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RIM Promotes Calcium Channel Accumulation at Active Zones of the Drosophila Neuromuscular Junction

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Synaptic communication requires the controlled release of synaptic vesicles from presynaptic axon terminals. Release efficacy is regulated by the many proteins that comprise the presynaptic release apparatus, including \( \text{Ca}^{2+} \) channels and proteins that influence \( \text{Ca}^{2+} \) channel accumulation at release sites. Here we identify \textit{Drosophila} RIM (Rab3 interacting molecule) and demonstrate that it localizes to active zones at the larval neuromuscular junction. In \textit{Drosophila} RIM mutants, there is a large decrease in evoked synaptic transmission because of a significant reduction in both the clustering of \( \text{Ca}^{2+} \) channels and the size of the readily releasable pool of synaptic vesicles at active zones. Hence, RIM plays an evolutionarily conserved role in regulating synaptic calcium channel localization and readily releasable pool size. Because RIM has traditionally been studied as an effecter of Rab3 function, we investigate whether RIM is involved in the newly identified function of Rab3 in the distribution of presynaptic release machinery components across release sites. Bruchpilot (Brp), an essential component of the active zone cytomatrix T bar, is unaffected by RIM disruption, indicating that Brp localization and distribution across active zones does not require wild-type RIM. In addition, larvae containing mutations in both \textit{RIM} and \textit{rab3} have reduced \( \text{Ca}^{2+} \) channel levels and a Brp distribution that is very similar to that of the \textit{rab3} single mutant, indicating that RIM functions to regulate \( \text{Ca}^{2+} \) channel accumulation but is not a Rab3 effecter for release machinery distribution across release sites.

Introduction

Synaptic vesicle exocytosis occurs at specialized regions of the presynaptic membrane, termed active zones, in which presynaptic release machinery proteins cluster opposite postsynaptic neurotransmitter receptors. The complement of presynaptic proteins associated with each active zone is a determinant of the release properties at each release site (Fejtova and Gundelfinger, 2006; Sigrist and Schmitz, 2011). In particular, proteins that determine the number of \( \text{Ca}^{2+} \) channels at each release site control synaptic efficacy.

At the \textit{Drosophila} larval neuromuscular junction (NMJ), Bruchpilot (Brp) and RIM-binding protein (RIM-BP) regulate \( \text{Ca}^{2+} \) channel accumulation (Kittel et al., 2006; Liu et al., 2011). RIM (Rab3 interacting molecule) enhances \( \text{Ca}^{2+} \) channel levels at mammalian synapses (Han et al., 2011; Kaeser et al., 2011); however, RIM has not been characterized previously in flies. RIM is an active zone protein that acts as an organizer of the presynaptic release apparatus via its interactions with multiple core active zone components, including \( \alpha \)-liprins, Munc-13, RIM-BPs, and \( \text{Ca}^{2+} \) channels (Wang et al., 2000; Betz et al., 2001; Coppola et al., 2001; Schoch et al., 2002; Dulubova et al., 2005; Kiyonaka et al., 2007). RIM also interacts with CAST/ERC in mice (Ohtsuka et al., 2002; Wang et al., 2002) and ELKS in \textit{Caenorhabditis elegans} (Deken et al., 2005), the mammalian and worm orthologs of Brp. Hence, RIM may have a similar function in \textit{Drosophila}.

In rodents and \textit{C. elegans}, RIM binds to and is an effecter of the small GTPase Rab3. We demonstrated previously that Rab3 dynamically controls the presynaptic protein composition of individual active zones at the \textit{Drosophila} NMJ (Graf et al., 2009). At wild-type (WT) NMJs, release machinery proteins are distributed across all active zones, resulting in the formation of hundreds of low probability release sites peppered with higher probability sites (Marrus and DiAntonio, 2004; Peled and Isacoff, 2011). Conversely, in the \textit{rab3} mutant, key constituents of the presynaptic active zone, including Brp and \( \text{Ca}^{2+} \) channels, are enriched at a subset of active zones, leaving the remaining release sites apparently devoid of such proteins (Graf et al., 2009). The molecular mechanisms by which Rab3 mediates this function remain essentially unknown; however, RIM is a potential effecter for Rab3 in this process.

To test the role of RIM at \textit{Drosophila} active zones, we cloned the single \textit{Drosophila} ortholog of RIM and generated RIM excision mutants (Müller et al., 2012). We show that \textit{Drosophila} RIM...
localizes to active zones and that its distribution requires Rab3. Mutant analysis demonstrates that full-length RIM is not necessary for the proper localization of Brp or for the altered distribution of Brp across active zones in rab3 mutants. Rather, RIM enables robust Ca$^{2+}$-dependent synaptic release by promoting the accumulation of Ca$^{2+}$ channels at release sites and synaptic vesicles in the readily releasable pool (RRP). Hence, WT RIM is not an essential effector of Rab3 for the control of protein composition across active zones but instead promotes synaptic efficacy by enhancing Ca$^{2+}$ channel density and the size of the RRP at the Drosophila NMJ.

**Materials and Methods**

**Fly stocks.** Flies were maintained at 25°C on standard fly food. WT flies were Canton S or Canton S outcrossed to elav–Gαd (Yao and White, 1994) or dvglt$^{NMX}$. Gαd (Daniels et al., 2008) based on the experiment. The following fly lines were obtained from the Bloomington Stock Center: the P-element line P{EPgy2}RimEY05246, the deficiency line Df(3R)ED5785, the UAS–Cacophony–GFP line P[UAS–cac1–GFP]422A (Kawasaki et al., 2004), and UAS–Dcr2 (Dietz et al., 2007). The rab3$^{ex73}$ mutant was described previously (Graf et al., 2009). The line containing the UAS–rab3 RNAi transgene was obtained from the Vienna Drosophila RNAi Center (Dietz et al., 2007). The line containing the UAS–brp RNAi transgene was obtained from Stephan Sigrist (University of Berlin, Berlin, Germany) (Kittel et al., 2006; Wagh et al., 2006).

**Cloning of RIM cDNA.** To generate the 7.4 kb full-length RIM cDNA, we cloned three overlapping cDNA fragments spanning the entire RIM opening reading frame and then sequentially ligated individual fragments together. In brief, total RNA was extracted from adult Drosophila heads, and reverse transcription (RT) reactions were performed using gene-specific primers to obtain three cDNA fragments that were ligated together using the unique internal cutting sites MluI and EcoRI. 5’ end to MluI (2288), MluI (2288) to EcoRI (5162), and EcoRI (5162) to 3’ end. NotI and XbaI sites were introduced onto the 5’ end and 3’ end, respectively, to facilitate subcloning into the pUAST vector (Brand and Perrimon, 1993) to generate untagged UAS–RIM. The RIM cDNA was also subcloned into the pUAST–EGFP (enhanced green fluorescent protein) vector (Parker et al., 2001) to generate the UAS–RIM::EGFP transgene that results in the fusion of EGFP fused to the N terminus of RIM.

**Generation of RIM mutants.** Deletions of the RIM gene were generated by imprecise excision of the P-element P{EPgy2}RimEY05246, which is located within an intron near the 3’ end of the RIM gene. To initiate the excision, homozygous y,w$: P{EPgy2}RimEY05246 females were crossed to y,w$: w-;Xa/CyO, D2–3,Sb males carrying the transposase. F1 male progeny y,w$: y,w$: P{EPgy2}RimEY05246/D2–3,Sb were collected and crossed to y,w$: y,w$: Tm6B,Scr to set up stocks that were subsequently screened by PCR to identify those in which DNA surrounding the P-element had been excised. Two mutants with substantial excisions, RIM$^{ex73}$ and RIM$^{ex89}$, were identified, and the following primer pairs were chosen to span the respective excised regions, creating PCR products that were sequenced to determine the precise nature of each excision: RIM$^{ex73}$, 5’–CCCGGACCTTACACCGGTATTCCGAGGCTGC–3’; RIM$^{ex89}$, 5’–CACCTACGGACCCCGACACCGAGG–3’; and 5’–CTGAAAGGAGATTCTTGCGGAAAGG–3’.

**Immunohistochemistry.** Third-instar larvae were dissected in PBS and fixed in either Bouin’s fixative for 5 min or 4% formaldehyde for 30 min. Larvae were washed with PBS containing 0.1% Triton X-100 (PBT) and blocked in 5% NGS in PBT for 30 min, followed by overnight incubation in primary antibodies in 5% NGS in PBT, three washes in PBT, incubation in secondary antibodies in 5% NGS in PBT for 45 min, three final washes in PBT, and equilibration in 70% glycerol in PBS. Samples were mounted in VectaShield (Vector Laboratories). The following primary antibodies were used: mouse α–Brp, 1:250 (Developmental Studies Hybridoma Bank); rabbit α–DGlurRIII, 1:2500 (Marrus et al., 2004); and rabbit α–Rab3, 1:1000 (Graf et al., 2009). Goat Cy3-conjugated secondary antibodies against mouse (Jackson ImmunoResearch) and Alexa Fluor 488-conjugated secondary antibodies against rabbit and mouse (Invitrogen) were used at 1:1000. Alexa Fluor 488-conjugated rabbit α–GFp (1:1000; Invitrogen) was used to visualize GFp-tagged fusion proteins. Antibodies obtained from the Developmental Studies Hybridoma Bank were developed under the auspices of the National Institute of Child Health and Human Development and maintained by the Department of Biological Sciences of the University of Iowa (Iowa City, IA).

**Imaging and analysis.** Samples were imaged using a Carl Zeiss LSM 5 Pascal confocal microscope or a Nikon C1 confocal microscope. All genotypes for an individual experiment were imaged at the same gain and set such that signals from the brightest genotype for a given experiment were not saturating. Images for quantifying average Cacophony:GFp intensity and percentage GlurIIII clusters apposed to Brp were randomized and analyzed using Velocity software (PerkinElmer Life and Analytical Sciences). For quantification of percentage apposition, GlurIIII clusters and Brp puncta were defined as objects, and the percentage of GlurIIII objects that overlapped with a Brp object was determined. For measurements of Cacophony:GFp intensity, Brp objects were first defined. The average intensity of Cacophony:GFp signal within each defined Brp object was then calculated, and the average muscle background intensity was subtracted. Images for quantifying average Brp area were randomized and analyzed using MetaMorph software (Universal Imaging Corporation). Statistical analysis was performed using ANOVA for comparison of samples within an experimental group. All histograms and measurements are shown as mean ± SEM.

**Electrophysiology.** Electrophysiological experiments were performed as described previously (Daniels et al., 2006). Male third-instar larvae were dissected in HL-3 saline containing 0.42 msc Ca$^{2+}$ and then washed with HL3 solution. Two-electrode voltage-clamp recordings were performed in HL-3 saline containing 0.42 msc Ca$^{2+}$ unless indicated otherwise from muscle 6 in segments A3 and A4. Muscle cells were clamped at −70 mV with less than −1.0 nA holding current, and then both spontaneous miniature excitatory junction currents (mEJC) and EJCs were recorded. One hundred consecutive miniature events were measured per cell and averaged to determine mean mEJC amplitude for each cell. For evoked recordings, the end of the cut segmental nerve was sucked into an electrode and stimulated with a 1 ms pulse. The amplitude of the stimulus was set to ensure recruitment of both nerves innervating muscle 6. EJC amplitude was determined by averaging 10 consecutive EJCs delivered at 0.2 Hz using Clampfit 9.0. Quantal content was estimated by dividing the mean EJC by the mean mEJC. Statistical analysis was performed using ANOVA for comparison of samples within an experimental group. Histograms and measurements are shown as mean ± SEM.

**Estimation of the RRP was performed according to the method of cumulative amplitude analysis (Schneegensburger et al., 2002; Baldelli et al., 2007; Weyhersmüller et al., 2011). Peak EJCs were summed from 20 repetitive stimuli applied at a frequency of 40 Hz in high Ca$^{2+}$ (2 mM) HL3 solution. The amplitude of the EJC was determined as the difference between the peak current and the baseline before the stimulation onset. This analysis assumes that the depression during the high-frequency stimulation is limited by the recycling of synaptic vesicles. After the RRP has been depleted, equilibrium is reached between released and recycled vesicles. To determine the initial size of the RRP, the last 10 points were fitted by a linear regression and back–extrapolated to time 0 to determine the initial size of the RRP. The quantal content for each peak was determined by dividing the EJC amplitude by the mean mEJC amplitude for that cell. Our method for assessing RRP uses lower-frequency stimulation but higher Ca$^{2+}$ concentration than previous studies, yet the estimation of the RRP size in WT animals is very similar to those reported previously (Weyhersmüller et al., 2011), indicating that the technique is robust to small changes in protocol.

**Electron microscopy.** Electron microscopy (EM) was performed as described previously (Liu et al., 2011). Briefly, dissected third-instar larvae were fixed overnight at 4°C in modified Trump’s fixative (1% glutaraldehyde and 4% paraformaldehyde in 0.1 M cacodylate buffer, pH 7.4) and postfixed in 1% OsO, in cacodylate buffer and stained en bloc with saturated aqueous uranyl acetate. Samples were then dehydrated and embedded in Spurr resin (Electron Microscopy Sciences). NMJ 6/7 terminals in abdominal segments A2 and A3 were serially sectioned with a Leica UC6
RIM::GFP under the control of a neuronally expressed galactosidase-4 (Gal4) in a WT background results in detection of GFP signal in the axon terminals of larval motor neurons. Furthermore, RIM::GFP concentrates at release sites, colocalizing with Brp puncta (Fig. 1B). When expressed in a WT background in which endogenous RIM is present, RIM::GFP is only mildly enriched at active zones. If there were a limited number of RIM molecules that can associate with active zones, then RIM::GFP may have to compete with endogenous RIM for localization to the active zone. We therefore assessed whether RIM::GFP localization to release sites would be enhanced if driven in a RIM mutant background. To address this, we drove UAS–RIM::GFP

ultramicrotome, stained with uranyl acetate and Sato’s lead, and observed using a FEOL 1400 electron microscope. Cross-sections through the center of a bouton with at least one T-bar and clear presynaptic and postsynaptic densities were selected for quantitative analysis. Active zone length was defined by the length of synaptic densities with a T-bar attached. Images of at least 19 boutons from four larvae of each genotype were analyzed.

**Results**

**RIM localizes to active zones**

As a potential regulator of Ca$^{2+}$ channel accumulation and Rab3 effector at the larval NMJ, we wanted to determine whether RIM functions at the *Drosophila* active zone. A single ortholog of *Drosophila* RIM had been predicted by analysis of genomic sequences that contain transcription units encoding typical RIM domains (Wang and Südhof, 2003); however, no full-length RIM cDNA clone has been identified. Therefore, we began by cloning RIM in three overlapping cDNA fragments from mRNA extracted from adult *Drosophila* heads and ligating the three fragments together to form a full-length 7.4 kb RIM cDNA (Fig. 1A). As predicted, the *Drosophila* RIM transcript contains coding regions for all four of the standard RIM Zn$^{2+}$ finger, PDZ, C$_{A}$, and C$_{B}$ domains (Fig. 1A, exons labeled in red), although the 7.4 kb *Drosophila* open reading frame is significantly longer than the 4.7 kb open reading frames in rodent and *C. elegans* RIM because of a lengthening of regions separating the domains. Furthermore, two RIM splice variants were identified, each containing one of two alternatively spliced exons near the C-terminal end of the transcript (exons S1 and S2 shown in blue in Fig. 1A).

Identification of a full-length *Drosophila* RIM transcript indicates that an evolutionarily conserved RIM protein is expressed in *Drosophila*. Studies of RIM in rodents and *C. elegans* indicate that RIM proteins localize to presynaptic release sites and bind to proteins associated with presynaptic active zones (Wang et al., 1997; Koushika et al., 2001; Schoch et al., 2002; Deeken et al., 2005). To characterize the localization of *Drosophila* RIM protein, we attempted to generate antibodies against multiple regions of the RIM protein. However, despite several attempts, we were unable to identify antibodies that recognized RIM via either immunoblot or immunohistochemistry. Therefore, to analyze the neuronal localization pattern of RIM, we fused GFP to the N terminus of both RIM splice variants and used the UAS/Gal4 system to express transgenic RIM::GFP in *Drosophila* larvae. To determine whether RIM preferentially localizes to active zones within the *Drosophila* NMJ, we stained UAS–RIM::GFP expressing neurons with antibodies against the presynaptic active zone protein Brp. Expression of UAS–RIM::GFP in a WT background results in detection of GFP signal in the axon terminals of larval motor neurons. Furthermore, RIM::GFP concentrates at release sites, colocalizing with Brp puncta (Fig. 1B). When expressed in a WT background in which endogenous RIM is present, RIM::GFP is only mildly enriched at active zones. If there were a limited number of RIM molecules that can associate with active zones, then RIM::GFP may have to compete with endogenous RIM for localization to the active zone. We therefore assessed whether RIM::GFP localization to release sites would be enhanced if driven in a RIM mutant background. To address this, we drove UAS–RIM::GFP

**Figure 1.** Transgenically expressed RIM concentrates at active zones. A, Schematic representations of the cloned WT *Drosophila* RIM gene and the RIM$^{\text{Ex73}}$ and RIM$^{\text{Ex98}}$ mutant genes generated by P-element excision. Red exonic regions correspond to the standard RIM Zn$^{2+}$ finger (Zn), PDZ, C$_{A}$, and C$_{B}$ domains. Blue exonic regions represent the two alternatively spliced exons (S1 and S2). Boxed regions of the RIM$^{\text{Ex73}}$ and RIM$^{\text{Ex98}}$ mutant genes denote the excised region of each mutant. In the RIM$^{\text{Ex73}}$ mutant, the gray exonic sequences downstream of the excised region denote exons that are thrown out of frame because of a frame shift. B–E, Confocal images of third-instar larval NMJs transgenically expressing the S1 splice variant of RIM::GFP (green) and costained with antibodies against Brp (red). UAS–RIM::GFP expression is driven by the neuronal elav–Gal4 driver in a WT background (B) (UAS–RIM::GFP/+; elav–Gal4), the RIM$^{\text{Ex73}}$ mutant background (C) (UAS–RIM::GFP/+; RIM$^{\text{Ex73}}$, elav–Gal4/Df(3R)ED5785), UAS–Rab3 RNAi expressing neurons (D) (UAS–Rab3 RNAi, UAS–DCR2/UAS–RIM::GFP;elav–Gal4/+), and control (UAS–DCR2+/UAS–RIM::GFP/+; elav–Gal4/+). UAS–RIM::GFP expressing neurons (E) (UAS–DCR2+/UAS–RIM::GFP/+; Brp RNAi/elav–Gal4). UAS–DCR2 is coexpressed with the UAS–RNAi transgenes to enhance the Brp knockdown of Rab3 and Brp. Scale bars, 2 μm.
Interestingly, RIM::GFP is localized to active zones in a much higher percentage of Brp puncta than observed for both the S1 and S2 splice variants (data not shown). In the absence of Brp, RIM::GFP localizes in a punctate pattern that is similar to that seen in WT (Fig. 1E). These findings suggest that Brp is not required for the localization of RIM to release sites. Similar observations have been made in C. elegans in which the Brp homolog ELKS is also not required for localizing RIM to active zones (Deken et al., 2005). Together, our results indicate that transgenically expressed RIM::GFP localizes to release sites, that its localization is regulated by Rab3, and demonstrate that Rab3 provides an important context for Brp localization.

Figure 2. RIM is not required for Brp localization and normal synaptic ultrastructure. A, Images of muscle 4 NMJs from WT and RIM(k012453)Df(3R)ED5785 mutants stained for the presynaptic active zone protein Brp (red) and the postsynaptic receptor DGlurIII (green). Scale bar, 2 μm. B, C, Micrographs of NMJ6/7 boutons from WT (B) and RIM(k012453)Df(3R)ED5785 mutants (C). Electron-dense membranes demarcated by arrowheads indicate synaptic vesicle release sites (active zones). T-bars are observed at some active zones (arrow). M, Mitochondria. D, E, Higher-magnification views of the active zones containing T-bars from WT (B) and RIM(k012453)Df(3R)ED5785 mutants (C). T-bar pedestals are marked with arrows.

in RIM excisions mutants (which are described in detail below). Interestingly, RIM::GFP is localized to active zones in a much brighter, more punctate manner when driven in the RIM mutant (Fig. 1C), consistent with our hypothesis that RIM::GFP must compete with endogenous RIM for space at the active zone when driven in a WT background. A similar localization pattern is observed for both the S1 and S2 splice variants (data not shown).

Comparison of RIM::GFP intensity at individual active zones in the RIM mutant background indicates that RIM::GFP preferentially localizes to active zones with brighter α-Brp signal. Binning Brp puncta into three groups based on Brp average intensity (dim Brp, 49 ± 0.2 a.u., n = 568 puncta; moderate Brp, 60 ± 0.1 a.u., n = 593 puncta; bright Brp, 75 ± 0.4 a.u., n = 212 puncta; p << 0.001 for moderate vs both bright and dim) and measuring the average intensity of RIM::GFP colocalized with Brp within each bin indicates that Brp and RIM::GFP intensities are correlated (RIM::GFP at dim Brp, 28 ± 0.6 a.u., n = 568 puncta; RIM::GFP at moderate Brp, 35 ± 0.6 a.u., n = 593 puncta; RIM::GFP at bright Brp, 48 ± 1.2 a.u., n = 212 puncta; p << 0.001 for moderate vs both bright and dim).
RIM is not essential for the clustering of RIM::GFP to the active zone.

**RIM is dispensable for proper Brp localization and active zone ultrastructure**

Because RIM::GFP localizes to active zones, it is in a position to influence the clustering of other presynaptic active zone components. How are active zones affected when endogenous RIM is disrupted? To address this question, we generated mutants by excision of a transposable P-element located within the RIM gene. Imprecise excision of P[EPy2]/RimEy05246 generated two large independent excisions, RIM\textsuperscript{Df}[3R]/H11006 and RIM\textsuperscript{Df}[3R]/H11022, that each result in the deletion of the majority of the central region of the RIM gene (Fig. 1A). These mutant alleles all contain the beginning and end of the RIM gene; however, 13 and 19.4 kb of genomic sequence has been excised from RIM\textsuperscript{Df}[3R] and RIM\textsuperscript{Df}[3R], respectively. The genomic sequences that code for the PDZ and C2A domains are fully removed in both mutant alleles; however, sequences coding for the Zn\textsuperscript{2+} finger and C2B domains remain.

Both the N-terminal and C-terminal domains exist in the RIM mutant alleles, so the RIM mutants could express a truncated RIM protein containing these domains. Indeed, RT-PCR analysis of RIM transcripts expressed in each mutant demonstrates that, in both RIM\textsuperscript{Df}[3R] and RIM\textsuperscript{Df}[3R], the exons to either side of the excised region can splice together, creating mRNA transcripts that contain the exonic coding sequences that remain in the excised alleles. In RIM\textsuperscript{Df}[3R], such splicing may result in a correctly translated C2B domain. However, in RIM\textsuperscript{Df}[3R], this splicing results in a frame shift such that coding sequences downstream of the RIM\textsuperscript{Df}[3R] excision are thrown out of frame (indicated in gray in Fig. 1A). In RIM\textsuperscript{Df}[3R], a frame shift occurs regardless of whether the alternatively spliced S1 or S2 exons are included in the transcript, so if a truncated mutant RIM protein were created in RIM\textsuperscript{Df}[3R], it would not contain the C2B domain. Because an antibody is not available to test whether a mutant version of RIM protein is expressed in either RIM mutant, we have used the RIM\textsuperscript{Df}[3R] mutant in the majority of our studies based on this transcript analysis.

Homozygous RIM\textsuperscript{Df}[3R] and RIM\textsuperscript{Df}[3R] mutants are viable and fertile and show no locomotor defects when compared with WT in a weak geotaxis assay ($p > 0.1$). Furthermore, no gross morphological defects are observed at the NMJs of homozygous mutant larvae. Placing RIM\textsuperscript{Df}[3R] and RIM\textsuperscript{Df}[3R] mutant chromosomes in trans to the Df(3R)/ED5785 deficiency chromosome that uncovers the entire RIM gene also results in viable animals with no gross morphological defects.

Because transgenic RIM localizes to and may function at active zones, we wanted to determine whether active zone components are affected by the absence of WT RIM protein. As an essential component of T-bars, Brp is an important constituent of the Drosophila presynaptic active zone (Kittel et al., 2006; Wagh et al., 2006; Fouquet et al., 2009). Therefore, we first asked whether Brp localization is disrupted in the RIM\textsuperscript{Df}[3R] mutant. We hypothesized that (1) RIM may be required for Brp localization or (2) RIM may work with Rab3 to control protein composition across active zones. If Brp localization requires RIM, we would predict a general reduction in Brp at RIM mutant active zones. Alternatively, if RIM acts as a Rab3 effector to control active zone protein composition, we would predict a RIM mutant phenotype similar to that observed in the rab3 mutant. At WT NMJs, immunostaining with antibodies against Brp and the essential glutamate receptor subunit GluRIII reveals a localization pattern in which discrete presynaptic Brp puncta are directly apposed to nearly all postsynaptic glutamate receptor clusters. In RIM\textsuperscript{Df}[3R] mutant NMJs, Brp and GluRIII immunostaining appears indistinguishable from WT (Fig. 2A). Quantification of Brp and GluRIII staining indicates no significant difference between WT and RIM\textsuperscript{Df}[3R] mutant NMJs both in terms of the percentage of GluRIII clusters that are apposed to Brp puncta [WT, 95 ± 1% puncta, $n = 13$; RIM\textsuperscript{Df}[3R]/ED5785, 97 ± 1%, $n = 17$ NMJs; $p > 0.3$] and the average area of individual Brp puncta [WT, 0.18 ± 0.01 $\mu m^2$, $n = 13$ NMJs; RIM\textsuperscript{Df}[3R]/ED5785, 0.19 ± 0.01 $\mu m^2$, $n = 16$; $p > 0.5$]. Because there is no decrease in Brp levels and the distribution of Brp across active zones is normal, neither hypothesis is supported.

As a necessary component of T-bars, Brp immunostaining appears normal in the RIM mutant. However, to determine whether RIM mutation affects T-bar and active zone ultrastructure in a manner that is undetectable by light microscopy, we analyzed RIM mutant NMJs from muscles 6/7 via EM (Fig. 2B–F). No obvious ultrastructural differences are observed between WT and RIM\textsuperscript{Df}[3R]/ED5785 mutant boutons. Active zones, visualized by electron dense membranes (arrowheads), and T-bars (arrows) with their associated vesicles are readily observed at both WT and RIM mutant boutons (Fig. 2B–E). Quantification of EM images of WT and RIM mutant synapses reveals no morphological difference in active zone length [WT, 0.73 ± 0.05 $\mu m$, $n = 21$; RIM\textsuperscript{Df}[3R]/ED5785, 0.64 ± 0.02 $\mu m$, $n = 22$; $p > 0.1$], T-bar width [WT, 171.3 ± 6.4 nm, $n = 57$; RIM\textsuperscript{Df}[3R]/ED5785, 165.7 ± 5.7 nm, $n = 51$; $p > 0.3$], and number of T-bars per active zone [WT, 0.46 ± 0.05, $n = 19$; RIM\textsuperscript{Df}[3R]/ED5785, 0.40 ± 0.03, $n = 21$; $p > 0.5$]. However, RIM mutant NMJs have a small decrease in the number of T-bars per bouton perimeter [WT, 0.26 ± 0.03 $\mu m$, $n = 19$; RIM\textsuperscript{Df}[3R]/ED5785, 0.18 ± 0.02 $\mu m$, $n = 21$; $p < 0.05$] and active zones per bouton perimeter [WT, 0.56 ± 0.03 $\mu m$, $n = 19$; RIM\textsuperscript{Df}[3R]/ED5785, 0.46 ± 0.03 $\mu m$, $n = 21$; $p < 0.05$]. A comparable reduction in the density of Brp puncta at muscle 4 NMJs is also observed at the light level [WT, 1.66 ± 0.03 puncta per $\mu m^2$, $n = 13$; RIM\textsuperscript{Df}[3R]/ED5785, 1.42 ± 0.03 puncta per $\mu m^2$, $n = 13$; $p < 0.001$] because of a 15% decrease in the number of Brp puncta per NMJ [WT, 377 ± 16 puncta, $n = 13$; RIM\textsuperscript{Df}[3R]/ED5785, 327 ± 10, $n = 13$; $p < 0.05$]. Thus, active zone ultrastructure is normal in the RIM mutant, but analysis at both the EM and light levels reveals a modest reduction in the number of release sites.

**RIM is required for Ca\textsuperscript{2+} channel accumulation at active zones**

The observation that Brp is unaffected by the RIM mutation suggests that WT RIM is unnecessary for the formation of the presynaptic active zone at the Drosophila NMJ. However, recent studies in mice indicate that rodent RIM is required for the proper localization of Ca\textsuperscript{2+} channels to presynaptic terminals (Kaeser et al., 2011). In RIM-deficient neurons, P/Q-type Ca\textsuperscript{2+} channel levels are reduced by ~40%, whereas the presynaptic active zone protein Bassoon remains unaffected (Kaeser et al., 2011). Interestingly, the PDZ domain of RIM is required for rescuing Ca\textsuperscript{2+} channel localization, suggesting an essential interaction between the RIM PDZ domain and Ca\textsuperscript{2+} channels, a domain that is fully deleted in our mutants. Thus, we asked whether the RIM mutation in Drosophila results in a similar decrease in Ca\textsuperscript{2+} channels at active zones. At WT NMJs, driving the GFP-tagged UAS–Cacophony, a calcium channel subunit, with the motor neuron DVGlu–Gal4 driver, results in the colocalization of Cacophony::GFP with Brp at active zones. The continued accu-
mutation of Cacophony at release sites requires Brp and is decreased in brp mutants (Kittel et al., 2006). However, in RIM<sup>Ex73</sup> mutants, Cacophony::GFP intensity at active zones is reduced, although Brp is unchanged (Fig. 3). Quantification of average Cacophony::GFP intensity per active zone, measured within regions of Brp signal, shows a decrease in Cacophony intensity of ~25% in RIM<sup>Ex73</sup>/Df compared with WT (Fig. 3B).

To ensure that reduced calcium channel levels are attributable to the absence of WT RIM, we coexpressed an untagged version of UAS–Cacophony::GFP in the RIM<sup>Ex73</sup>/Df background and measured Cacophony::GFP intensity at active zones. Expression of UAS–RIM increases Cacophony::GFP intensity to WT levels, rescuing the calcium channel phenotype of the RIM mutant (Fig. 3). Similar results were found for the other RIM<sup>Ex98</sup> excision and for each RIM allele over a genetically unrelated deficiency chromosome that deletes the RIM gene. Average Cacophony intensity per active zone is reduced in both homozygous RIM<sup>Ex73</sup> and RIM<sup>Ex73</sup>/Df(3R)ED5785 mutants (WT, 100 ± 4 a.u., n = 13; RIM<sup>Ex73</sup>, 60 ± 4 a.u., n = 14; RIM<sup>Ex73</sup>/Df, 72 ± 4 a.u., n = 13; p < 0.001 for WT vs both RIM<sup>Ex73</sup> and RIM<sup>Ex73</sup>/Df) and homozygous RIM<sup>Ex98</sup> and RIM<sup>Ex98</sup>/Df(3R)ED5785 mutants (WT, 100 ± 3 a.u., n = 12; RIM<sup>Ex98</sup>, 59 ± 2 a.u., n = 13; RIM<sup>Ex98</sup>/Df, 68 ± 1 a.u., n = 11; p < 0.001 for WT vs both RIM<sup>Ex98</sup> and RIM<sup>Ex98</sup>/Df). For both RIM mutants, average Cacophony::GFP intensity per active zone is similar when comparing homozygous RIM and RIM/Df(3R)ED5785 NMJs (RIM<sup>Ex73</sup>, p > 0.08; RIM<sup>Ex98</sup>, p > 0.3), suggesting that RIM<sup>Ex73</sup> and RIM<sup>Ex98</sup> behave as genetic nulls for this phenotype. Our results indicate that, in <i>Drosophila</i> WT RIM is necessary for the proper accumulation of calcium channels to active zones, although it is not required for localization of Brp to those same release sites. This is also consistent with the reduced amplitude of Ca<sup>2+</sup> transients observed by Ca<sup>2+</sup> imaging in RIM mutants (Müller et al., 2012).

**RIM mutation results in impaired synaptic vesicle release**

Immunohistochemical analysis suggests that fewer calcium channels are accumulated at RIM mutant active zones. How does this affect the function of RIM mutant NMJs? To assess the functional consequences of RIM mutation, we examined evoked and spontaneous neurotransmitter release in WT and RIM<sup>Ex73</sup>/Df(3R)ED5785 mutants in low external calcium (0.42 mM). There is a dramatic defect in synaptic transmission: the average evoked EJC amplitude in the RIM<sup>Ex73</sup>/Df(3R)ED5785 mutant is reduced by ~75% compared with WT (Fig. 4A, B). This physiological defect is partially rescued after transgenic expression of RIM with an approximate doubling of the average EJC amplitude when UAS–RIM is expressed in the RIM<sup>Ex73</sup>/Df(3R)ED5785 background (Fig. 4A, B). Partial rescue by transgenic RIM suggests
that the precise timing or levels of RIM expression are important for full function. In contrast to the reduction in EJC amplitude, the average amplitude of spontaneous mEJCs is not significantly different between WT, RIM mutant, and UAS–RIM rescue NMJs (Fig. 4A, C), indicating that there is no change in quantal size, the postsynaptic response to the release of a single vesicle. There is also no significant difference in frequency of spontaneous miniature events. Estimates of quantal content based on the direct method (EJC/mEJC) indicate that approximately one-quarter as many vesicles are released in RIM mutant as compared with WT after a single action potential, and this too is partially rescued by neuronal expression of UAS–RIM in RIM mutant (Fig. 4D).

Because calcium influx triggers the evoked release of synaptic vesicles, the reduction in the number of released vesicles in the RIM mutant is consistent with decreased calcium channel levels at release sites. To determine whether the defective vesicle release observed in the RIM mutant might be explained solely by reduced Ca\(^{2+}\) influx, we assayed evoked release in high external calcium. At 2 mM Ca\(^{2+}\), the average evoked EJCs in the RIM mutant are decreased by only 30% compared with WT [WT, 146.9 ± 13.4 nA, n = 7; RIM mutant, 99.9 ± 12.6 nA, n = 5; p < 0.05]. The more modest reduction in evoked EJC amplitude in high calcium concentrations (30% compared with 74% in low concentrations) suggests that increased calcium influx can partially restore vesicle release in RIM mutants. However, because high calcium does not fully rescue evoked release, other defects associated with vesicle release likely exist at RIM mutant NMJs.

In rodents, RIM not only promotes calcium channel localization but also regulates the size of the RRP of vesicles at the active zone (Han et al., 2011). Therefore, we assessed whether defective vesicle release in the RIM mutant might be explained solely by reduced Ca\(^{2+}\) influx, we assayed evoked release in high external calcium. At 2 mM Ca\(^{2+}\), the average evoked EJCs in the RIM mutant are decreased by only 30% compared with WT [WT, 146.9 ± 13.4 nA, n = 7; RIM mutant, 99.9 ± 12.6 nA, n = 5; p < 0.05]. The more modest reduction in evoked EJC amplitude in high calcium concentrations (30% compared with 74% in low concentrations) suggests that increased calcium influx can partially restore vesicle release in RIM mutants. However, because high calcium does not fully rescue evoked release, other defects associated with vesicle release likely exist at RIM mutant NMJs.

In rodents, RIM not only promotes calcium channel localization but also regulates the size of the RRP of vesicles at the active zone (Han et al., 2011). Therefore, we assessed whether defective vesicle release in the RIM mutant could be partially attributed to a reduced RRP. To estimate the size of the RRP in RIM mutant and WT animals, we used the method of cumulative amplitude analysis that has been successfully applied to the Drosophila NMJ (Schneeggenburger et al., 2002; Baldelli et al., 2007; Weyhersmüller et al., 2011). Stimulating at 40 Hz stimulation in 2 mM Ca\(^{2+}\) saline, we find that the size of the RRP is significantly reduced in the RIM mutant with UAS–RIM rescue [UAS–RIM rescue, 178 ± 39 vesicles, n = 5; RIM mutant, 178 ± 39 vesicles, n = 5; p < 0.05] (Fig. 4E), which likely accounts for some of the defect in evoked release observed at RIM mutant NMJs. There is also a slight decrease in the slope of the steady-state cumulative release indicating that the observed change in the RRP size is not the only defect in RIM, consistent with a change in the release probability and/or refilling rate. Moreover, as described above, the 15% decrease in active zone number in the RIM mutant may also contribute to the decreased EJC size. Altered release properties after RIM disruption are examined in detail in a companion manuscript (Müller et al., 2012).

**RIM is required for enhanced Ca\(^{2+}\) channel accumulation but not altered Brp distribution in the rab3 mutant**

We have shown previously that one role of Rab3 is to ensure that release components are distributed appropriately across release sites, potentially by influencing the proper nucleation of Brp at active zones (Graf et al., 2009). Because RIM is a known effector of Rab3 function in other organisms, and we show here that RIM protein can preferentially localize to release sites in Drosophila NMJs, we wanted to determine whether RIM is required for this...
function of Brp. The observation that Brp localizes normally in RIM mutants suggests that RIM is not required for Brp distribution across sites. However, RIM could initiate the altered distribution of release machinery in the Rab3 mutant. In C. elegans, enhanced expression of the Rim PDZ domain disrupts ELKS localization (Deken et al., 2005). In a related manner, the lack of appropriate Rab3–RIM interaction in the rab3\textsuperscript{opp} mutant may result in disrupted RIM function or localization and subsequently induce an altered Brp distribution. The observation that RIM::GFP localizes to the same subset of release sites as Brp in the rab3 mutant (Fig. 1D) is consistent with such a scenario.

We first asked whether full-length WT RIM is required for Rab3 localization at the NMJ. Rab3 staining in WT NMJs reveals overlapping synaptic vesicle-like and active zone-like patterns. We have shown previously that Rab is essential for the concentration of Rab3 at active zones (Graf et al., 2009). Is RIM similarly required for the active zone-like localization of Rab3? Staining for Rab3 in RIM\textsuperscript{Ex73}\textsl{Df(3R)ED5785} mutants reveals a Rab3 localization pattern that is similar to WT (Fig. 5). In the RIM mutant, Rab3 is still observed to colocalize with Brp in punctate-like concentrations (Fig. 5, inset) in regions of lower levels of synaptic vesicle-like staining. We see no difference in the fraction of NMJs showing punctate Rab3 staining between WT and the RIM mutant (p < 0.6), indicating that RIM is unnecessary for Rab3 localization to release sites.

Although WT RIM is not essential for Rab3 localization, is it required for the formation of the rab3 mutant active zone phenotype? To investigate this, we created rab3\textsuperscript{opp} and RIM\textsuperscript{Ex73} double mutants and compared Brp and GluRIII staining in WT, rab3\textsuperscript{opp} mutant, and rab3\textsuperscript{opp}–RIM\textsuperscript{Ex73} double mutant larvae. Brp distribution is similar between rab3\textsuperscript{opp} single mutants and rab3\textsuperscript{opp}–RIM\textsuperscript{Ex73} double mutants (Fig. 6A). In both single and double mutants, the majority of GluRIII clusters are unopposed to Brp (Fig. 6C), and average Brp puncta area is twice that of WT (Fig. 6D). rab3\textsuperscript{opp} single mutants and rab3\textsuperscript{opp}–RIM\textsuperscript{Ex73} double mutants are not significantly different in terms of the percentage of GluRIII clusters apposed by Brp (p > 0.7) or average Brp area (p > 0.5), indicating that RIM is not required for the altered distribution of Brp in the rab3\textsuperscript{opp} mutant.

Because RIM is necessary for appropriate Ca\textsuperscript{2+} channel accumulation at WT active zones, we asked whether RIM is similarly required for the enhanced concentration of Ca\textsuperscript{2+} channels observed at rab3\textsuperscript{opp} mutant release sites. Whereas average Caconophony::GFP intensity per Brp puncta region is significantly increased in the rab3\textsuperscript{opp} mutant compared with WT, RIM disruption in the rab3\textsuperscript{opp}–RIM\textsuperscript{Ex73} double mutant prevents enhanced Ca\textsuperscript{2+} channel accumulation, resulting in an average Caconophony::GFP intensity per active zone that is similar to WT (Fig. 6B, E). Because Brp is also involved in Ca\textsuperscript{2+} channel accumulation, the fact that Caconophony::GFP levels in the rab3\textsuperscript{opp}–RIM\textsuperscript{Ex73} double mutant are greater than in the RIM\textsuperscript{Ex73} single mutant is likely attributable to increased levels of Brp per active zone. Thus, although WT RIM is unnecessary for Rab3 localization at WT NMJs and Brp distribution in both WT and rab3 mutant NMJs, it is required for appropriate Ca\textsuperscript{2+} channel accumulation in both WT and rab3 mutants.

**Discussion**

We show that RIM localizes to active zones at the Drosophila NMJ and is required for the normal accumulation of Ca\textsuperscript{2+} channels. In RIM excision mutants, Ca\textsuperscript{2+} channel levels are significantly reduced at release sites, and there is a reduction in the size of the RRP of synaptic vesicles and a modest decrease in active zone number. As a result, RIM mutants show a dramatic impairment of evoked synaptic vesicle release. Conversely, the synaptic ultrastructure and localization and distribution of the release machinery protein Brp remains unaffected by the lack of WT RIM. Furthermore, the reduction in Ca\textsuperscript{2+} channels with no apparent effect on Brp is observed in both a WT background and in a rab3 mutant background that exhibits an altered distribution of active zone components across potential release sites.

Our findings are consistent with recent studies in rodents that show decreased accumulation of Ca\textsuperscript{2+} channels and reduced Ca\textsuperscript{2+} currents at synapses of conditional knock-out RIM mutant mice (Han et al., 2011; Kaeser et al., 2011), indicating an evolutionarily conserved role for RIM in regulating Ca\textsuperscript{2+} channel density at release sites. In rodents, this role requires the direct interaction of the PDZ domain of RIM with the C termini of N- and P/Q-type Ca\textsuperscript{2+} channels (Kaeser et al., 2011). In our studies, the entire central region of the Drosophila RIM gene is excised in both the RIM\textsuperscript{Ex73} and RIM\textsuperscript{Ex88} mutants, including the genomic
sequence that encodes for the PDZ domain. It is unknown whether an evolutionarily conserved interaction exists between the PDZ domain of Drosophila RIM and Cacophony; however, the complete removal of the PDZ domain in RIMEx73 and RIMEx98 ensures that such direct interactions cannot exist in the mutant, even if truncated versions of RIM protein are potentially expressed in RIMEx73 and RIMEx98. Thus, Ca\(^{2+}\) channel accumulation is likely reduced in RIMEx73 and RIMEx98 mutants because of the inability of RIM and Ca\(^{2+}\) channels to directly interact.

Several proteins have now been identified in Drosophila as regulators of Ca\(^{2+}\) channel accumulation, including Brp and the recently reported Drosophila RIM-binding protein (DRBP) (Kittel et al., 2006; Wagh et al., 2006; Liu et al., 2011). Disruption of any one of the three proteins fails to completely disrupt Ca\(^{2+}\) channel accumulation in the rab3 mutant. A, Images of NMJs from WT, rab3 mutant (rab3\(^{396}\)/rab3\(^{396}\)), and rab3–RIM double-mutant (rab3\(^{396}\)/rab3\(^{396}\); RIMEx73/RIMEx73) larvae stained with α-Brp (red) and α-DGluRIII (green). Scale bar, 2 μm. B, Images of NMJs expressing the Ca\(^{2+}\) channel subunit Cacophony::GFP (green) and costained with α-Brp (red) from WT (dyglut\(^{NMJX}\); Gal4\(^{UAS}\)–Cacophony::GFP\(^{+/+}\)), the rab3 mutant (dyglut\(^{NMJX}\); Gal4\(^{UAS}\)–Cacophony::GFP\(^{-/+}\)), and the rab3–RIM double mutant (dyglut\(^{NMJX}\); Gal4\(^{+/+}\); UAS–Cacophony::GFP, rab3\(^{396}\)/rab3\(^{396}\); RIMEx73/RIMEx73). Scale bar, 2 μm. C, Histogram shows the average percentage of DGluRIII clusters apposed to Brp puncta per NMJ for the genotypes listed in A. WT, n = 16 NMJs; rab3 mutant, n = 14 NMJs; rab3–RIM double mutant, n = 13 NMJs; *p < 0.001 versus WT. D, Histogram shows the average area of individual Brp puncta for the genotypes listed in A. n = 10 NMJs for all genotypes; *p < 0.001 versus WT. E, Histogram shows the average intensity of Cacophony::GFP per Brp puncta area for the genotypes listed in B. WT, n = 17 NMJs; rab3 mutant, n = 16 NMJs; rab3–RIM double mutant, n = 19 NMJs; *p < 0.001 versus WT and rab3–RIM double mutant.
channel localization to release sites, suggesting that they may play partially overlapping roles. Evidence in Drosophila, rodents, and C. elegans indicates that all three proteins directly interact with Ca$^{2+}$ channels (Hibino et al., 2002; Kiyonaka et al., 2007; Fouquet et al., 2009; Kaeser et al., 2011; Liu et al., 2011) but also interacting with each other and several other components to form the active zone cytomatrix (Wang et al., 2000; Betz et al., 2001; Coppola et al., 2001; Ohtsuka et al., 2002; Schoch et al., 2002; Wang et al., 2002; Deken et al., 2005; Dulubova et al., 2005). Consistent with protein binding studies, we show that GFP-tagged RIM concentrates at release sites and colocalizes with Brp. Interestingly, whereas DRBP levels are reduced in brp mutants (Liu et al., 2011), RIM::GFP localizes in an active zone-like pattern even in the absence of Brp. This finding is consistent with studies in C. elegans (Deken et al., 2005) and suggests that Brp is unnecessary for the localization of RIM to release sites. Furthermore, Brp localization to active zones is maintained in both brp mutant mice and indicates that multiple functions of RIM are evolutionarily conserved (Han et al., 2011; Kaeser et al., 2011).

**RIM is not an effector of Rab3 for the control of protein composition across release sites**

We originally hypothesized that RIM may act as an effector of Rab3 to regulate active zone protein composition across release sites. RIM has been studied previously in rodents and C. elegans as an important effector in the Rab3-mediated docking of synaptic vesicles to active zones during the synaptic vesicle cycle (Koushika et al., 2001; Südhof, 2004). The active zone localization and extensive interactions of RIM with multiple active zone proteins place it in an attractive position to potentially mediate this newly described role for Rab3 in the control of active zone protein composition (Graf et al., 2009). Moreover, our observation that RIM can localize in an active zone-like manner in the absence of Brp suggests that RIM can potentially act upstream of other presynaptic cytomatrix proteins to regulate the formation of the presynaptic release machinery. Brp is a central component of Drosophila active zones and is dramatically redistributed in rab3 mutant NMJs, so we would expect a downstream effector of Rab3 to control Brp localization to active zones. However, analysis of Brp in the Rab3 mutant background has no effect on the altered distribution of Brp that results after rab3 disruption, demonstrating that RIM is not involved in the molecular mechanisms that control Brp re-distribution in the rab3 mutant. The reduction of Ca$^{2+}$ channels in the rab3--RIM double mutant does suggest that RIM may be required for the enhanced efficacy of “super active zones” formed in the rab3 mutant (Graf et al., 2009; Peled and Isacoff, 2011); however, the molecular mechanism by which Rab3 controls the general distribution of active zone components across release sites remains unknown.

**References**


