). (D) As in C, but for nuclear c-Fos (****P < 0.0001; one-way ANOVA followed by Dunnett’s post hoc test versus TAT-Scr 4.7K). (E) Nuclear pCREB levels in hippocampal neurons transfected with the indicated constructs and treated with 4.7 mM K+ or 45 mM K+. Each point represents a single cell; bars are mean ± SEM (****P < 0.0001 and **P = 0.0070; one-way ANOVA followed by Tukey’s post hoc test). (F) As in F, but for nuclear c-Fos levels (**P = 0.0079 and *P = 0.0295; one-way ANOVA followed by Tukey’s post hoc test). (H) Working model. The Kv2.1 CCAD promotes the targeting of LTCCs to somatic ER-PM junctions to enhance Ca2+ signals that trigger gene expression.

Fig. 7. LTCCs associated with Kv2.1-mediated ER-PM junctions play a prominent role in excitation–transcription coupling. (A) Schematic of experimental protocols for 45 mM K+ depolarization-induced phosphorylation of CREB at Ser133 (pCREB) or the expression of c-Fos (NC medium: neuronal culture medium). (B) Representative images of nuclei and pCREB immunofluorescence in neurons treated with TAT-Scr or TAT-CCAD under each condition (4.7K: 4.7 mM K+; 45K: 45 mM K+; and 45K + nim: 45 mM K+ + 10 μM nimodipine) (Scale bar, 10 μm). (C) Nuclear pCREB levels in hippocampal neurons treated as indicated. Each point represents a single cell (solid circles) or the mean value from each of three independent experiments (open circles); bars are the mean ± SEM (***P < 0.0001; one-way ANOVA followed by Dunnett’s post hoc test). (D) As in C, but for nuclear c-Fos (****P < 0.0001; one-way ANOVA followed by Dunnett’s post hoc test versus TAT-Scr 4.7K). (E) Nuclear pCREB levels in hippocampal neurons transfected with the indicated constructs and treated with 4.7 mM K+ or 45 mM K+. Each point represents a single cell; bars are mean ± SEM (****P < 0.0001 and **P = 0.0070; one-way ANOVA followed by Tukey’s post hoc test). (F) As in F, but for nuclear c-Fos levels (**P = 0.0079 and *P = 0.0295; one-way ANOVA followed by Tukey’s post hoc test). (H) Working model. The Kv2.1 CCAD promotes the targeting of LTCCs to somatic ER-PM junctions to enhance Ca2+ signals that trigger gene expression.

at these sites (35), which could further amplify somatic Ca2+ signals and impact nuclear transcription factor activation. Decoupling LTCCs from Kv2.1 and somatic ER-PM junctions by CCAD interference could impair the LTCC-dependent activation of critical signaling proteins, reducing the subsequent phosphorylation of CREB and gene expression, although our data do not rule out additional effects on other components of the signaling pathway. The future use of the TAT-CCAD cell-penetrating peptide as a selective modulator of LTCCs on the soma and proximal dendrites, but not of those in distal dendrites at/near dendritic spines, could provide further insights into the contribution of these distinct populations of LTCCs in neuronal function and plasticity.

A key finding in our study is that Kv2.1, through its CCAD, plays a nonconducting structural role to regulate LTCCs that does not require its K+ conductance. Our data suggest that Kv2.1–Cav1.2 interactions require an intact CCAD and that the TAT-CCAD peptide interferes with this association. One limitation of our current study is that it does not reveal the precise molecular mechanism of this interaction, including whether it is direct or indirect. The CCAD sequence identified here has also been shown to be necessary for Kv2.1 binding to the SNARE protein Stx-1A (44–46), raising the possibility that Stx-1A may influence the association of LTCCs with Kv2.1 and mitochondrial Ca2+ overload in ischemia-induced neuronal cell death (76), it will be of interest to determine whether the impact of TAT-CCAD peptide on suppressing Cav1.2-mediated Ca2+ signaling plays any role in its neuroprotective effects.

In summary, we have identified VAP proteins (29) and numerous LTCC subunits (35) in our proteomic analyses of components of native brain Kv2.1-containing ion channel complexes, we have not detected Stx-1A. However, Cav1.2 has been shown to interact with Stx-1A in neuroendocrine cells (72, 73) and when heterologously expressed in HEK293 cells (74). These observations raise the possibility that in certain cell types Stx-1A may influence the association of Kv2.1 with Cav1.2 by acting as a binding intermediary, a competitive inhibitor, or in another role. The TAT-CCAD peptide has been previously shown to exhibit neuroprotective effects in in vitro and in vivo ischemic stroke models (44). Our data show that this cell-penetrating peptide reduces Cav1.2 channel activity and the spatial and functional association of Cav1.2 with RyRs at neuronal ER-PM junctions, sites that are also rich in mitochondria (75). Given the prominent role of increased intracellular Ca2+ and mitochondrial Ca2+ overload in ischemia-induced neuronal cell death (76), it will be of interest to determine whether the impact of TAT-CCAD peptide on suppressing Cav1.2-mediated Ca2+ signaling plays any role in its neuroprotective effects.
of LTCC-dependent Ca\textsuperscript{2+} signals at somatic ER-PM junctions in initiating depolarization-triggered gene transcription in neurons.

Materials and Methods

Experimental procedures are described in detail in SI Appendix.

Animals. All procedures using mice and rats were approved by the University of California, Davis Institutional Animal Care and Use Committee and performed in accordance with the NIH Guide for the Care and Use of Laboratory Animals (77). Animals were maintained under standard light-dark cycles and allowed to feed and drink ad libitum. Experiments using hippocampal neurons were performed using cultures obtained from pooling neurons from Sprague-Dawley rat fetuses of both sexes. Adult male mice were used in immunohistochemistry experiments.

Cell Culture, Transfection, and TAT Peptide Treatments. HEK293T cells (ATCC No. CRL-3216) were cultured and transfected using Lipofectamine 2000 using standard procedures (details in SI Appendix). Neuronal cultures were prepared and maintained using a standard protocol (78) (details in SI Appendix). TAT-Scr and TAT-CCAD peptides were synthesized by GenScript at over 95% purity (peptide sequences are provided in SI Appendix, Table S2) and diluted to 1 \(\mu\)M in culture medium. For the evaluation of 45 mM K\textsuperscript{+}-induced CREB S133 phosphorylation or c-Fos expression, DIV20 neurons were treated overnight with TAT peptides (details in SI Appendix).

Immunolabeling and Imaging of Fixed Brain Sections, Cultured Neurons, and Cell Lines. Preparation and immunolabeling of mouse brain sections was as previously described (27) (details in SI Appendix). Cultured neurons and HEK293T cells were immunolabeled as described previously (35) (details in SI Appendix).

Live Cell Ca\textsuperscript{2+} Imaging and Bimolecular Fluorescence Complementation. HEK293T cells transfected with the respective Cav channel subunits were imaged 48 h after transfection and after overnight treatments with TAT peptides (details in SI Appendix). Electrophysiological recordings were acquired at room temperature using an Axopatch200B amplifier and a Digidata1440 digitizer using pClamp 10.2 software (Molecular Devices) (details in SI Appendix).

Sparklets. Ca\textsuperscript{2+} sparklets were recorded using a dual-TIRF imaging/patch clamp system using INS-1 cells 48 h after transfection with respective Kv2.1 plasmids, employing Fluor-Sf (Invitrogen No. FI4221) as a Ca\textsuperscript{2+} reporter (details in SI Appendix).

PLA. The Duolink In Situ PLA kit (Sigma) was used to detect Cav1.2 and Kv2.1 in DIV20 neurons treated overnight with 1 \(\mu\)M TAT-CCAD or TAT-Scr and fixed 15 h later (details in SI Appendix).

Experimental Design and Statistical Analysis. For all datasets presented in this study for which statistical analyses were performed, measurements were imported into GraphPad Prism for statistical analysis and presentation. Reported values are means \pm SEM. Exact \(P\) values are reported in each figure legend. Paired datasets were compared using a Student’s t test if the data passed a normality test; a nonparametric test was used otherwise. For all experiments, at least two independent cultures were used for experimentation; the number of samples analyzed is noted in each figure or figure legend.

Data Availability. All study data are included in the article and/or SI Appendix.

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