Nanoscaled RIM clustering at presynaptic active zones revealed by endogenous tagging

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Chemical synaptic transmission involves neurotransmitter release from presynaptic active zones (AZs). The AZ protein Rab3-interacting molecule (RIM) is important for normal Ca2+–triggered release. However, its precise localization within AZs of the glutamatergic neuromuscular junctions of Drosophila melanogaster remains elusive. We used CRISPR/Cas9-assisted genome engineering of the rim locus to incorporate small epitope tags for targeted super-resolution imaging. A V5-tag, derived from simian virus 5, and an HA-tag, derived from human influenza virus, were N-terminally fused to the RIM zinc finger. Whereas both variants are expressed in co-localization with the core AZ scaffold Bruchpilot, electrophysiological characterization reveals that AP-evoked synaptic release is disturbed in rimV5

Introduction


At the Drosophila melanogaster neuromuscular junction (NMJ), RIM promotes VGCC accumulation within the AZ, controls the readily releasable pool of synaptic vesicles, and is essential for presynaptic homeostatic plasticity (Graf et al, 2012; Müller et al, 2012; Paul et al, 2022), and thus the dynamic regulation of synaptic strength (Davis & Müller, 2015). Whereas remarkable reorganization of RIM during synaptic plasticity was described in cultured murine neurons (Tang et al, 2016; Müller et al, 2022), the nanoscale organization of RIM at Drosophila AZs, remains unclear. Previous confocal and stimulation emission depletion imaging relied on overexpression or the endogenous expression of GFP-fused constructs (Graf et al, 2012; Petzoldt et al, 2020). During the last decade, various tools for the generation of genetically marked constructs emerged, offering new labeling strategies for super-resolution imaging.

Here, we used the clustered regularly interspaced short palindromic repeats (CRISPR)/CRISPR-associated protein 9 (Cas9) system (Bassett et al, 2013; Gratz et al, 2013, 2014; Kondo & Ueda, 2013; Yu et al, 2013; Port et al, 2014) to introduce small epitope tags into RIM to enable its precise localization within presynaptic AZs. We inserted either a V5-tag derived from the P and V proteins of the simian virus 5 (GPIPNPLLGLDST, Hanke et al, 1992) or an HA-tag derived from the human influenza virus (YPYDVPDYA) into the endogenous rim ORF to tag the N-terminal end of the zinc finger domain (rimV5

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extension, neurotransmitter release. Electrophysiological characterizations revealed that the HA-tag, in contrast to the V5-tag, leaves spontaneous and AP-evoked synaptic release and the expression of presynaptic plasticity undisturbed. We found efficient expression of RIMV5−Znf and RIMHA−Znf in distinct co-localization with the AZ scaffold protein Bruchpilot (Brp) in the peripheral and central nervous systems of Drosophila larvae. Thus, we investigated the nanotopology of RIMHA−Znf at the AZ using two-color direct stochastic optical reconstruction microscopy (dSTORM; Heilemann et al, 2008; van de Linde et al, 2011) in combination with hierarchical density-based spatial clustering of applications with noise (HDBSCAN). Our super-resolution approach reveals information about the RIMHA−Znf nano-assembly at the Drosophila AZ into distinct clusters of ~130 nm² size in ~120 nm distance from the AZ center. Furthermore, these RIMHA−Znf clusters change in response to an acute homeostatic challenge by shrinking, increasing in localization density, and in absolute numbers.

Results
CRISPR/Cas9-engineering of rim

Previous RIM imaging at the Drosophila NMJ utilized overexpression or endogenous expression of GFP-fused proteins (Graf et al, 2012; Petzoldt et al, 2020). In the latter, a recombinase-mediated cassette exchange-derived line based on the Minos-mediated integration cassette collection (Nagarkar-Jaiswal et al, 2015) targeting only one of the 14 predicted splice variants was employed (Petzoldt et al, 2020). Here, we genomically engineered alleles of the single Drosophila rim gene to investigate the population of RIM at AZ scaffolds at levels under endogenous cis- and trans-regulatory control (Fig 1A). We combined CRISPR/Cas9-assisted genome editing (Gratz et al, 2014) and φC31-mediated recombine-mediated cassette exchange (Huang et al, 2009) as previously applied to expedite generation of alleles encoding RIM C2A domain mutations (Paul et al, 2022). To this end, we targeted a genome fragment spanning exons 3–7 that cover the coding sequence for the Rab3-interacting/zinc finger (Znf) domain of RIM and replaced it by a rescue genomic fragment with the full removed sequence (rimrescue-Znf), or genomic fragments with additional coding sequences for the V5- or HA-tag before the common start codon (rimV5-Znf and rimHA-Znf, respectively, Fig 1B), leading to N-terminally tagged fusion proteins (Fig 1C).

Baseline synaptic transmission at rimV5−Znf and rimHA−Znf terminals

Next, we performed two-electrode voltage clamp recordings for evaluation of spontaneous and evoked synaptic transmission at larval NMJs of the epitope-tagged rim variants (Fig 2 and Tables S1 and S2). We recorded miniature excitatory postsynaptic currents (mEPSCs) to examine spontaneous release in wt and rimrescue−Znf third instar larvae to control for normal synaptic release in the latter. In addition, mEPSCs of rimV5−Znf and rimHA−Znf larvae were compared with rimrescue−Znf (Fig 2A, all homozygous). mEPSC amplitudes and the frequency of spontaneous fusion events were unchanged at both rimV5−Znf and rimHA−Znf NMJs (Fig 2B). Next, we measured evoked excitatory postsynaptic currents (eEPSCs) in response to nerve stimulation (Fig 2C). eEPSC amplitudes were unaltered in rimrescue−Znf compared with wt (Fig 2D). Interestingly, eEPSC amplitudes were significantly decreased in rimV5−Znf compared with
rim^{rescue-Znf}, whereas they were unchanged in rim^{HA-Znf} (Fig 2D), indicating that the V5-tag but not the HA-tag interferes with evoked synaptic release. We also tested whether insertion of the tags alters synaptic short-term plasticity but found paired pulse ratios unchanged at both rim^{V5-Znf} and rim^{HA-Znf} NMJs (Fig 2D). Taken together, these data show that both spontaneous and evoked synaptic transmissions are intact in rim^{rescue-Znf} and rim^{HA-Znf} variants. Furthermore, as evoked synaptic release is significantly decreased in rim^{V5-Znf} but not in rim^{HA-Znf}, the HA-tag is an appropriate tool for further assessment of structure–function relationships at the Drosophila NMJ.

Undisturbed presynaptic homeostatic potentiation (PHP) at rim^{HA-Znf} NMJs

RIM is required for PHP by modulation of the readily releasable vesicle pool (RRP, Müller et al, 2012). Thus, we probed if our genetically engineered rim variants carrying an epitope tag at the N-terminal zinc finger domain still exhibit PHP at normal levels. To test if the HA-tagged RIM is still functional, we measured the electrophysiological response to an acute homeostatic challenge using Philanthotoxin (PhTx) in rim^{HA-Znf} animals compared with rim^{rescue-Znf} larvae (Fig 2E and Tables S3 and S4). We found that upon PhTx treatment, rim^{HA-Znf} showed the same increase in quantal content and thus evoked EPSC restoration as rim^{rescue-Znf}. This indicates that rim^{HA-Znf} NMJs exhibit unperturbed PHP. We conclude that genomic HA-tag insertion into the endogenous rim ORF leaves presynaptic plasticity intact at larval Drosophila NMJs.

Expression of tagged rim alleles in the Drosophila nervous system

After verification that rim^{HA-Znf} but not rim^{V5-Znf} larvae show normal synaptic release upon AP-evoked stimulation and at homeostatic challenge, we investigated the expression of RIM with the N-terminally fused epitope tags within the Drosophila nervous system.
Suitable antibodies for the detection of RIM at presynaptic Drosophila AZs are still lacking. Thus, implementation of reliable RIM imaging in the fly is highly demanding, especially regarding the AZ nano-arrangement. To first probe the overall expression of RIMHA−Znf in the Drosophila central and peripheral nervous systems, we performed immunostainings using a monoclonal antibody against HA for detection of the N-terminal HA-tag (see the Material and Methods section) and a well-characterized, highly specific monoclonal antibody BrpNc82 mapping to the C-terminal region of Brp (Kittel et al., 2006; Fouquet et al., 2009; Mrestani et al., 2021; Fig 3). RIMHA−Znf was strongly expressed in the larval central nervous system (Fig 3A). We also detected considerable co-expression of RIMHA−Znf and BrpNc82 at third instar larval NMJs (Fig 3B and C). In addition, co-expression of RIMV5−Znf and BrpNc82 at NMJs was observed (Fig S1), although, in rimV5−Znf animals, baseline synaptic transmission was disturbed (Fig 2C and D). We conclude that RIMHA−Znf is ubiquitously expressed in the fly central and peripheral nervous systems and specifically co-localizes with the AZ scaffold protein. Thus, super-resolution analysis of RIMHA−Znf at pre-synaptic AZs is feasible. To analyze if NMJ morphology itself is altered in rimHA−Znf animals, we performed immunostainings of α-HRP and BrpNc82 in rimHA−Znf and rimrescue−Znf (Fig 3D and Table S5). The number of BrpNc82 puncta per NMJ and NMJ area were unaltered, however, the number of boutons per NMJ slightly decreased in rimHA−Znf. We conclude that the overall NMJ morphology remains mostly unaltered in the tagged rim variant.

Identification of RIMHA−Znf clusters at presynaptic AZs

Next, we performed two-color dSTORM localization microscopy of RIMHA−Znf at larval NMJs (Fig 4 and Table S6; Heilemann et al., 2008;
Acute PHP results in compaction and addition of RIM<sup>HA−Znf</sup> subclusters

The induction of presynaptic homeostasis is associated with structural reorganization of presynaptic AZs (Weychersmüller et al., 2011; Böhme et al., 2019; Mrestani et al., 2021). As RIM is essential for PHP expression (Müller et al., 2012), we wondered whether nanoscaled reorganization of RIM<sup>HA−Znf</sup> occurs at the Drosophila NMJ (Fig 5 and Tables S6 and S7). Thus, we compared Brp<sup>HCA2</sup> and RIM<sup>HA−Znf</sup> localization data using HDDBSCAN-based algorithms in PhTx-treated preparations (phTx, Fig 5A) and DMSO controls (ctrl). We found no difference in RIM<sup>HA−Znf</sup> localization numbers per SC between phtx and ctrl, however, acute PHP reduced RIM<sup>HA−Znf</sup> SC areas and increased localization density (Fig 5B). Interestingly, the number of SCs per AZ along with the total number of RIM<sup>HA−Znf</sup> localizations per AZ increased in phtx (Fig 5C), corroborating earlier results from cultured murine neurons (Müller et al., 2022). The radial distance between SC c.o.m.s and the AZ c.o.m. was unchanged in phtx and the total AZ area occupied by RIM<sup>HA−Znf</sup> remained the same (Fig 5C and Table S6). We tested if the slightly decreased AZ circularity in phtx

van de Linde et al, 2011; Löscherberger et al, 2012; Mrestani et al, 2021; Paul et al, 2022; Dannhäuser et al, 2022). Using a combination of Brp<sup>HCA2</sup> and a monoclonal antibody against the HA-tag for detection of RIM<sup>HA−Znf</sup> we observed distinct co-localization of both epitopes at presynaptic AZs of type Ib boutons (Fig 4A). Application of HDDBSCAN and Ripley analyses as established in previous work (Mrestani et al, 2021; Dannhäuser et al, 2022) extracted individual RIM<sup>HA−Znf</sup> subclusters (SCs) with diameters of ~13 nm (Fig 4B and C). Using alpha shapes for area determination (Mrestani et al, 2021), we obtained RIM<sup>HA−Znf</sup> SCs per AZ (n = 893 AZs from 18 NMJs in nine animals) in all AZs without selection according to AZ circularity (see the Material and Methods section). Inset in the left panel highlights the range between 0 and 400 nm<sup>2</sup> SC area. Solid red line indicates median, dashed red lines, 25<sup>th</sup> and 75<sup>th</sup> percentiles. Scale bars in (A) 1 μm, in (C) 100 nm.

**Figure 4.** RIM<sup>HA−Znf</sup> clusters at presynaptic AZs. (A) Two-channel dSTORM localizations for a rim<sup>HA−Znf</sup> type Ib bouton. Left: RIM<sup>HA−Znf</sup> stained with α-HA antibody and Alexa Fluor647 conjugated F(ab')2 fragments (magenta). Middle: overlay with Brp<sup>NC82</sup> labeled with Alexa Fluor 532-conjugated IgGs (green). (C) Asterisk marks enlarged AZ in (C). Right: RIM<sup>HA−Znf</sup> localizations from left panel with all localizations with Euclidian distance >20 nm to Brp localizations removed. The removed signal is considered noise. Individual RIM<sup>HA−Znf</sup> subclusters (SCs) were extracted by HDDBSCAN and assigned to nearest AZs by color. Colored lines indicate alpha shapes used for area determination. The center of mass (c.o.m.) of the corresponding AZ (x) is indicated. Dashed red line shows the Euclidian distance between the AZ c.o.m and an SC c.o.m., referred to as radial distance. Parameters in HDDBSCAN were: minimum cluster size = 100 localizations, minimum samples = 25 localizations for Brp<sup>NC82</sup>; minimum cluster size = 2 localizations, minimum samples = 2 localizations for RIM<sup>HA−Znf</sup>, α-value Brp<sup>NC82</sup> = 800 nm<sup>2</sup>, α-value RIM<sup>HA−Znf</sup> = 300 nm<sup>2</sup>; exclusion criteria for Brp<sup>NC82</sup> clusters were area ≤0.03 μm<sup>2</sup> and ≥0.3 μm<sup>2</sup>. (D) Distributions of RIM<sup>HA−Znf</sup> SC area (11,094 SCs from 18 NMJs in nine animals) and the number of RIM<sup>HA−Znf</sup> SCs per AZ (n = 893 AZs from 18 NMJs in nine animals) in all AZs without selection according to AZ circularity (see the Material and Methods section). Inset in the left panel highlights the range between 0 and 400 nm<sup>2</sup> SC area. Solid red line indicates median, dashed red lines, 25<sup>th</sup> and 75<sup>th</sup> percentiles. Scale bars in (A) 1 μm, in (C) 100 nm.
influences structural parameters (Table S6, Mrestani et al, 2021). However, all relative changes between experimental groups were present in filtered AZs in planar view indicated by AZ circularity ≥ 0.6 (Tables S6 and S7). To further control the robustness of our findings, we established an analysis routine alternative to our previous algorithm (Dannhäuser et al, 2022), now relying on HDBSCAN to account for noise. Single-channel HDBSCAN analysis of RIM^MA-Znf (Fig S2A and B) delivers less intuitive segmentation opposed to Brp^Nc82 (Fig S2C, compare Fig 1B in Mrestani et al [2021]). However, it accounts for noise in the data, as an alternate way to denoising by distance to the Brp^Nc82 signal (compare the Material and Methods section and Dannhäuser et al [2022]). Furthermore, after AZ assignment (Fig S2D), RIM^MA-Znf SCs outside the AZ are accessible for quantification (Fig S2E and F). Whereas analysis of intrasyaptic RIM^MA-Znf SCs confirmed compaction during PHP, no differences between ctrl and phtx were detectable for extrasynaptic SC populations (Fig S2G and Table S8). Interestingly, extrasynaptic SCs displayed similar localization numbers, increased areas, and lower localization densities opposed to their intrasyaptic counterparts (Fig S2G), implying a nanotopological differentiation of these two populations. To address whether increased RIM^MA-Znf protein per AZ during homeostasis (Fig 5C) arises from recruitment from the AZ vicinity, we quantified the effect of PHP on RIM^MA-Znf SC numbers and localizations in the extrasynaptic SC population in 400 nm distance around the AZ and found no difference, however, RIM^MA-Znf SC radial distance was slightly increased (Fig S2H).

Strikingly, analyzing these parameters for intrasyaptic SCs using the two different denoising approaches delivered identical results (Figs 5C and S2I). Lastly, we employed a second level HDBSCAN analysis to investigate RIM^MA-Znf superclusters (SpCs, Fig 5D, Dannhäuser et al, 2022). Remarkably, in this analysis, the percentage of SCs clustered into SpCs was increased after PHP (Fig 5E), and the fraction of AZs displaying superclustering at all (45.6% versus 53.32% in ctrl and phtx, respectively). Furthermore, nearest neighbor analysis revealed decreased distances between SC c.o.m.s in phtx (Fig S3A and B), suggesting enhanced clustering, that is, compaction of RIM^MA-Znf during the homeostatic challenge. In addition, in both ctrl and phtx most AZs contained about 2–3 SpCs and about 4 SCs per SpC which displayed similar sizes (Figs 5F and S3C and D). In summary, PHP leads to more RIM^MA-Znf SCs per AZ with increased localization density. Furthermore, the proportion of SCs clustered into SpCs is larger in phtx and distances between individual SCs are decreased, suggesting compaction of RIM^MA-Znf during a homeostatic challenge.

**Discussion**

We used CRISPR/Cas9-assisted genome engineering of D. rim and fused two established epitope tags N-terminally to the zinc finger domain (Fig 1). We show that both rim variants RIM^V5-Znf and
RIM\(^{HA-Znf}\) are efficiently expressed at presynaptic AZs in co-localization with Brp (Figs 3 and S1). Using electrophysiology and PhTx to induce PHP, we demonstrate that release is diminished in rim\(^{V5-Znf}\), whereas, baseline synaptic transmission and PHP remain intact in rim\(^{HA-Znf}\) (Fig 2). We determine the localization of RIM within the AZ nano-scaffold in Drosophila, applying a combination of two-channel local microscopy and HDBSCAN analysis (Figs 4, S2, and S3). We detect ~10 RIM\(^{HA-Znf}\) clusters per AZ of ~130 nm\(^2\) size in ~120 nm distance from the AZ center, which compact and increase in numbers during acute PHP (Figs S5 and S4A and B).

Endogenous epitope tagging of rim

Previous RIM imaging at AZs of Drosophila NMJs utilized endogenous Minos-mediated integration cassette-based tagging of a single splice variant (Petzoldt et al, 2020) or overexpression of a full-length N-terminally GFP-tagged fusion protein (Graf et al, 2012). While establishing a method for deciphering RIM nanoscale arrangement at the AZ, we found disturbed synaptic transmission in the GFP-tagged fusion protein (data not shown). Through the application of a previously introduced genomic editing platform (Paul et al, 2022), we fused a V5-tag inserted N-terminally to the RIM C2A domain which did not yield specific imaging results (data not shown), possibly through lack of antibody accessibility of the epitope at this protein position. Therefore, we decided to focus on tag insertion at RIM’s N-terminus, because this position principally worked in earlier studies (Graf et al, 2012). Furthermore, only the Rab3-binding domain is present in all D. rim splice variants, enabling visualization of the whole rim population. To reduce the possibility to disturb the RIM–Rab3 interaction via RIM’s zinc finger domain through an adjacent fusion, we resorted to smaller V5 and HA epitope tags. Strikingly, whereas both peptide tags are composed of only a few amino acids, we found differences in their effect on synaptic transmission. In animals expressing the V5-tagged rim variant, we observed reduced evoked release, whereas RIM\(^{HA-Znf}\) displayed intact neurotransmission and PHP. Reduced eEPSC amplitudes in rim\(^{V5-Znf}\) animals fit with disturbed vesicle recruitment via a Rab3 interaction but the HA-tag insertion at the same position without discernible effects on synaptic function argues against these assumptions. How can the effect of the V5-tag on rim be explained?

Both epitope tags are small and have a low molecular weight (V5: 14 amino acids, ~1.4 kD; HA: 9 amino acids, ~1.1 kD), but differ in their substituents and charged residues and, accordingly, in their isoelectric points (V5: pI = 5.84, 1 positive and 1 negative residue; HA: pI = 3.56, 2 negative residues). V5 contains more amino acids with aliphatic substituents compared with HA, which is composed of more aromatic residues. In combination with the higher isoelectric point this leads to less hydrophilicity of the V5-tag which may interfere with folding of the Rab3/Znf-domain and, thus, RIM function. In contrast, the HA-tag contains two negatively charged amino acids and, because of the aromatic residues that can form π–π interactions, the HA peptide may adopt a more compact conformation. These differences can lead to an increased hydrophilicity of the HA-tag, with less steric interactions with the protein, preserving RIM function (Fig 2). In summary, our results indicate that not only epitope tag position and size but also other, yet ill-defined properties of the tag, profoundly influence RIM. At AZ scaffolds, RIM recruits presynaptic vesicles via Rab3 interaction to VGCCs through multiple direct and indirect interactions with its C-terminal domains (Wang et al, 1997; Kaeser et al, 2011). Assuming a central VGCC arrangement at Drosophila AZs (Hallermann et al, 2010; Ghelani & Sigrist, 2018; Ghelani et al, 2023), the RIM C-terminus should localize closer to AZ centers than the ~120 nm reported for N-terminal RIM SCs in the present study. Thus, simultaneous tagging and mapping of RIM N- and C-termini might be informative. However, the different D. rim gene products display considerable structural heterogeneity at the C-terminus with only a fraction possessing the C-terminal C\(_2\)B domain, inevitably leading to deviant variant subgroups during simultaneous imaging. Another tagging option is the relative mapping of RIM’s N- and C-termini to VGCCs. However, the two-color dSTORM approach as presented here using Alexa Fluor532 to establish Brp as reference signal does not meet the high-resolution requirements for such mapping studies. The described technical limitations might be overcome with more advanced dyes and more favorable photophysics and/or applying spectral demixing single-molecule localization microscopy of spectrally close far-red dyes (Lehmann et al, 2016; Wang et al, 2022).

Rapidly reorganizing RIM nanoclusters during homeostatic plasticity

RIM is required for many forms of presynaptic plasticity, including short-term plasticity, long-term potentiation, and PHP (Castillo et al, 2002; Schoch et al, 2002; Fourcaudot et al, 2008; Pelkey et al, 2008; Müller et al, 2012). PHP is among the best characterized plasticity patterns of the Drosophila NM; however, despite emerging evidence for remarkable reorganization of RIM nanoclusters during synaptic plasticity at mammalian synapses (Tang et al, 2016), structural reorganization of RIM at Drosophila synapses has not been investigated yet. We report increased numbers of RIM SCs in PhTx-induced acute PHP at Drosophila NMJ AZs (Fig 5C), in agreement with earlier findings from cultured hippocampal neurons (Müller et al, 2022). Increased RIM amounts for enhanced vesicle traffic in PHP appear plausible regarding its role in vesicle priming and Ca\(^{2+}\)-triggered release (Schoch et al, 2006; Deng et al, 2011) mediated through liquid–liquid phase separation (Wu et al, 2019, 2021). However, the changes of AZ proteins in PHP appear to be differentially regulated. In earlier studies, we and others demonstrated compaction of Brp, RBP, and Unc-13 and VGCCs at AZs in PHP (Mretoni et al, 2021; Dannhäuser et al, 2022; Ghelani et al, 2023). Whereas SC areas of Brp, RBP, and RIM decreased and localization density consecutively increased, SC area and localization density of Unc-13 remained unchanged. Furthermore, Brp and Unc-13 SCs move towards the AZ center in PHP (reduced radial distance). This is a second level of compaction, and it appears that in addition to SC compaction (for some proteins), the entire AZ is compacted in PHP. Remarkably, RIM is the only epitope so far with more SCs per AZ and therefore higher overall localization numbers per AZ in PHP, consistent with a mechanism of protein recruitment to the AZ, possibly involving the kinesin-associated axonal cargo machinery (Goel et al, 2019) or altered proteostasis during homeostasis (Baccino-Calace et al, 2022). The decreased area and enhanced molecule density of RIM\(^{HA-Znf}\) SCs described here (Figs S5B and S4).
fits well with the AZ compaction pattern implicated in structurally defining high release probability terminals (He et al., 2022 Preprint). In an earlier study, we determined a higher-level organization of Unc-13 SCs into ~2–3 SpCs per AZ matching the number of docked vesicles (Böhme et al., 2016). Protein level changes in PHP are not observed for Brp and Unc-13. Hence, increased formation of vesicles (Böhme et al., 2016). Protein level changes in PHP are generally defined by relatively fix Unc-13 SCs, whereas active release sites are discernable by additionally recruited RIM SCs, supporting increased vesicle traffic.

**Perspectives on RIM nanotopology at AZ scaffolds**

We provide a nanoscale quantification of the crucial AZ component RIM in *Drosophila*, utilizing a novel endogenously tagged genetic tool in combination with two-color dSTORM and HDBSCAN algorithms. In principle, it is possible to retrieve quantitative information on protein numbers from dSTORM data (Löschberger et al., 2012; Ehmann et al., 2014). Using Alexa Fluor 647 to quantitatively assess Brp distribution at *Drosophila* AZs, a conversion factor of 0.134 ± 0.028 (SEM) between localizations and molecules was determined (Ehmann et al., 2014). Regarding the similar staining and imaging conditions in the present study, this factor can be used for a rough approximation of RIM molecules at the AZ scaffold. Accordingly, six localizations per RIM SC may translate in ~1–2 molecules. RIM SCs are smaller than Brp, RBP, and Unc-13 SCs and at the resolution limit of our imaging approach; however, we still observe compaction in PHP. In addition, it appears promising to use this imaging strategy for studying RIM structure and function in phasic type IS versus tonic type Ib boutons. The HA-tag used in this study is incorporated in all 14 variants present in *Drosophila*; however, recent work suggested differential isoform expression in these bouton types (Jetti et al., 2023 Preprint). Finally, considering the remarkable structural reorganization of RIM (Yao et al., 2007; Tang et al., 2016; Müller et al., 2022), probing RIM redistribution in the context of human disease relevant point mutations, for example, the arginine to histidine substitution in the 3C helix of the CA domain causing human CORD7 syndrome (Paul et al., 2022), will be informative.

**Materials and Methods**

**Fly stocks**

*Drosophila melanogaster* were raised on a standard cornmeal and molasses medium at 25°C. Male third instar larvae were used for experiments.

**Fly stocks generated in this study**

Internal stock IDs indicated in brackets after the genotype:

- rim<sup>Rob3/Znf</sup>, DsRed<sup>+</sup>; +; rim<sup>Rob3/Znf</sup> attP(DsRed<sup>+</sup>/TM3, Sb (RIM76, RIM98)
- rim<sup>Rob3/Znf</sup>, DsRed<sup>+</sup>; +; rim<sup>Rob3/Znf</sup> attP(DsRed<sup>+</sup>/TM3, Sb (LAT471, LAT473)

**RIM nanoclusters at presynaptic active zones**

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acid V5-tag (GGTAAACCATCCCTACCCCTCCTCGGTCGATATTAGC-CCGGGGCGCCGC; sequence in 5’–3’ order, additional XmaI and NotI sites are underlined) directly 5’ to the rim start codon, giving rise to the fusion of the amino acids MGKPIPNPLLGLDSTPGGGR to the RIM N-terminus. pTLB21 was BstBI/XhoI-digested to release the 1.1 kb fragment which was subsequently ligated into a 10.4-kb fragment of BstBI/XhoI-cut pLM6, generating the rim\textsuperscript{[V5-2nd]} vector pLM11.

rim\textsuperscript{[HA-2nd]} attrB plasmid

Primers am\_212F and am\_213R were annealed and ligated into the multiple cloning site of the AvrII/NcoI-digested subcloning vector pMCS5, generating plasmid pAM60-containing BstBI/XhoI sites. A 1.1-kb BstBI/XhoI fragment of pLM11 was ligated into BstBI/XhoI-digested pAM60, giving rise to plasmid pAM61. A 0.2-kb genomic DNA fragment containing a HindIII site located 5’ to the rim start codon and the subsequent 5’-UTR followed by an additional start codon and the coding sequence of the 9 amino acid HA-tag (TACCCT-TACCGACTCTCCCCCCCCTGGGGGGGGGGCGGCCGC; sequence in 5’–3’ order, additional XmaI and NotI sites are underlined, results in the fusion of the amino acids MYPYDVPDYAPGGGR N-terminal to the RI sites are underlined) directly 5’ to the rim start codon, and the subsequent 5’-UTR followed by an additional start codon and the coding sequence of the 9 amino acid HA-tag (TACCCT-TACCGACTCTCCCCCCCCTGGGGGGGGGGCGGCCGC; sequence in 5’–3’ order, additional XmaI and NotI sites are underlined, results in the fusion of the amino acids MYPYDVPDYAPGGGR N-terminal to the

CRISPR targeting

All transgenesis steps were performed at BestGene Inc. The gRNA plasmids (pMH1, pMH2) and the HDR donor plasmid (pMH10) were injected into embryos of a Drosophila strain with germline expression of vasa-Cas9 (Gratz et al., 2014), producing the rim\textsuperscript{[A2b3/2nd]} allele. Correct transgene incorporation was confirmed by sequencing of PCR fragments covering breakpoints between genomic/transgenic DNA amplified from genomic DNA of respective adult transgenic flies and across the deleted fragment. The 3p3X-DsRed\textsuperscript{red} transformation marker was removed by expressing a germline Cre source and confirmed by PCR genotyping. Subsequent insertion of the different attB transgenes (pLM6, pLM11, pAM62) into rim\textsuperscript{[A2b3/2nd]} by pC31-mediated transgenesis followed by Cre-driven removal of the mW\textsuperscript{+} selection marker was performed by BestGene Inc.

Fixation, staining, and immunofluorescence

For immunofluorescence imaging of larval NMJs and ventral nerve cords (VNCs), larvae were dissected in ice-cold hemolymph-like solution (HL-3, Stewart et al., 1994), fixed with 4% PFA in PBS for 10 min, and blocked for 30 min with PBT (PBS containing 0.05% Triton X-100, Sigma-Aldrich) including 5% normal goat serum (Dianova). Primary antibodies were added for overnight staining at 4°C. After two short and three 20-min-long washing steps with PBT, preparations were incubated with secondary antibodies for 3 h at room temperature, followed by two short and three 20-min-long washing steps with PBT. Preparations were kept in PBS at 4°C until imaging.

All NMJ data were obtained from abdominal muscles 6/7 in segments A2 and A3. Directly compared data were obtained from larvae stained in the same vial and measured in one imaging session.

Confocal microscopy and structured illumination microscopy (Apotome, SIM)

Preparation, fixation, and antibody staining were performed as described above. Primary antibodies were used in the following concentrations: mouse monoclonal α-Brp (Brp\textsubscript{N/C2}, 1:100; AB_2314866; Developmental Studies Hybridoma Bank), rabbit monoclonal α-HA (1:500; C294F catalog #3724; Cell signaling technology), and mouse monoclonal α-V5 (1:100; R960-25; Invitrogen). Secondary antibodies were used in the following concentrations: goat α-rabbit conjugated Alexa Fluor 488 (1:500; A-11088; Invitrogen), goat α-mouse conjugated Alexa Fluor 488 (1:250 for SIM imaging and 1:500 for Apotome imaging. A-32723, Invitrogen), goat α-mouse conjugated Cy3 (1:500, RRID: AB_2338690; Jackson ImmunoResearch). Directly conjugated antibodies were incubated together with secondary antibodies in the following concentrations: goat α-horseradish peroxidase (α-HRP) labeled with Alexa Fluor647 (1:500; AB_2338967; Jackson ImmunoResearch), goat α-HRP labeled with Cy3 (1:250; AB_2338959; Jackson ImmunoResearch). Larval preparations were mounted in Vectashield (Vector Laboratories). Images were acquired at room temperature. Confocal images were obtained with a Leica SP8 system (Leica Microsystems) equipped with HC PL APO Lasers with 488 and 531 nm were used. Again, images were manually corrected and brightness and contrast were manually adjusted in FIJI. For NMJs, images were obtained with 300 nm z-stack size, maximum projected, background subtracted using the rolling ball method (rolling ball radius 50 pixels) with brightness and contrast manually adjusted. To assess NMJ morphology, images were acquired using an Apotome System (Axioview 200M, objective 63x, NA 1.4, oil; Zeiss). Brp puncta per NMJ and NMJ area were measured using a thresholding algorithm in FIJI (Schindelin et al., 2012), essentially as described previously (Mrestani et al., 2021; Paul et al., 2022). Boutons per NMJ were counted manually. SIM imaging was performed as previously described (Dannhäuser et al., 2022) using a Zeiss Elyra S.1 structured illumination microscope equipped with a sCMOS camera (pco.edge 5.5 m) and an oil-immersion objective (Plan-Apochromat 63x, 1.4 NA). Lasers with 488 and 531 nm were used. Again, images were maximum-projected and brightness and contrast were manually adjusted.

dSTORM

dSTORM imaging of the specimen was performed essentially as previously reported (Ehmann et al., 2014; Paul et al., 2015, 2022; Mrestani et al., 2021; Dannhäuser et al., 2022). The same primary antibodies as described above were used in the following concentrations: mouse α-Brp (Brp\textsubscript{N/C2}, 1:100), rabbit α-HA (1:500). The following secondary antibodies were used: goat α-rabbit F(ab')\textsubscript{2} fragments labeled with Alexa Fluor 647 (1:500; A21246; Thermo Fisher Scientific) and goat α-mouse IgGs labeled with Alexa Fluor 532
(1:500; A11002; Thermo Fisher Scientific). After staining, larval preparations were incubated in 100 mM mercaptoethanolamine (Sigma-Aldrich) in a 0.2 M sodium phosphate buffer, pH 7.9, to allow reversible switching of single fluorophores during data acquisition (van de Linde et al., 2008). Images were acquired on an inverted microscope (IX-71, 60x, NA 1.49, oil immersion; Olympus) equipped with a nosepiece-stage (IX2-NPS; Olympus). 647 nm (F-04306-113; MBP Communications Inc.) and 532 nm (gem 532; Laser Quantum) lasers were used for excitation of Alexa Fluor647 and Alexa Fluor532, respectively. Laser beams were passed through clean-up filters (Brightline HC 642/10 and Semrock, ZET 532/10, respectively), combined by two dichroic mirrors (LaserMUX BS 514–543 and LaserMUX BS 473–491R, 1064R, F38-M03; AHF Analysentechnik), and directed onto the probe by an excitation dichroic mirror (HC Quadband BS R405/488/532/635, F73-832; AHF Analysentechnik). The emitted fluorescence was filtered with a bandwidth filter (HC-quadband 446/523/600/677; Semrock) and a long pass (Edge Basic 635; Semrock) or bandpass filter (Brightline HC 582/75; Semrock) for the red and green channels, respectively, and divided onto two cameras (iXon Ultra DU-897-U; Andor) using a dichroic mirror (HC-BS 640 imaging; Semrock). For the red channel, image resolution was 127 nm × 127 nm per pixel to obtain super-resolution of RIMHA-Znf. For the green channel, image resolution was 130 nm × 130 nm per pixel. Localization of single fluorophores and high-resolution image reconstruction was performed with rapidSTORM (Heilemann et al., 2008; Wolter et al., 2010; van de Linde et al., 2011; Wolter et al., 2012; https://www.biozentrum.uni-wuerzburg.de/super-resolution/archiv/rapidstorm/). Only fluorescence spots with an A/D count over 5,000 were analyzed and a subpixel binning of 10 nm px⁻¹ was applied.

**Analysis of localization data**

Localization data were analyzed essentially as described previously (Mrestani et al., 2021; Dannhäuser et al., 2022), using custom-written Python code (https://www.python.org/), language version 3.6) and the Python interface Jupyter (Kluyver et al., 2016) to directly load and analyze localization tables from rapi. The Python interface Jupyter (Kluyver et al., 2016) to directly load and analyze localization tables from rapi, the Python interface Jupyter (Kluyver et al., 2016) to directly load and analyze localization tables from rapi, and six boutons, were masked in the reconstructed, binned images from based analysis, the regions of interest, corresponding to terminal RIM nanoclusters at presynaptic active zones. RIM localization data were analyzed essentially as described previously (van de Linde et al., 2008). After staining, larval preparations were incubated in 100 mM mercaptoethanolamine (Sigma-Aldrich) in a 0.2 M sodium phosphate buffer, pH 7.9, to allow reversible switching of single fluorophores during data acquisition (van de Linde et al., 2008). Images were acquired on an inverted microscope (IX-71, 60x, NA 1.49, oil immersion; Olympus) equipped with a nosepiece-stage (IX2-NPS; Olympus). 647 nm (F-04306-113; MBP Communications Inc.) and 532 nm (gem 532; Laser Quantum) lasers were used for excitation of Alexa Fluor647 and Alexa Fluor532, respectively. Laser beams were passed through clean-up filters (Brightline HC 642/10 and Semrock, ZET 532/10, respectively), combined by two dichroic mirrors (LaserMUX BS 514–543 and LaserMUX BS 473–491R, 1064R, F38-M03; AHF Analysentechnik), and directed onto the probe by an excitation dichroic mirror (HC Quadband BS R405/488/532/635, F73-832; AHF Analysentechnik). The emitted fluorescence was filtered with a bandwidth filter (HC-quadband 446/523/600/677; Semrock) and a long pass (Edge Basic 635; Semrock) or bandpass filter (Brightline HC 582/75; Semrock) for the red and green channels, respectively, and divided onto two cameras (iXon Ultra DU-897-U; Andor) using a dichroic mirror (HC-BS 640 imaging; Semrock). For the red channel, image resolution was 127 nm × 127 nm per pixel to obtain super-resolution of RIMHA-Znf. For the green channel, image resolution was 130 nm × 130 nm per pixel. Localization of single fluorophores and high-resolution image reconstruction was performed with rapidSTORM (Heilemann et al., 2008; Wolter et al., 2010; van de Linde et al., 2011; Wolter et al., 2012; https://www.biozentrum.uni-wuerzburg.de/super-resolution/archiv/rapidstorm/). Only fluorescence spots with an A/D count over 5,000 were analyzed and a subpixel binning of 10 nm px⁻¹ was applied.

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**Electrophysiology**

Two-electrode voltage clamp recordings (Axoclamp 2B amplifier, Digidata 1440A; Molecular Devices) were obtained from abdominal muscle 6 in segments A2 and A3 as previously described.
(Dannhäuser et al., 2022; Paul et al., 2022). All measurements were obtained at room temperature in HL-3 (Stewart et al., 1994) with the following composition (in mM): NaCl 70, KCl 5, MgCl2 20, NH4Cl 10, trehalose 5, sucrose 115, Heps 5, and CaCl2 1, pH adjusted to 7.2. Intracellular electrodes had resistances of 10–30 MO and were filled with 3 M KCl. For analysis, only cells with an initial membrane potential of ~50 mV or less and a membrane resistance of ≥ 4 MO were included. During recordings, cells were clamped at a holding potential of ~80 mV (miniature EPSCs, mEPSCs) or ~60 mV (evoked EPSCs, eEPSCs). Signals were low-pass filtered at 10 kHz and analyzed in Clampfit (Version 11.1, Molecular Devices). mEPSCs were recorded for 90 s and the occurrence rate of mEPSCs determined mEPSC frequency. Amplitude, rise time, and decay time constants were determined using an average of all mEPSCs recorded within one time period. To evoke synaptic currents, nerves were stimulated via a suction electrode with pulses of 300 μs length and typically at 12 V (Grass S48 stimulator and isolation unit SIU5; Astro-Med). We applied a paired-pulse protocol with 0.2 Hz frequency and 30 ms interpulse intervals. For analysis, 5–10 traces per interval were averaged. The quantal content was estimated by dividing the mean eEPSC amplitude by the mean mEPSC amplitude measured in one cell. mEPSC amplitudes were corrected for the more hyperpolarized holding potential (Hallermann et al., 2010).

**PhTx treatment**

PhTx 433 tris (trifluoroacetate) salt (PhTx, CAS 276684-27-6; Santa Cruz Biotechnology) was dissolved in dimethyl sulfoxide (DMSO) to obtain a stock solution of 4 mM and stored at ~20°C. For each experiment, the respective volume was further diluted with freshly prepared HL-3 to a final PhTx concentration of 20 μM in 0.5% DMSO. Control experiments were performed with the same DMSO concentration in HL-3. PhTx treatment of semi-intact preparations was performed essentially as described previously (Frank et al., 2006; Mrestani et al., 2021; Dannhäuser et al., 2022). In brief, larvae were pinned down in calcium-free, ice-cold HL-3 at the anterior and posterior endings, followed by a dorsal incision along the longitudinal axis. Larvae were incubated in 10 μl of 20 μM PhTx in DMSO for 10 min at room temperature. After this incubation time, PhTx was replaced by HL-3 and larval preparations were completed, followed by electrophysiological measurements or dSTORM imaging.

**Statistics**

Statistical analyses were performed with Sigma Plot 13 (Systat Software) or GraphPad Prism 9. D’Agostino & Pearson (electrophysiology) or Shapiro–Wilks (imaging data) were used to test normality. If data were not normally distributed, we used the non-parametric Mann–Whitney rank sum test (eEPSC and mEPSC amplitudes in rim<sup>HA-Znf</sup> ctrl versus phtx; dSTORM parameters of RIM<sup>HA-Znf</sup> and Brp<sup>PS2</sup> in rim<sup>HA-Znf</sup> ctrl versus phtx), the Kruskal–Wallis test (mEPSC frequency, mEPSC amplitude, eEPSC rise time, paired pulse ratios of wt, rim<sup>rescue-Znf</sup>, rim<sup>Y5-Znf</sup> and rim<sup>HA-Znf</sup>) or one-way ANOVA (mEPSC amplitude, mEPSC tau decay, eEPSC tau decay in wt, rim<sup>rescue-Znf</sup>, rim<sup>Y5-Znf</sup> and rim<sup>HA-Znf</sup>) and reported as mean ± SEM. In box plots, horizontal lines represent median, boxes, quartiles, and whiskers 10<sup>th</sup> and 90<sup>th</sup> percentiles, unless indicated otherwise. Scatter plots show individual data points. Bin counts in histograms were normalized to the total number of observed events which was set to 1. All plots were produced with SigmaPlot. Figures were assembled using Adobe Illustrator (Adobe Creative Cloud 2022). Tables S1–S8 and S10 contain all numerical values stated or not stated in text and figure legends including P-values and sample sizes.

**Data Availability**

The authors declare that the custom-written Python code and all datasets supporting the findings of this work are available from the corresponding authors.

**Supplementary Information**

Supplementary information is available at [https://doi.org/10.26508/lsa.202302021](https://doi.org/10.26508/lsa.202302021).

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**Author Contributions**

A Mrestani: conceptualization, data curation, formal analysis, funding acquisition, investigation, visualization, and writing—original draft.
S Dannhäuser: data curation, formal analysis, investigation, and writing—review and editing.
M Pauli: software, formal analysis, validation, and investigation.
P Kollmannsberger: software, formal analysis, and writing—review and editing.
M Hübsch: investigation.
L Morris: investigation.
T Langenhan: conceptualization, funding acquisition, and writing—review and editing.
M Heckmann: conceptualization, resources, supervision, funding acquisition, validation, methodology, and writing—original draft and project administration.
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