RIM-binding proteins recruit BK-channels to presynaptic release sites adjacent to voltage-gated Ca\(^{2+}\)-channels

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Abstract

The active zone of presynaptic nerve terminals organizes the neurotransmitter release machinery, thereby enabling fast Ca\(^{2+}\)-triggered synaptic vesicle exocytosis. BK-channels are Ca\(^{2+}\)-activated large-conductance K\(^+-\)channels that require close proximity to Ca\(^{2+}\)-channels for activation and control Ca\(^{2+}\)-triggered neurotransmitter release by accelerating membrane repolarization during action potential firing. How BK-channels are recruited to presynaptic Ca\(^{2+}\)-channels, however, is unknown. Here, we show that RBPs (for RIM-binding proteins), which are evolutionarily conserved active zone proteins containing SH3- and FN3-domains, directly bind to BK-channels. We find that RBPs interact with RIMs and Ca\(^{2+}\)-channels via their SH3-domains, but to BK-channels via their FN3-domains. Deletion of RBPs in calyx of Held synapses decreased and decelerated presynaptic BK-currents and depleted BK-channels from active zones. Our data suggest that RBPs recruit BK-channels into a RIM-based macromolecular active zone complex that includes Ca\(^{2+}\)-channels, synaptic vesicles, and the membrane fusion machinery, thereby enabling tight spatio-temporal coupling of Ca\(^{2+}\)-influx to Ca\(^{2+}\)-triggered neurotransmitter release in a presynaptic terminal.

Keywords active zone; BK-channels; nano-domain coupling; neurotransmitter release; voltage-gated calcium channels

Subject Categories Membrane & Intracellular Transport; Neuroscience; Physiology

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Introduction

At the active zone of presynaptic terminals, action potentials (APs) cause voltage-gated Ca\(^{2+}\)-channels to open. The resulting transient increase in cytoplasmic Ca\(^{2+}\) induces neurotransmitter release by triggering synaptic vesicle exocytosis (Walter et al, 2011; Sudhof, 2013). The number and location of voltage-gated Ca\(^{2+}\)-channels at the active zone and the coupling of voltage-gated Ca\(^{2+}\)-channels to synaptic vesicles are critical in determining the strength and plasticity of synapses (Meinrenken et al, 2002; Modchang et al, 2010; Eggermann et al, 2011). Several mechanisms control the size and duration of the Ca\(^{2+}\)-signals at the active zone, thereby generating temporally restricted micro- and nano-domains of Ca\(^{2+}\) that determine the extent of release (e.g., see Fedchyshyn & Wang, 2005; Bucurenciu et al, 2008; Vyleta & Jonas, 2014). Among the most important of these mechanisms is the control of the AP duration by BK-channels (Fakler & Adelman, 2008; Contet et al, 2016; Griguoli et al, 2016). BK-channels are Ca\(^{2+}\)-activated large-conductance K\(^+-\)channels that limit the duration of an AP, thereby controlling the extent of Ca\(^{2+}\)-channel opening and of neurotransmitter release per AP. BK-channels are enriched in presynaptic terminals and are present in a complex with voltage-gated Ca\(^{2+}\)-channels, but it is unknown how BK-channels are recruited to presynaptic terminals and how they are molecularly connected with voltage-gated Ca\(^{2+}\)-channels (Roberts, 1993; Berkefeld et al, 2006; Indriati et al, 2013).

Voltage-gated Ca\(^{2+}\)-channels are localized to active zones by binding to a multi-domain protein complex composed of RIMs and RIM-binding proteins (RBPs); this complex also directly interacts with synaptic vesicles and the exocytotic machinery and forms the core of the active zone (Deng et al, 2011; Han et al, 2011; Kaeser et al, 2011; Liu et al, 2011; Davydova et al, 2014; Acuna et al, 2015, 2016; Muller et al, 2015; Grauel et al, 2016; reviewed in Sudhof, 2012). In mammals, RBPs are encoded by three genes: RBP1 and RBP2 are differentially expressed throughout the brain, while RBP3 is primarily expressed outside the brain (Wang et al, 2000; Mittelstaedt & Schoch, 2007). RIM-binding proteins are composed of three SH3-domains and three fibronectin-type III domains (FN3-domains) that are flanked by variable interspersed sequences. The SH3-domains of RBPs directly bind to proline-rich sequences in RIMs and in L-, and P/Q- and N-type voltage-gated Ca\(^{2+}\)-channels (Wang et al, 2000; Hibino et al, 2002; Kaeser et al, 2011). Via these interactions, RBPs contribute to the presynaptic recruitment of voltage-gated Ca\(^{2+}\)-channels at Drosophila neuromuscular junctions (Liu et al, 2011), of N- and P/Q-type Ca\(^{2+}\)-channels at standard chemical synapses (Acuna et al, 2015, 2016; Grauel et al, 2016),
and of L-type Ca\(^{2+}\)-channels at ribbon synapses (Krinner et al., 2017; Luo et al., 2017). RIMs interact with Ca\(^{2+}\)-channels both indirectly via binding to RBPs and directly via their PDZ-domain that bind to the C-terminus of N- and P/Q-type Ca\(^{2+}\)-channels (Kaeser et al., 2011). In addition, RIMs interact with Rab3-proteins on synaptic vesicles, with Bassoon in the cytomatrix, and with Munc13 in the vesicle fusion machinery (Wang et al., 1997, 2000; Betz et al., 2001; Schoch et al., 2002; Deng et al., 2011; Kaeser et al., 2011; Davydova et al., 2014). Thus, RIMs nucleate a macromolecular complex that includes Ca\(^{2+}\)-channels, synaptic vesicles, and the release machinery in addition to RBPs (Sudhof, 2012).

At standard chemical mammalian synapses in which N- and P/Q-type Ca\(^{2+}\)-channels mediate presynaptic Ca\(^{2+}\)-influx, RIMs are the primary drivers for synaptic recruitment of Ca\(^{2+}\)-channels (Kaeser et al., 2011), whereas RBPs support RIMs in organizing nano-scale coupling of Ca\(^{2+}\)-influx to release (Acuna et al., 2015; Grauel et al., 2016). As a result, at these synapses (as measured at the calyx of Held synapse), deletion of RIMs massively decreases presynaptic Ca\(^{2+}\)-currents (Han et al., 2011; Kaeser et al., 2011), deletion of RBPs decreases the fidelity of neurotransmitter release without changing overall Ca\(^{2+}\)-currents (Acuna et al., 2015), and deletion of both RIMs and RBPs nearly abolishes presynaptic Ca\(^{2+}\)-currents (Acuna et al., 2016). At ribbon synapses that use L-type voltage-gated Ca\(^{2+}\)-channels, however, RBPs are primarily responsible for recruiting voltage-gated Ca\(^{2+}\)-channels because RBPs but not RIMs bind to L-type Ca\(^{2+}\)-channels (Hibino et al., 2002; Kaeser et al., 2011). Here, deletion of RBPs has a much more pronounced effect on release at ribbon synapses than at standard chemical synapses (Luo et al., 2017).

BK-channels are formed by tetramers of BK\(\alpha\)-subunits that have seven transmembrane regions and a long cytoplasmic sequence containing two regulator-of-K+-conductance (RCK) domains (the RCK1- and RCK2-domains; Atkinson et al., 1991; Adelman et al., 1992; Butler et al., 1993; Lee & Cui, 2010). The RCK-domains regulate gating of BK-channels via Ca\(^{2+}\)-binding, phosphorylation, and interactions with other proteins (Xia et al., 2002; Yusifov et al., 2008; Yuan et al., 2010). BK\(\alpha\) proteins associate with cytoplasmic auxiliary BK\(\beta\)- and BK\(\gamma\)-subunits that affect the gating properties of BK-channels and may contribute to their subcellular targeting (Meera et al., 1996; Tseng-Crank et al., 1996; Wallner et al., 1996; Orío et al., 2002). BK-channels are ubiquitously expressed in the brain (Sausbier et al., 2006) and enriched in presynaptic terminals (Robitaille et al., 1993a; Knaus et al., 1996; Zhou et al., 1999; Hu et al., 2001; Ishikawa et al., 2003; Misonou et al., 2006; Nakamura & Takahashi, 2007), where they form a molecular complex with Ca\(^{2+}\)-channels, and are thus optimally located for coupling Ca\(^{2+}\)-influx to the Ca\(^{2+}\) - regulation of the duration of an AP (Roberts, 1993; Robitaille et al., 1993b; Berkefeld et al., 2006; Loane et al., 2007; Wang, 2008).

High Ca\(^{2+}\)-concentrations (~10 \(\mu\)M) are required for activation of BK-channels, implying that BK-channels are physiologically activated only in close proximity to voltage-gated Ca\(^{2+}\)-channels and preferentially during AP trains (Berkefeld & Fakler, 2013). As a result, it has been suggested that nano-domain coupling of BK-channels to voltage-gated Ca\(^{2+}\)-channels is necessary to trigger opening of BK-channels during APs (Berkefeld et al., 2006; Fakler & Adelman, 2008). Selective block of BK-channels with iberiotoxin (IbTX) or paxilline increased the probability of glutamate release in CA3-CA3 synapses (Raffaelli et al., 2004) and regulated neurotransmitter release in A17 amacrine cells in the retina (Grimes et al., 2009). Moreover, at the calyx of Held synapse (Ishikawa et al., 2003; Nakamura & Takahashi, 2007), in hippocampal mossy fibre synapses (Alle et al., 2011), and at CA3-CA1 Schaffer collateral synapses (Hu et al., 2001), a presynaptic function of BK-channels in regulating neurotransmitter release was uncovered when voltage-gated K\(^{-}\)-channels were blocked with 4-aminopyridine. Overall, the current evidence suggests that BK-channels are an important mechanism of regulating neurotransmitter release especially during AP trains, but the molecular interactions that recruit BK-channels to presynaptic terminals are unclear.

Here, we identified BK-channels in an unbiased yeast two-hybrid screen as novel interaction partner of RBPs. The binding of RBPs to BK-channels required the FN3-domains of RBPs and the RCK-domains of BK-channels. These findings led us to propose that RBPs function as scaffolds at the release site, and that by simultaneously binding to BK-channels (through the FN3-domains) and voltage-gated Ca\(^{2+}\)-channels and RIMs (through their SH3-domains), RBPs contribute to the nano-domain coupling of Ca\(^{2+}\)-influx to Ca\(^{2+}\)-triggered synaptic vesicle exocytosis. Our results thus describe a function for the conserved FN3-domains of RBPs and suggest a mechanism by which BK-channels are recruited to voltage-gated Ca\(^{2+}\)-channels at the release site in the active zone.

Results

RBPs bind to BK\(\alpha\)-channels

RB1 and RB2 are large multi-domain proteins that contain one N-terminal SH3-domain, three central FN3-domains, and two C-terminal SH3-domains (Fig 1A; Mittelstaedt & Schoch, 2007; Wang et al., 2000). To identify new binding partners for RBPs, we performed unbiased large-scale yeast two-hybrid screens using three fragments of rat RB2 as baits: a fragment (residues 247–859) containing the FN3-domains and the linker region; a fragment (residues 1–859) containing the N-terminal SH3-domains, the FN3-domains, and the long central linker sequence; and a fragment (residues 247–1,068) containing the FN3-domains, linker region, and C-terminal SH3-domains (Fig 1A, Appendix Fig S1A). We isolated and re-confirmed a total of 71 prey clones (Appendix Fig S1B, Tables S1 and S2). Four of the prey clones that were isolated with the RBP2-247-859 bait encoded the cytoplasmic RCK2-domain of BK\(\alpha\)-channels (Fig 1A, Appendix Table S1). Moreover, as previously described (Hibino et al., 2002; Kaeser et al., 2011), multiple clones representing the L-type Ca\(^{2+}\)-channel \(\alpha\)-subunit (Cacna1b) were recovered with baits containing the SH3-domains, validating the overall screen (Appendix Fig S1B and Table S1). In addition, several other preys of potential interest were recovered (Appendix Table S1), but were not pursued further because of the potential importance of a direct interaction of RBPs with BK\(\alpha\)-channels.

To independently assess the interaction between RBP2 and BK\(\alpha\)-channels, we performed co-immunoprecipitation experiments with proteins expressed in HEK293T cells (Fig 1B and C). We co-expressed myc-tagged full-length RB2 with the YFP-tagged BK\(\alpha\) RCK2-domain that we had isolated in the yeast two-hybrid screen. As negative and positive controls, respectively, we used YFP alone
or YFP-tagged full-length RIM1α (Fig 1B). We then confirmed by co-immunoprecipitations with GFP antibodies (which recognize YFP) or with RBP2 antibodies that myc-tagged RBP2 can bind to both RIM and BKα-channels, whereas YFP alone could not bind to either molecule (Fig 1C).

Next, we examined the interaction between RBP2 and BKα-channels by imaging their localizations in transfected HEK293T cells (Fig 1D and E). We expressed the YFP-tagged RCK2-domain of BKα alone or together with myc-tagged full-length RBP2 in HEK293T cells, immunostained the cells with myc-specific antibodies, and then quantified fluorescence signals relative to the nucleus, using DAPI as a marker of nuclei. The YFP-tagged RCK2-domain of BKα and YFP, when expressed alone, was localized to all cellular compartments (as is often observed for overexpressed proteins), whereas RBP2 expressed alone quantitatively localized to the plasma membrane (Fig 1D). Co-expression of RBP2 with the RCK2-domain, however, caused the translocation of the RCK2-domain to the membrane, confirming binding (Fig 1D). Co-expression of RBP2 with YFP, which is also localized in all cellular compartments, had no effect. Quantifications using correlation analyses confirmed that YFP-tagged RCK2-domain and YFP co-localized with the nuclear DAPI-stain in the absence of RBP2, and that YFP-tagged RCK2-domain but not YFP were recruited to the plasma membrane in the presence of RBP2, thus revealing their interaction in the context of the expressing cell (Fig 1E).
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The experiments up to now establish an in vitro interaction of RBP2 with the RCK2-domain of BKα. To test whether endogenous RBP2 and BKα-channels form a complex in vivo, we immunoprecipitated RBP2 or RIMs from brain homogenates, and immunoblotted the immunoprecipitated proteins for BKα, RBP2, and RIMs (Fig 1F and G, Appendix Fig S2). The challenge inherent to these experiments is that RIMs and RBPs are part of the active zone protein complex that is largely detergent-insoluble, prompting us to explore a variety of detergents for the immunoprecipitations, including low concentrations of SDS. We found that in these immunoprecipitations, BKα was co-enriched with both RBP2 and RIMs, whereas syntaxin-1 (used as a negative control) was not (Fig 1G, Appendix Fig S2B). These results are consistent with the presence of a complex composed of endogenous BKα, RBP2, and RIMs. Importantly, no signal for BKα was detected when immunoprecipitations were performed with samples from constitutive RBP1,2 double KO mice, confirming the specificity of the interaction (Appendix Fig S2C).

RBP sequences containing FN3-domains bind to both RCK-domains of BKα-channels

To map the domains of RBP2 that bind to BKα-channels, we co-expressed different fragments of myc-tagged RBP2 with YFP-tagged BKα-channels in HEK293T cells and tested binding by immunoprecipitation and imaging (Fig 2A). As assayed by co-immunoprecipitations, full-length RBP2 (aa 1–1,068) as expected bound robustly to BKα-protein (Fig 2B). A truncated version of RBP2 containing the FN3-domains and adjacent linker sequences (aa 247–859) bound reproducibly but weakly to BKα-channels, whereas a RBP2 construct containing a deletion of the FN3-domains and linker sequences (Δ247–859) failed to bind (Fig 2B). The same binding relationships were observed when measured by imaging the RBP2-dependent translocation of the YFP-tagged RCK2-domain of BKα out of the nucleus (Fig 2C and D). These results confirm the yeast two-hybrid experiment and suggest that the FN3-domains of RBPs are required for RB binding to BKα-channels.

We next examined the region of BKα-channels that mediates BKα-binding to RBPs. The intracellular sequences of BKα-channels are composed of tandem RCK-domains, the RCK1-domain (aa 391–692) and RCK2-domain (aa 761–1,178), that are homologous to each other (Appendix Fig S3A and B). We therefore tested if the interaction of RBP2 with the RCK2-domain, as shown above, is
To confirm the presence of a direct interaction between RBPs and BKα, we produced recombinant GST-fused fragments of RBP1 (aa 771–1,086) and RBP2 (aa 298–604) containing only the FN3 domains (Appendix Fig S4A and B). We found that these fragments were sufficient to successfully pull-down YFP-BKα (aa761–1,178) containing the RCK2 domain from HEK293T cell lysates (Appendix Fig S4C and D). The presence of this interaction was further strengthened by co-immunoprecipitation experiments from transfected HEK293T cells co-expressing full-length RBP2 with Ca2+-channels (Cav2.2) and BKα-transfected HEK293T cells expressing either myc-tagged RBP2 alone or together with YFP or the indicated YFP-tagged BKα RCK-domains. Input fractions (ln; 1% of total) or immunoprecipitates (IP) were analysed by immunoblotting with antibodies to myc (red) or CFP (green, recognizes YFP). For more experiments, see Appendix Fig S3.

C Representative images (top; scale bar, 5 µm) or line scans (bottom) of HEK293T cells expressing myc-tagged RCK1 (top) and RCK2 domains (bottom) and nuclear DAPI (blue). Note that the BKα-RCCK domains (green YFP fluorescence) are soluble when expressed alone, but are recruited to the cell membrane when co-expressed with RBP2. See also Fig 1D (experiments shown in Figs 1D and 3 were done in the same batch).

D Cumulative plots and bar graphs of the correlation coefficients between YFP-tagged BKα domains (top) and of the two BKα isoforms expressed in the brain (RBP1 and RBP2) can bind directly and simultaneously to Ca2+-channels via their SH3-domains and to BKα-channels via their FN3-domains.
blue trace) caused a shift in the voltage dependence of BKα-channels, whereas co-expression of a deleted version of RBP2 in which the FN3-domains and the linker region were removed had no effect (Fig 4B, green trace). Quantification of the conductance–voltage (G–V) relationships for BKα-mediated currents under the various conditions demonstrated that the observed changes were significant (Fig 4C). Specifically, RBP2 variants that bound to BKα-protein in vitro shifted the voltage dependence of the BKα-current activation by 15–20 mV without affecting the slope of the voltage dependence, whereas the RBP2 variant that did not bind to BKα-protein in vitro had no effect (Fig 4D). Therefore, these experiments confirm with a functional assay that the interaction between BKα-channels and RBPs requires the FN3-domains of RBPs.

**Deletion of RBPs decreases presynaptic BK-currents**

To test whether the interaction between RBPs and BKα-channels is physiologically significant, we turned to the calyx of Held, a giant synaptic terminal in the auditory system that allows direct access to the presynaptic terminal (Schneggenburger & Forsythe, 2006). Because presynaptic terminals of the calyx of Held synapse can be patched, this synapse enables direct measurements of various presynaptic ion currents, revealing among others prominent presynaptic BK-currents (Ishikawa et al., 2003; Nakamura & Takahashi, 2007). We deleted RBPs in presynaptic calyx of Held terminals by crossing RBP1,2 conditional double KO mice with Krox-20 Cre-driver mice that express Cre-recombinase in presynaptic neurons of the calyx of Held synapse (Voiculescu et al., 2000; Acuna et al., 2015; Fig 5A). Since only RBP1 and RBP2 but not RBP3 are detectable in the brain, the double deletion of RBP1 and RBP2 completely removes RBPs from calyx synapses. In the following, we analysed littermate RBP1,2 conditional double KO mice either lacking (referred to as “WT”) or expressing Cre-recombinase (referred to as “DKO”).

We prepared acute brainstem slices from P10-12 WT and DKO mice, and patched presynaptic calyx of Held terminals (Fig 5B). To monitor BK-currents, we blocked voltage-gated Na⁺- and
K⁺-channels with tetrodotoxin (TTX) and 4-aminopyridine (4-AP), respectively, and measured currents in response to progressive step depolarizations with 10 mV intervals (Fig 5C; Nakamura & Takahashi, 2007). Under these conditions, high-voltage activated Ca²⁺-currents manifested as inward currents with a rapid onset, and 4-AP-insensitive K⁺-currents manifested as outward currents with a delayed onset (Fig 5C). 4-AP-insensitive K⁺-currents are comprised of BK-currents and of other slowly activating K⁺-currents (Nakamura & Takahashi, 2007). Subsequent bath application of iberiotoxin (IbTx, 0.2 μM), a specific inhibitor of BK-channels (Galvez et al, 1990), suppressed the majority of the 4-AP-insensitive K⁺-currents that correspond to BK-currents (Fig 5C, compare traces (i) and (ii); quantified in Fig 5D). Treatment with IbTx retained a low-amplitude 4-AP-insensitive K⁺-current that is probably due to TEA-sensitive K⁺-conductances, and had no effect on Ca²⁺-currents (Fig 5C and E–G). Overall, these data indicate that most of the 4-AP-insensitive K⁺-currents in the presynaptic calyx terminal are due to BK-currents.

Using this protocol, we first analysed 4-AP-insensitive K⁺-currents in WT and DKO calyx of Held terminals (Fig 6). We found that the density of the 4-AP-insensitive K⁺-current was significantly decreased in RBP-deficient terminals depolarized to 10 mV, and that the time constant of the fast component of the current was increased more than fivefold (Fig 6A–C). The remaining 4-AP-insensitive K⁺-current after addition of IbTx, however, was not impaired in RBP-deficient presynaptic terminals (Fig 6D–F).

We then calculated the BK-current for each terminal by subtracting the IbTx-insensitive K⁺-current from the total 4-AP-insensitive K⁺-current (Fig 6G–M). Strikingly, the BK-current density was significantly reduced in RBP-deficient presynaptic terminals compared to WT terminals from littermate controls (Fig 6G–I). Moreover, we found that deletion of RBP1,2 rendered no change in the voltage dependence of the BK-current activation (Fig 6J), but induced a large increase in the rise times of BK-currents (i.e., their activation kinetics, Fig 6L) and caused a delay in BK-current onset as reflected in an increased latency (Fig 6M). Note that the nerve terminal capacitance was unaltered, suggesting that these changes are not due to a difference in the architecture of the terminal (Fig 6N).

Viewed together, these results suggest that deletion of RBP1 from the presynaptic calyx terminal causes a decrease in the concentration of BK-channels. However, the gating of BK-channels is dependent on Ca²⁺, which in turn depends on the activity of voltage-gated Ca²⁺-channels in the presynaptic calyx of Held terminals (Berkefeld & Fakler, 2013). Therefore, it is possible that the decrease in BK-currents observed in the RBP-deficient terminals is not due to an impairment in BK-channels, but caused by a decrease in voltage-gated Ca²⁺-influx. We tested this possibility directly and found that, consistent with previous results (Acuna et al, 2015), the Ca²⁺-current density and rise times (as a measure of activation kinetics) were not significantly different between WT and RBP1,2 DKO calyx of Held terminals (Appendix Fig S6). This result shows that the phenotype of reduced BK-currents we observed in RBP-deficient presynaptic terminals is not due to a reduction in the presynaptic Ca²⁺-influx.

Since the RBP DKO causes a loss of tight coupling of Ca²⁺-channels to the release machinery even though it does not produce a loss of Ca²⁺-channels as such (Acuna et al, 2015), it is conceivable that the decrease in BK-currents in RBP1,2 DKO terminals is an indirect effect of this function of RBPs. To test this remote possibility, we measured BK-currents in control and RBP1,2 DKO terminals in the

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**Figure 5. Experimental approach to recording presynaptic K⁺-currents mediated by BK-channels at the calyx of Held synapse.**

A Breeding strategy to obtain mice with a conditional deletion of RBP1 and RBP2 at the calyx of Held synapse using Krox-20 Cre-driver mice (Voiculescu et al, 2000). Matings of RBP1,2 conditional DKO (cDKO) mice containing or lacking Krox-20 Cre alleles produced littermate DKO and WT mice.

B Schematic of whole-cell patch-clamp recordings from presynaptic calyx terminals.

C Presynaptic currents recorded in response to stepwise 50-ms depolarizations of presynaptic terminals from –50 to +50 mV (10 mV increments) in the presence of tetrodotoxin (TTX, 1 μM) and 4-aminopyridine (4-AP, 25 mM), without (i) and with (ii) addition of 0.2 μM iberiotoxin (IbTx). BK-currents are calculated by subtracting (ii) from (i) as IbTx-sensitive K⁺-currents. Note that in these recordings, inward Ca²⁺-currents (ICa²⁺) and outward K⁺-currents (IK⁺) are monitored, of which only BK-currents are IbTx-sensitive.

D Time course of 4-AP-insensitive K⁺-currents as recorded in a representative experiment as a function of the application of iberiotoxin (IbTx, 0.2 μM).

E Same as (D), but for Ca²⁺-currents to illustrate that IbTx has no effect on Ca²⁺-channels.

F Summary plot of the Ca²⁺-current amplitude before and after treatment with IbTx. Single cell values are represented with dotted lines, while the average is shown by full line.

G Same as (F) but for Ca²⁺-current rise times.

Data information: Data are means ± SEM. Significance was calculated using Student’s t-test, n = 19, P < 0.05.
Figure 6. Deletion of RBPs decreases and decelerates presynaptic BK-currents. All experiments were performed by patch-clamp recordings from presynaptic calyx of Held terminals from littermate RBP1.2 conditional DKO mice that lacked (WT) or expressed Krox-20 Cre (DKO) at P10-12. Low concentration of EGTA (0.2 mM) in the internal solution was used for these experiments.

A Sample traces of Ca²⁺-currents and 4-AP-insensitive K⁺-currents induced by step depolarizations (from −50 mV to +10 mV in 10 mV increments, monitored in 2.5 mM 4-AP). Arrows indicate 30-ms time points, used for analysis in (B). The yellow dotted line superimposed to the trace represents fitting of the curve with a double exponential function, for analysis in (C).

B Summary plot of the mean density of 4-AP-insensitive K⁺-currents as a function of voltage measured at 30 ms after depolarization as described in (A).

C Summary graphs of the fast (t_fast, left) and slow time constants (t_slow, right) of 4-AP-insensitive K⁺-currents monitored at +10 mV depolarization. Currents were fitted with a double exponential equation to calculate time constants.

D–F Same as (A–C), but for currents measured after additional application of iberiotoxin (IbTx, 0.2 µM). Current density was measured at the peak.

G Sample traces of BK-currents calculated by subtracting currents produced by depolarizations in the presence of both iberiotoxin and 4-AP from currents produced in the presence of only 4-AP for a given calyx terminal.

H Summary plots of the mean density of BK-currents in presynaptic calyx of Held terminals; BK-current density is plotted as a function of the depolarization voltage, measured at the peak amplitude.

I Cumulative distribution plot (left) and mean value at 10 mV (right) of the BK-current density as described in (H).

J Same as (H), but normalized to the peak current to demonstrate that the deletion of RBPs has no effect on the voltage dependence of BK-channel activation.

K Sample traces for BK-currents monitored at −50 mV and +10 mV to illustrate measurements of 20–80% rise times and latencies of BK-currents.

L Summary plot of mean 20–80% rise times of BK-channels in presynaptic calyx of Held terminals monitored at −30 mV, 0 mV, and +10 mV.

M Summary plot of mean latencies of BK-channels analysed as in (L).

N Summary graph of the capacitance of calyx of Held terminals.

Data information: Data are means ± SEM. Significance was calculated using Student’s t-test (panels C, F, I, and N) or two-way ANOVA for repetitive measurements followed by Bonferroni post hoc test (panels B, E, H, J, L, and M). n = 9 terminals/5 mice [WT] or 10 terminals/6 mice [DKO]. *P < 0.05, **P < 0.01, ***P < 0.001. For Ca²⁺-current quantifications, see Appendix Fig S6.
The presence of high concentration of the slow Ca$^{2+}$-chelator EGTA (10 mM) in the patch pipette. Moreover, with this experiment, we further explored whether the changes in gating kinetics and latency of BK-currents observed in RBP1,2 DKO calyces were due to alterations in the coupling within voltage-gated Ca$^{2+}$-channels and BK-channels. EGTA is a Ca$^{2+}$-chelator with slow Ca$^{2+}$-binding rate constant that has been used to selectively interfere with processes that require Ca$^{2+}$-microdomains, and therefore are loosely coupled to Ca$^{2+}$-channels (at distances higher than 50 nm from Ca$^{2+}$-channels). However, processes that require Ca$^{2+}$-nano-domains and are placed at a distance within 20–50 nm from the Ca$^{2+}$ source are not sensitive to EGTA, but can be blocked by fast Ca$^{2+}$-chelators, such as BAPTA (Fakler & Adelman, 2008; Acuna et al., 2015; Luo et al., 2017). We found that at the calyx of Held, application of high concentration of EGTA (10 mM) completely blocks BK-currents in RBP1,2 DKO mice, confirming that removal of RBPs affects the coupling of these channels to voltage-gated Ca$^{2+}$-channels (Fig 7A, B and D). Somewhat surprisingly, BK-currents in WT mice were also decreased by EGTA, suggesting that at the age tested (P10-12), a fraction of BK-channels is only loosely coupled to Ca$^{2+}$-channels. As expected from previous results, no differences in the capacitance (Fig 7C) and the Ca$^{2+}$-currents (Fig 7E) were observed in WT and RBP1,2 DKO mice, suggesting that the phenotype was specific for BK-channels.

These experiments let us to conclude that removal of RBPs contributes the correct localization of BK-channels at the presynaptic terminal, controlling their coupling to voltage-gated calcium channels.

**RBPs do not affect expression, but impair presynaptic localization of Bkx-channel protein**

The reduction in BK-currents we observed upon deletion of RBPs in the calyx of Held (Fig 6G–I) could potentially be explained by changes in the expression levels or by a loss of presynaptic BK-channels, i.e., a mislocalization of Bkx-channels away from the active zone.

In order to examine whether removal of RBPs interferes with the expression of BK-channels, we performed immunoblotting analyses. We prepared total brain lysates from WT and constitutive RBP1,2 DKO mice and analysed the protein levels of the different subunits of BK-channels (Fig 8A). BK-channels are formed by tetramerization of pore-forming Bkx-subunits that recruit auxiliary BKβ-subunits (β1–β4; Atkinson et al., 1991; Adelman et al., 1992; Butler et al., 1993; Tseng-Crank et al., 1996; Behrens et al., 2000; Brenner et al., 2000; Uebele et al., 2000; Poulsen et al., 2009). In the brain, BKβ2 and BKβ4 subunits are most abundant, while BKβ3 is undetectable (Poulsen et al., 2009). We found that the RBP1,2 DKO had no effect on the levels of Bkx, BKβ1, BKβ2, and BKβ4 as measured by quantitative immunoblotting of whole-brain homogenates (Fig 8A), suggesting that RBPs are not regulating the expression levels of BK-channels or their auxiliary subunits. Similarly, the ventral cochlear nucleus (VCN), which contains the cell bodies of the neurons forming calyx synapses, exhibited no significant differences in the level of Bkx and BKβ subunits (Appendix Fig S7B). These data support the idea that genetic removal of RBPs does not affect the overall expression of BK channel subunits. Interestingly, when we performed immunoblotting in the micro-dissected medial nucleus of the trapezoid body (MNTB) containing calyx synapses, we found a trend towards reduced Bkx levels and a significant decrease in the presence of high concentrations of the slow Ca$^{2+}$-chelator EGTA (10 mM) in the patch pipette.
of Held synapse, BKα-channels partially co-localized with vGlut1, confirming their presynaptic localization (Fig 8B). In RBP-deficient DKO mice, the presynaptic area containing BKα-channels at the calyx of Held synapse was significantly decreased (~45%; Fig 8C and D). These results confirm that RBPs are necessary to ensure the correct localization of BK-channels at the presynaptic terminal.

**Discussion**

Presynaptic BK-channels fine-tune the temporal pattern of APs, thereby controlling the dynamics of neurotransmitter release from nerve terminals (Storm, 1987; Hu et al, 2001; Fakler & Adelman, 2008). In the brain, BK-channels are largely present in a complex with voltage-gated Ca2+-channels that are also partly presynaptic, thus enabling tight coupling of Ca2+-influx to both Ca2+-dependent exocytosis and Ca2+-dependent BK-channel activation (Roberts, 1993; Robitaille et al, 1993a; Knaus et al, 1996; Zhou et al, 1999; Hu et al., 2001; Ishikawa et al., 2003; Misonou et al, 2006; Nakamura & Takahashi, 2007). How BK-channels are recruited into a complex with Ca2+-channels to presynaptic release sites, however, was unknown. Presynaptic release sites are formed by active zones, which are composed of large complexes of multi-domain proteins (Sudhof, 2012). RBPs are a component of active zones that, among others, control the fidelity of neurotransmitter release by enabling tight coupling of voltage-gated Ca2+-influx to Ca2+-triggered synaptic vesicle exocytosis (Acuna et al, 2015, 2016; Grael et al., 2016; Krinner et al, 2017; Luo et al, 2017). In addition, RBPs redundantly support the essential functions of RIMs, their eponymous binding partners, in the tethering and priming of synaptic vesicles prior to Ca2+-triggered exocytosis (Acuna et al, 2016). RBPs contain two major domains, SH3- and FN3-domains, consistent with a scaffold that operates by simultaneous binding of

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**Figure 8. Deletion of RBPs causes partial loss of BKα-channels from presynaptic terminals without altering BKβ expression levels.**

A Immunoblot analysis of the levels of BKα, BKβ1-, BKβ2-, and BKβ4-subunits of BK-channels in total brain homogenates from littermate WT and constitutive RBP1,2 DKO mice (left, representative blots; right, summary graph of levels normalized to tubulin (Tuji) and to WT levels; n = 6 WT and DKO mice). B Illustration of the strategy for quantitative immunofluorescence analyses of the presynaptic localization of vGlut1 (green) and BKα (magenta) in the calyx of Held synapse in control mice (top, representative images [dashed arrows = position of optical sections for quantitative analyses], bottom, normalized profiles of the vGlut1 and BKα staining intensity in the calyx of Held terminals as revealed by optical sections). C Immunofluorescence measurements of the BKα levels in presynaptic terminals of calyx of Held synapses in littermate RBP1,2 conditional DKO mice that lacked (WT) or expressed Krox-20 Cre (DKO). Brainstem sections of the medial nucleus of the trapezoid body (MNTB) containing calyx synapses were stained for vGlut1 (green) and BKα-channels (magenta), and BKα levels were determined using optical sections as illustrated in (B). BKα staining area was normalized for that of vGlut1 as an internal standard in the same sections to control for differences in staining efficiencies and section plane. D Summary graph of the mean presynaptic BKα staining area normalized to the vGlut1 staining area as described in (C) (n = 8 WT and 9 DKO mice, averaging multiple images from each mouse as described in the Appendix). Data information: Data in (A and D) are means ± SEM. Significance was calculated using Student’s t-test (*P < 0.05). See Appendix Fig S7 for more details.

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level of BKβ4, while the other subunits were unchanged (Appendix Fig S7C). In the MNTB, we also observed no changes in the level of synaptotagmin-2 (Syt2) and parvalbumin (PV), used as markers for the calyx of Held synapses.

The immunoblotting data suggest that deletion of RBPs may lead to a specific loss of BK-channels from calyx of Held terminals. To further explore this possibility, we measured presynaptic BKα-protein levels by immunofluorescence, using the presynaptic marker vGlut1 as an internal control. We found that the wild-type calyx

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RIM-BPs recruit presynaptic BK-channels

...multiple interactors. Multiple binding partners for RBPs are known, including RIMs, voltage-gated Ca\(^{2+}\)-channels, and bassoon (Wang et al., 2000; Hibino et al., 2002; Kaeser et al., 2011; Davydova et al., 2014). All of these proteins bind to the SH3-domains of RBPs, however, and no ligands for the FN3-domains of RBPs were known. To search for potential interactors for the RBP FN3-domains, we here performed an unbiased yeast two-hybrid screen that identified BK\(_{\alpha}\)-channels as candidate binding partners for RBPs (Fig 1, Appendix Fig S1, Table S1 and S2). We found that BK\(_{\alpha}\)-channels physically interact with RBPs (Figs 2–4) and that deletion of RBPs leads to a decrease in BK\(_{\alpha}\)-currents in presynaptic terminals (Figs 5–7). Thus, we propose that BK\(_{\alpha}\)-channels are recruited as a component of the macromolecular active zone complex by direct binding to RBPs, which in turn bind to RIMs and to voltage-gated Ca\(^{2+}\)-channels (Fig 9).

The evidence in support of this proposal can be summarized as follows: First, BK\(_{\alpha}\)-channels were found to directly bind to RBPs by four different protein interaction measurements, namely yeast two-hybrid assays (Fig 1A, Appendix Fig S1), co-immunoprecipitations (using both expression systems and native brain homogenates, Fig 1, Appendix Fig S2), imaging of co-expressed proteins (Fig 1D and E), and pull-down experiments (Appendix Fig S4). Each individual measurement may be considered inconclusive owing to the limitations of the underlying methods, but in combination these measurements provide strong evidence for an interaction. Second, mapping of binding with defined BK\(_{\alpha}\)- and RBP2-domains revealed that the RCK-domains of BK\(_{\alpha}\) bound specifically to RBP2 sequences containing its FN3-domains, whereas the SH3-domains were dispensable (Figs 2 and 3). This conclusion was further supported by pull-down experiments using recombinant FN3-domains of both RBP1 and RBP2 (Appendix Fig S4). Third, co-expression of RBP2 sequences that contained the BK\(_{\alpha}\)-binding site but not co-expression of RBP2 sequences that lacked the BK\(_{\alpha}\)-binding impacted the function of BK\(_{\alpha}\)-channels reconstituted in HEK293T cells (Fig 4). We interpret these results purely as evidence for a specific interaction, but not as indicative of a particular function of this interaction since the conditions of the HEK293T cell expression of the BK\(_{\alpha}\)-channel are too artificial for the experiment to be functionally meaningful. Fourth, deletion of RBPs from the calyx of Held synapse produced a significant reduction in both BK-current amplitude and kinetics, demonstrating that endogenous RBPs are essential for full recruitment of BK-channels to presynaptic terminals (Figs 6 and 7). Fifth, immunofluorescence showed that deletion of RBPs from the calyx of Held synapse caused a physical loss of BK\(_{\alpha}\)-channel protein from the presynaptic nerve terminal, which was confirmed by immunoblotting analyses of micro-dissected MNTB samples (Fig 8, Appendix Fig S7).

Although we believe that these data strongly support the proposal that RBPs serve to recruit BK-channels to presynaptic active zones in proximity to Ca\(^{2+}\)-channels (Fig 9), several questions may be raised about the interpretation of these data. Is it possible that the loss of presynaptic BK\(_{\alpha}\)-channels caused by the RBP1,2 DKO in the calyx of Held synapse is an indirect consequence of the mislocalization of voltage-gated Ca\(^{2+}\)-channels (Acuna et al., 2015; Grauel et al., 2016)? This possibility seems highly unlikely because deletion of RBPs in DKO synapses by itself does not cause a change in total presynaptic Ca\(^{2+}\)-currents, but only a modest uncoupling of release from APs during high-frequency stimulus trains (Appendix Fig S5; Acuna et al., 2015). Thus, this possibility would not explain the physical loss of BK\(_{\alpha}\)-channels from presynaptic terminals. Another question is whether our results obtained at the unusually large calyx of Held synapse are generally applicable to all synapses. Although we cannot rule out that other mechanisms guide recruitment of BK-channels to presynaptic terminals in other synapses, this possibility again appears unlikely given that the function of RBPs operates at all synapses (Acuna et al., 2015; Grauel et al., 2016; Kinner et al., 2017; Luo et al., 2017), and that in total brain, RBPs appear to be present in a complex both with BK\(_{\alpha}\)-channels and with Ca\(^{2+}\)-channels (Fig 1, Appendix Fig S2). Yet another question regards the relation of RBP binding to BK\(_{\alpha}\)-channels to the interaction of BK\(_{\alpha}\)-channels with BK\(_{\beta}\)-subunits. Although in vivo BK\(_{\alpha}\)-channels operate likely as a multi-subunit complex containing BK\(_{\alpha}\) and auxiliary BK\(_{\beta}\)-subunits, our in vitro studies were performed in the absence of BK\(_{\beta}\)-subunits. Thus, we do not know whether BK\(_{\beta}\)-subunits affect RBP binding to BK\(_{\alpha}\)-channels either positively or negatively, a subject that requires extensive additional studies. However, the RBP1,2 DKO phenotype reflects a change in the entire BK-complex, not just BK\(_{\alpha}\)-channels alone, suggesting that RBP clearly is essential for the proper localization of the entire complex.
In summary, we here propose that BKα-channels are recruited into the active zone complex by a direct interaction with RBPα, which also bind to voltage-gated Ca\(^{2+}\)-channels and to RIMs as central active zone components (Fig 9). As a result, the active zone protein complex is much more extensive as previously envisioned and includes not only Ca\(^{2+}\)-channels and cell adhesion molecules that are coupled to the synaptic vesicle fusion machinery (Sudhof, 2012), but also with BKα-channels proteins that limit the extent of membrane depolarization during an AP and thereby the amount of neurotransmitter release. This molecular architecture of the active zone complex is likely to have significant implications for the regulation of release under diverse physiological and pathological conditions.

Materials and Methods

All experiments were performed on anonymized coded samples. For detailed experimental procedures, see Appendix. All animal procedures conformed to National Institutes of Health Guidelines for the Care and Use of Laboratory Animals and were approved by the Stanford University Administrative Panel on Laboratory Animal Care.

Yeast two-hybrid experiments

Yeast two-hybrid screens of a rat cDNA library were performed as described (Wang et al, 1997; Kaeser et al, 2011; see Appendix for details). Three bait vectors expressing: (i) truncated RBP2 containing residues 247–859, encoding the N-terminal SH-domain and the FN3-domains of RBP2; (ii) truncated RBP2 containing residues 1–859, encoding the N-terminal SH-domain and the FN3-domains of RBP2; and (iii) truncated RBP2 containing residues 247–1,068, encoding the FN3-domains and the C-terminal SH3-domains. All preys were validated by re-assessing their interaction in yeast two-hybrid assays.

Immunoprecipitations and immunoblotting experiments

Protein interactions were validated by immunoprecipitations of proteins expressed in HEK293T cells. Proteins were co-expressed in HEK293T cells in various combinations as specified in the figures and figure legends by co-transfection. Immunoprecipitation and immunoblotting experiments were performed using standard procedures (see Appendix). For analyses of endogenous brain proteins, experiments used brain homogenates solubilized in Triton X-100 from control and RBP1/RBP2 constitutive DKO mice (Acuna et al, 2015).

Analyses of the RBP interaction with BKα using imaging of transfected HEK293T cells

HEK293T cells were co-transfected with constructs encoding various RBP2, BKα, and YFP proteins as described in the figures and figure legends, and analysed by immunocytochemistry and imaging 24 h later. Protein localizations were quantified in line scans of the fluorescence signal of a cell by measuring the correlation of nuclear DAPI signals with signals emitted by YFP or YFP-tagged RCK-domains from BKα.

RBP1 and RBP2 double-mutant mice

RBP1/RBP2 conditional and constitutive DKO mice were described previously (Acuna et al, 2015, 2016). To delete RBP1 and RBP2 in the calyx of Held, conditional DKO mice were crossed with Krox-20-Cre mice (Voiculescu et al, 2000).

Electrophysiology

Presynaptic BK-currents were recorded in whole-cell voltage clamp configuration from calyx of Held terminals. The internal solution contained (in mM): 97.5 potassium gluconate, 32.5 KCl, 10 HEPES, 1 MgCl\(_2\), 12 Na\(_2\) phosphocreatine, 2 ATP-Mg, and 0.5 GTP-Na (295–305 mosmol/l, pH 7.3 adjusted with KOH; final K\(^+\) concentration, 143.5 mM). The internal solution was supplemented with the calcium chelator EGTA at the concentration of 0.2 mM (for Figs 5 and 6) and 10 mM (for Fig 7). The external solution (standard ACSF) contained (mM): 125 NaCl, 2.5 KCl, 25 NaHCO\(_3\), 1.25 NaH\(_2\)PO\(_4\), 2 CaCl\(_2\), 1 MgCl\(_2\), 10 glucose, 3 myo-inositol, 2 sodium pyruvate, and 0.5 ascorbic acid, pH 7.4 when bubbled with 95% O\(_2\) and 5% CO\(_2\). Tetrodotoxin (TTX; 1 μM) and 4-aminopyridine (4-AP; 2.5 mM) were added to the external solution to block, respectively, Na\(^+\) and K\(^+\) channels. BK-channels were blocked by application of Iberiotoxin (IbTX; 200 nM) to ACSF, perfusing slices at the rate of 1.0 ml/min.

Immunohistochemistry of brainstem sections

Immunohistochemistry experiments were performed as previously described (Zhang et al, 2015) (see Appendix). Serial confocal z-stack images were acquired using a Nikon confocal microscope (A1Rsi). Immunofluorescence was analysed with Nikon analysis software.

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Author contributions

CA performed the mouse genetics and electrophysiological experiments; FL performed the electrophysiological recordings with 10 mM EGTA; AS performed all other experiments; AS, CA, FL and TCS planned the experiments, analysed data, and wrote the paper.

Conflict of interest

The authors declare that they have no conflict of interest.

References

Acuna C, Liu X, Gonzalez A, Sudhof TC (2015) RIM-BPs mediate tight coupling of action potentials to Ca(2+)-triggered neurotransmitter release. Neuron 87: 1234–1247

Acuna C, Liu X, Sudhof TC (2016) How to make an active zone: unexpected universal functional redundancy between RIMs and RIM-BPs. Neuron 91: 792–807
Adelman JP, Shen KZ, Kavanaugh MP, Warren RA, Wu YN, Lagrutta A, Bond CT, North RA (1992) Calcium-activated potassium channels expressed from cloned complementary DNAs. Neuron 9: 209 – 216

Alle H, Kubota H, Geiger JR (2011) Sparse but highly efficient Kv3 outpace BKCa channels in action potential repolarization at hippocampal mossy fiber boutons. J Neurosci 31: 8001 – 8012

Atkinson NS, Robertson GA, Ganetzky B (1991) A component of calcium-activated potassium channels encoded by the Drosophila slo locus. Science 253: 551 – 555

Behrens R, Nolting A, Reimann F, Schwarz M, Waldschutz R, Pongs O (2000) hKCNNB3 and hKCNB4, cloning and characterization of two members of the large-conductance calcium-activated potassium channel beta subunit family. FEBS Lett 474: 99 – 106

Berkfeld H, Saier CA, Bild W, Rohde V, Thumfart JO, Eble S, Klugbauer N, Reisinger E, Bischofberger J, Oliver D, Knaus HG, Schulte U, Fakler B (2006) BKCa-Cav channel complexes mediate rapid and localized Ca\textsuperscript{2+}-activated \textit{K}\textsuperscript+ signaling. Science 314: 615 – 620

Berkfeld H, Fakler B (2013) Ligand-gating by Ca\textsuperscript{2+} is rate limiting for physiological operation of BK(Ca) channels. J Neurosci 33: 7358 – 7367

Betz A, Tahkur P, Junge HJ, Ashery U, Rhee JS, Rosenmund C, Retzig J, Brose N (2001) Functional interaction of the active zone proteins Munc13-1 and RIM1 in synaptic vesicle priming. Neuron 30: 183 – 196

Brenner R, Jegla TJ, Wickenden A, Liu Y, Aldrich RW (2000) Cloning and functional characterization of novel large conductance calcium-activated potassium channel beta subunits, hKCNB3 and hKCNB4. J Biol Chem 275: 6453 – 6461

Bucurenciu I, Kulik A, Schwaller B, Frotscher M, Jonas P (2008) Nanodom domain coupling between Ca\textsuperscript{2+} channels and Ca\textsuperscript{2+} sensors promotes fast and efficient transmitter release at a cortical GABAergic synapse. Neuron 57: 536 – 545

Butler A, Tsuoda S, McCobb DP, Wei A, Salkoff L (1993) mSlo, a complex mouse gene encoding “maxi” calcium-activated potassium channels. Science 261: 221 – 224

Contet C, Goulding SP, Kuljis DA, Barth AL (2016) BK channels in the central nervous system. Int Rev Neurobiol 128: 281 – 342

Davydova D, Marini C, King C, Klueva J, Bischof F, Romorini S, Montenegro-Venegas C, Heine M, Schneider R, Schroder MS, Altrock WD, Henneberger C, Rusakov DA, Gundelfinger ED, Fejtova A (2014) Bassoon specifically controls presynaptic P/Q-type Ca\textsuperscript{2+} channels via RIM-binding protein. Neuron 82: 181 – 194

Deng L, Kaeser PS, Xu W, Sudhof TC (2011) RIM proteins activate vesicle priming by reversing autoinhibitory homodimerization of Munc13. Neuron 69: 317 – 331

Eggermann E, Bucurenciu I, Goswami SP, Jonas P (2011) Nanodom domain coupling between Ca(2+)- and Ca\textsuperscript{2+}-sensors promotes fast and efficient transmitter release at a cortical GABAergic synapse. Neuron 57: 536 – 545

Fakler B, Adelman JP (2008) Control of K(Ca) channels by calcium nano-/microdomains. Neuron 59: 873 – 881

Fedchushyn MJ, Wang LY (2005) Developmental transformation of the release modality at the calyx of Held synapse. J Neurosci 25: 4131 – 4140

Galvez A, Gimenez-Gallego G, Reuben JP, Roy-Contancin L, Feigenbaum P, Kaczorowski Cj, Garcia ML (1990) Purification and characterization of a unique, potent, peptide probe for the high conductance calcium-activated potassium channel from venom of the scorpion Buthus tamulus. J Biol Chem 265: 11083 – 11090

Grauel MK, Maglione M, Reddy-Alla S, Willmes CG, Brockmann MM, Trimbach T, Rosenmund T, Pangalos M, Vardar G, Stumpf A, Walter AM, Rost BR, Eickholt BJ, Haucke V, Schmitz D, Signst SJ, Rosenmund C (2016) RIM-binding protein 2 regulates release probability by fine-tuning calcium channel localization at murine hippocampal synapses. Proc Natl Acad Sci USA 113: 11615 – 11620

Griguoli M, Sgritta M, Cherubini E (2016) Presynaptic BK channels control transmitter release: physiological relevance and potential therapeutic implications. J Physiol 594: 3489 – 3500

Grimes WN, Li W, Chavez AE, Diamond JS (2009) BK channels modulate pre- and postsynaptic signaling at reciprocal synapses in retina. Nat Neurosci 12: 585 – 592

Han Y, Kaeser PS, Sudhof TC, Schneggenburger R (2011) RIM determines Ca\textsuperscript{2+} channel density and vesicle docking at the presynaptic active zone. Neuron 69: 304 – 316

Hibino H, Pironkova R, Onwumere O, Vologodskia M, Hudspeth AJ, Lesage F (2002) RIM binding proteins (RIBPs) couple Rab3-interacting molecules (RIMs) to voltage-gated Ca(2+) channels. Neuron 34: 411 – 423

Hu H, Shaor LR, Chavoshy S, Gu N, Trieb M, Behrens R, Laka P, Pongs O, Knaus HG, Ottersen OP, Storm JF (2001) Presynaptic Ca\textsuperscript{2+}-activated \textit{K}\textsuperscript+ channels in glutamatergic hippocampal terminals and their role in spike repolarization and regulation of transmitter release. J Neurosci 21: 9585 – 9597

Indriati DW, Kamasawa N, Matsu K, Meredith AL, Watanabe M, Shimogero R (2013) Quantitative localization of Cav2.1 (P/Q-type) voltage-dependent calcium channels in Purkinje cells: somatodendritic gradient and distinct somatic coclustering with calcium-activated potassium channels. J Neurosci 33: 3668 – 3678

Ishikawa T, Nakamura Y, Saitoh N, Li WB, Iwasaki S, Takahashi T (2003) Distinct roles of Kv1 and Kv3 potassium channels at the calyx of Held presynaptic terminal. J Neurosci 23: 10445 – 10453

Kaeser PS, Deng L, Wang Y, Dulubova I, Liu X, Rizo J, Sudhof TC (2011) RIM proteins tether Ca\textsuperscript{2+} channels to presynaptic active zones via a direct PDZ-domain interaction. Cell 144: 282 – 295

Knaus HG, Schwarzer C, Koch RO, Eberhart A, Kaczorowski GJ, Glossmann H, Wunder F, Pongs O, Garcia ML, Sperr G (1996) Distribution of high-conductance Ca\textsuperscript{2+}-activated K\textsuperscript+ channels in rat brain: targeting to axons and nerve terminals. J Neurosci 16: 955 – 963

Krinne S, Butola T, Jung S, Wichmann C, Moser T (2017) RIM-binding protein 2 promotes a large number of Cav1.3 Ca\textsuperscript{2+}-channels and contributes to fast synaptic vesicle replenishment at hair cell active zones. Front Cell Neurosci 11: 334

Lee US, Cui J (2010) BK channel activation: structural and functional insights. Trends Neurosci 33: 415 – 423

Liu KS, Siebert M, Mertel S, Knoche E, Siebert M, Muette C, Buckers J, Hell SW, Muller M, Davis GW, Schmitz D, Sigrist SJ (2011) RIM-binding protein, a central part of the Ca\textsuperscript{2+} channels via RIM-binding protein. Neuron 82: 181 – 194

Loane DJ, Lima PA, Marrion NV (2007) Co-assembly of N-type Ca\textsuperscript{2+} and BK channels underlies functional coupling in rat brain. J Cell Sci 120: 985 – 995

Luo F, Liu X, Sudhof TC, Acuna C (2017) Efficient stimulus–secretion coupling at ribbon synapses requires RIM-binding protein tethering of L-type Ca\textsuperscript{2+} channels. Proc Natl Acad Sci USA 114: E8081 – E8090

Meera P, Wallner M, Jiang Z, Toro L (1996) A calcium switch for the functional coupling between alpha (hslo) and beta subunits (KV, Ca beta) of maxi K channels. FEBS Lett 382: 84 – 88

Meinrenken CJ, Borst JG, Sakmann B (2002) Calcium secretion coupling at calyx of Held governed by nonuniform channel-vesicle topography. J Neurosci 22: 1648 – 1667
Misonou H, Menegola M, Buchwalder L, Park EW, Meredith A, Rhodes KJ, Aldrich RW, Trimmer JS (2006) Immunolocalization of the Ca\(^{2+}\)-activated K\(^+\) channel Slo1 in axons and nerve terminals of mammalian brain and cultured neurons. *J Comp Neurol* 496: 289 – 302

Mittelstaedt T, Schoch S (2007) Structure and evolution of RIM-BP genes: identification of a novel family member. *Gene* 403: 70 – 79

Modchang C, Nadkarni S, Bartol TM, Triampo W, Sejnowski TJ, Levine H, Rappel WJ (2010) A comparison of deterministic and stochastic simulations of neuronal vesicle release models. *Phys Biol* 7: 026008

Muller M, Genc O, Davis GW (2015) RIM-binding protein links synaptic homeostasis to the stabilization and replenishment of high release probability vesicles. *Neuron* 85: 1056 – 1069

Nakamura Y, Takahashi T (2007) Developmental changes in potassium currents at the rat calyx of Held presynaptic terminal. *J Physiol* 581: 1101 – 1112

Oro P, Rojas P, Ferreira G, Latorre R (2002) New disguises for an old channel: maxK channel beta-subunits. *Neus Physiol Sci* 17: 156 – 161

Poulsen AN, Wulf H, Hay-Schmidt A, Jansen-Olesen I, Olesen J, Klaerke DA (2009) Differential expression of BK channel isoforms and beta-subunits in rat neuro-vascular tissues. *Biochem Biophys Acta* 1788: 380 – 389

Raffaelli G, Saviane C, Mohajerani MH, Pedarzani P, Cherubini E (2004) BK potassium channels control neurotransmitter release at CA3-CA3 synapses in the rat hippocampus. *J Physiol* 557: 147 – 157

Roberts WM (1993) Spatial calcium buffering in saccular hair cells. *Nature* 363: 74 – 76

Robitaille R, Adler EM, Charlton MP (1993a) Calcium channels and calcium-gated potassium channels at the frog neuromuscular junction. *J Physiol Paris* 87: 15 – 24

Robitaille R, Garcia ML, Kaczorowski GJ, Charlton MP (1993b) Functional colocalization of calcium and calcium-gated potassium channels in control of transmitter release. *Neuron* 11: 645 – 655

Saussier U, Saussier M, Sailer CA, Arintz C, Knaus HG, Neuhuber W, Ruth P (2006) Ca\(^{2+}\)-activated K\(^+\) channels of the BK-type in the mouse brain. *Histochem Cell Biol* 125: 725 – 741

Schneggenburger R, Forsythe ID (2006) The calyx of Held. *Cell Tissue Res* 326: 311 – 337

Schoch S, Castillo PE, Jo T, Mukherjee K, Ceppert M, Wang Y, Schmitz F, Malenka RC, Sudhof TC (2002) RIM1alpha forms a protein scaffold for regulating neurotransmitter release at the active zone. *Nature* 415: 321 – 326

Storm JF (1987) Action potential repolarization and a fast after-hyperpolarization in rat hippocampal pyramidal cells. *J Physiol* 385: 733 – 759

Sudhof TC (2012) The presynaptic active zone. *Neuron* 75: 11 – 25

Sudhof TC (2013) Neurotransmitter release: the last millisecond in the life of a synaptic vesicle. *Neuron* 80: 675 – 690

Tseng-Crank J, Godinot N, Johansen TE, Ahring PK, Strobaek D, Mertz R, Foster CD, Olesen SP, Reinhart PH (1996) Cloning, expression, and distribution of a Ca\((2+)\)-activated K\(^+\) channel beta-subunit from human brain. *Proc Natl Acad Sci USA* 93: 9200 – 9205

Ubele VN, Lagrutta A, Wade T, Figueroa DJ, Liu Y, McKenna E, Austin CP, Bennett PB, Swanson R (2000) Cloning and functional expression of two families of beta-subunits of the large conductance calcium-activated K\(^+\) channel. *J Biol Chem* 275: 23211 – 23218

Voiculescu O, Charnay P, Schneider-Maunoury S (2000) Expression pattern of a KroX-20/Cre knock-in allele in the developing hindbrain, bones, and peripheral nervous system. *Genesis* 26: 123 – 126

Vyleta NP, Jonas P (2014) Loose coupling between Ca\(^{2+}\) channels and release sensors at a plastic hippocampal synapse. *Science* 343: 665 – 670

Wallner M, Meera P, Toro L (1996) Determinant for beta-subunit regulation in high-conductance voltage-activated and Ca\((2+)\)-sensitive K\(^+\) channels: an additional transmembrane region at the N terminus. *Proc Natl Acad Sci USA* 93: 14922 – 14927

Walter AM, Groffen AJ, Sorensen JB, Verhage M (2011) Multiple Ca\(^{2+}\) sensors in secretion: teammates, competitors or autocrats? *Trends Neurosci* 34: 487 – 497

Wang Y, Okamoto M, Schmitz F, Hofmann K, Sudhof TC (1997) Rim is a putative Rab3 effector in regulating synaptic-vesicle fusion. *Nature* 388: 593 – 598

Wang Y, Sugita S, Sudhof TC (2000) The RIM/NIM family of neuronal C2 domain proteins. Interactions with Rab3 and a new class of Src homology 3 domain proteins. *J Biol Chem* 275: 20033 – 20044

Wang ZW (2008) Regulation of synaptic transmission by presynaptic CaMKII and BK channels. *Mol Neurobiol* 38: 153 – 166

Xia XM, Zeng X, Lingle CJ (2002) Multiple regulatory sites in large-conductance calcium-activated potassium channels. *Nature* 418: 880 – 884

Yuan P, Leonetti MD, Pico AR, Hsiung Y, MacKinnon R (2000) The RCK domain of the human BKCa channel is a calcium sensor. *Proc Natl Acad Sci USA* 105: 376 – 381

Zhang B, Chen LY, Liu X, Maxeiner S, Lee SJ, Gokce O, Sudhof TC (2015) Neurouilin1 sculpts Cerebellar Purkinje-cell circuits by differential control of distinct classes of synapses. *Neuron* 87: 781 – 796

Zhou Y, Schopperle WM, Murray H, Jaramillo A, Dagan D, Griffith LC, Levitan IB (1999) A dynamically regulated 14-3-3, Slob, and Slowpoke potassium channel complex in Drosophila presynaptic nerve terminals. *Neuron* 22: 809 – 818

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