Nanoscale Structural Plasticity of the Active Zone Matrix Modulates Presynaptic Function

Graphical Abstract

Highlights
- Dual-color STORM imaging of active zone (AZ) organization
- Neuronal activity bidirectionally regulates clustering of the AZ matrix
- Presynaptic machinery and function inversely correlate with AZ matrix clustering

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In Brief
Glebov et al. find that the clustered presynaptic matrix restricts the recruitment of presynaptic machinery to the active zone. Neuronal activity bidirectionally controls clustering of the AZ matrix to control presynaptic function. These data suggest that AZ structure may locally control function through molecular crowding.
Nanoscale Structural Plasticity of the Active Zone Matrix Modulates Presynaptic Function

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SUMMARY

The active zone (AZ) matrix of presynaptic terminals coordinates the recruitment of voltage-gated calcium channels (VGCCs) and synaptic vesicles to orchestrate neurotransmitter release. However, the spatial organization of the AZ and how it controls vesicle fusion remain poorly understood. Here, we employ super-resolution microscopy and ratiometric imaging to visualize the AZ structure on the nanoscale, revealing segregation between the AZ matrix, VGCCs, and putative release sites. Long-term blockade of neuronal activity leads to reversible AZ matrix unclustering and presynaptic actin depolymerization, allowing for enrichment of AZ machinery. Conversely, patterned optogenetic stimulation of postsynaptic neurons retrogradely enhanced AZ clustering. In individual synapses, AZ clustering was inversely correlated with local VGCC recruitment and vesicle cycling. Acute actin depolymerization led to rapid (5 min) nanoscale AZ matrix unclustering. We propose a model whereby neuronal activity modulates presynaptic function in a homeostatic manner by altering the clustering state of the AZ matrix.

INTRODUCTION

Synaptic transmission begins with the entry of calcium into the presynaptic terminal through voltage-gated calcium channels (VGCCs), followed by the fusion of neurotransmitter-filled synaptic vesicles (SV) with the presynaptic membrane. Both of these events occur at the active zone (AZ), a specialized site in presynaptic boutons that brings together VGCCs and synaptic vesicles within close proximity of each other and of release sites (Südhof, 2012).

Modulation of the AZ structure is thought to be an important regulatory site controlling the efficacy of synaptic transmission. The function of the AZ can be dynamically regulated in the context of homeostatic plasticity, resulting in increase in presynaptic Ca^{2+} influx (Zhao et al., 2011) and release probability (Murthy et al., 2001; Vitureira et al., 2011). Similarly, blockade of postsynaptic activity also results in structural changes in hippocampal boutons (Murthy et al., 2001) as well as numerous changes in the levels of AZ proteins in cortical synapses (Lazarevic et al., 2011), lending further support to the idea that the AZ structure is dynamic and controls presynaptic function. However, the connection between neuronal activity and the precise structure of the AZ has not been formally investigated.

Ultrastructurally, the AZ is formed of a dense scaffold containing hundreds of copies of dozens of different proteins. This tight arrangement is thought to result in a locally crowded molecular environment (Morales et al., 2000; Rothman et al., 2016; Wilhelm et al., 2014), that could lead to competition for space between AZ proteins. Due to its high molecular weight (410 kDa) and large estimated numbers (<400 molecules per AZ) (Wilhelm et al., 2014), the matrix protein Bassoon (Bsn) is estimated to account for a large proportion of the AZ matrix material. This is in contrast to the levels of other AZ proteins such as Rab3-interacting molecule (RIM) and VGCCs, whose numbers are estimated to be an order of magnitude lower (Holderith et al., 2012; Indriati et al., 2013; Nakamura et al., 2015; Wilhelm et al., 2014). As a result of its large bulk and high abundance, Bsn is a prime candidate for shaping the structure of the AZ matrix, with important implications for presynaptic function.

Theoretical simulations and electrophysiological and structural evidence suggest that the spatial organization of the AZ components may play an important role in coupling Ca^{2+} signaling and presynaptic release (Eggermann et al., 2011; Ernolyuk et al., 2013; Grauel et al., 2016; Holderith et al., 2012; Nakamura et al., 2015; Tang et al., 2016). Several key AZ proteins, including VGCCs, RIM, and Bsn, are non-randomly distributed in the AZ, raising the possibility that the structural arrangement of the AZ
may be important for synapse function. In agreement with this, correlation between RIM levels and release probability (Holdersh et al., 2012; Paul et al., 2015; Peled et al., 2014) and preferential occurrence of vesicle fusion in RIM-enriched domains (Tang et al., 2016) establishes RIM as a structural marker for release sites and of synapse strength. However, the relative spatial organization, as well as the dynamics, of the AZ components remains unknown.

Here, we hypothesized that the AZ architecture may be regulated by neuronal activity and that the structural plasticity of the AZ may be involved in controlling synapse function. To test this, we used high-resolution imaging to investigate the relationship between neuronal function and the nanoscale AZ structure in hippocampal synapses. Our results show that the AZ architecture features largely non-overlapping domains of Bsn-enriched matrix and either VGCCs or RIM-enriched areas located in close proximity to each other. We find that the clustering of Bsn is bidirectionally controlled by postsynaptic neuronal activity. Activity blockade results in Bsn unclustering and recruitment of multiple AZ proteins, while local Bsn clustering is negatively correlated with AZ protein enrichment and presynaptic function. Our data therefore suggest that clustering of the Bsn-enriched AZ matrix may act as a barrier to the recruitment of the presynaptic machinery, thus limiting synapse function. This restrictive modality constitutes a mechanism gating local presynaptic release, implicating molecular congestion in the regulation of synaptic function.

RESULTS

Nanoscale Imaging of the AZ Structure

Confocal microscopy imaging of Bsn and the canonical P/Q-type VGCC pore-forming subunit Ca_{2,1} shows clear colocalization in punctate structures (Figure 1A), indicating that they both localize to the AZ; this level of resolution is, however, insufficient to visualize their detailed distribution within the AZ. To this end, we imaged Bsn and Ca_{2,1} using dual-color direct stochastic optical reconstruction microscopy (dSTORM) (Winterflood et al., 2015) (Figures S1A–S1C). Although Ca_{2,1} and Bsn within the AZ were localized in close proximity, there was little spatial overlap between them, suggesting spatial segregation on the nanoscale (Figure 1A). The median nearest neighbor distance (NND) between Ca_{2,1} and Bsn localizations within the AZ was 36.2 nm (Bsn to Ca_{2,1}) and 36.3 nm (Ca_{2,1} to Bsn), with multiple instances of >100 nm distance (Figures 1B and 1C). Similar results were obtained for the distribution of Bsn and a release site marker RIM (Figures 1D–1F), although the median NNDs between Bsn and RIM localizations were consistently less than that between Bsn and Ca_{2,1} (Figures 1G and 1H). To assess the relationship between Bsn clustering and Ca_{2,1} recruitment, we plotted the AZ-specific enrichment of Ca_{2,1} as a function of Bsn-Bsn NND. There was a positive correlation between these two values, indicating that loosening of Bsn clustering may be associated with Ca_{2,1} accumulation (Figure 1I). Importantly, similar results were obtained for Bsn and RIM (Figure 1J). Together these data imply that Bsn clustering may limit the enrichment of other AZ proteins important for synapse function.

To assess the clustering and size of AZ protein domains, we binned the localization data into bins of 25, 50, and 100 nm and computed the Pearson’s correlation coefficients to measure the resulting spatial overlap (Figures S2A and S2B). As a positive control for overlap, we used samples that were doubly labeled for Bsn and either Ca_{2,1} or RIM was significantly different from Bsn-Bsn, while, at the 100 nm, all three labels showed nearly complete overlap (Figure S2C). Thus, Bsn and Ca_{2,1} as well as Bsn and RIM are spatially segregated on a 25- to 50-nm scale, in agreement with the NND measurements and confocal microscopy data.

Activity Blockade Unclusters the AZ Matrix

In contrast to a previous study reporting inactivity-induced decreases in Bsn levels in cortical neurons (Lazarevic et al., 2011), we found no evidence of such a decrease in hippocampal neurons (Glebov et al., 2016); furthermore, Bsn synthesis rate is independent of activity manipulation (Schanzenbächer et al., 2016), suggesting that activity-dependent regulation of the hippocampal AZ structure may differ from that in the cortex. We therefore used STORM to visualize the distribution of Bsn, Ca_{2,1}, and RIM following activity blockade.

Blockade of action potential firing with tetrodotoxin (2 μM, 48 hr) had no effect on the number of Bsn and RIM localizations in the AZ; on the other hand, the number of Ca_{2,1} localizations was increased (Figure S1D; data not shown), in agreement with previous studies (Lazarevic et al., 2011; Zhao et al., 2011) (also see Figure 4). Surprisingly, the corresponding NND measures were largely unchanged (Figures 2A–2D), with the exception of the slightly decreased Ca_{2,1}-Bsn NND, consistent with an increase in Ca_{2,1} recruitment. Furthermore, the relationship between Bsn NND and either Ca_{2,1} or RIM recruitment, as well as the correlation between binned datasets, were not affected by tetrodotoxin (TTX) treatment (Figures 2E, 2F, and S2D). Thus, we conclude that despite the apparent AZ enrichment of VGCCs after activity blockade, the degree of spatial segregation between Bsn- and VGCC- or RIM-enriched domains is independent of neuronal activity.

For a quantitative insight into the activity-dependent changes in the AZ structure, we used clustering analysis based on Ripley’s K-statistics (Williamson et al., 2011) (Figures 2G–2K). Extraction of clustering statistics from thresholded data revealed that approximately 42% of Bsn labeling was organized in significantly smaller clusters than an average AZ (Schikorski and Stevens, 1997) (Figure 2J), indicative of the non-random distribution of Bsn within the AZ (Tang et al., 2016). TTX treatment decreased the maximal value of the Ripley function (Figures 2H and 2I), the area of clusters and the proportion of Bsn molecules in clusters (Figures 2J and 2K). In contrast, the density of Bsn within the cluster, the overall levels of Bsn and cluster number remained unchanged (Figures S1D–S1F). This suggests that blockade of neuronal activity leads to dissociation of Bsn from the clusters within the AZ, resulting in their shrinkage.

Ratiometric Imaging of Protein Clustering

The long image acquisition times and substantial computational trade-offs associated with STORM imaging limit its use...
for investigation of large synapse populations. To circumvent these restrictions, we sought for an alternative approach allowing for rapid and high-throughput comparative visualization of protein clustering across multiple regions of interest, using ratiometric fluorescence resonance energy transfer (FRET) (Glebov and Nichols, 2004). In this approach (see Experimental Procedures for details), changes in distances between proteins can be visualized as changes in the ratio of intensities of the donor-acceptor fluorophores (Glebov and Nichols, 2004); we will hereafter refer to this as the Acceptor/Donor Ratio (RA/D) (Figure 3A).

To test the validity of this approach, we carried out the following control experiments (Figure S3). First, we performed ratiometric labeling using serial dilution of a primary anti-Bsn antibody, reasoning that increasingly diluted antibody concentrations should result in lower RA/D values due to the increased interfluorophore distances. Indeed, dilution of the secondary antibodies progressively lowered the RA/D (Figure S3A). Similar results were obtained for two other proteins, namely an endosome marker transferrin receptor (Figure S3B) and a synaptic vesicles protein vGlut1 (data not shown), highlighting the generalizable nature of the ratiometric approach. Second, to demonstrate that our approach can report acute decreases in clustering of a probe, we ratiometrically imaged limited actin depolymerization in neurons using a low concentration (500 nM) of the actin depolymerization drug vinblastine (Figure 3C). This resulted in a decrease in the RA/D ratio, as expected (Figure 3D). These results suggest that the ratiometric approach can be used to quantitatively assess changes in protein clustering in a rapid and high-throughput manner.

Figure 1. Dual-Color STORM Imaging of the AZ Structure

(A) Imaging of Bsn and Ca$_{2.1}$, Left, Ca$_{2.1}$ and Bsn colocalize in puncta as visualized using confocal light microscopy. Scale bar, 5 μm. Right, neurons were stained for Bsn (CF680) and Ca$_{2.1}$ (AF647) and imaged using dual-color STORM. Zoomed regions, AZs in the “face-up” orientation. Scale bar, 1 μm.

(B) Histogram of Bsn-to-Ca$_{2.1}$ NNDs in AZs. Dashed line denotes median value.

(C) Histogram of Ca$_{2.1}$-to-Bsn NNDs in AZs. Dashed line denotes median value.

(D–F) As in (A)–(C) but for RIM and Bsn.

(G) Comparison of the median values between Bsn-to-Ca$_{2.1}$ and Bsn-to-RIM NNDs.

(H) Comparison of the median values between Ca$_{2.1}$-to-Bsn and RIM-to-Bsn NNDs. *p < 0.05, **p < 0.01, Mann-Whitney test.

(I) Correlation plot for relative AZ enrichment of Ca$_{2.1}$ versus Bsn-to-Bsn NND.

(J) Correlation plot for relative AZ enrichment of RIM versus Bsn-to-Bsn NND. n = 3, ten synapses/experiment.
depolymerizing drug Latrunculin A (LatA) that does not reduce the overall levels of the polymerized filamentous (F)-actin (Glebov et al., 2015). Under these conditions, a decrease in $R_{A/D}$ was readily detected, consistent with decreased clustering of F-actin (Figure S3C). Third, to show that our assay can detect acute increases in clustering, we used a recombinant cell-surface glycospophatidylinositol (GPI)-GFP probe after inducing clustering by antibody crossing (Glebov and Nichols, 2004). 10 min crosslinking resulted in a visible clustering of the probe, concomitant with a robust increase in $R_{A/D}$ (Figure S3D). Taken together, these controls show that the ratiometric approach can be used to report nanoscale changes in clustering of a probe.

**Ratiometric Imaging Confirms Inactivity-Induced AZ Bsn Matrix Unclustering**

Having validated our assay, we used it to study the activity-dependent nanoscale changes of the AZ matrix using confocal microscopy. Bsn levels and the size of the Bsn-positive puncta were unaffected by TTX, suggesting that the gross morphology of the AZ remained intact (Figures 3D and S3E). However, in agreement with the STORM data (Figure 2), TTX treatment led to a decrease in $R_{A/D}$ for Bsn (Figures 3B and 3C). An $R_{A/D}$ decrease was also observed using two alternative anti-Bsn antibodies, further confirming the unclustering of the AZ matrix (Figure S3F). The FRET ratio measured using the sensitized emission approach (Glebov and Nichols, 2004) was also reduced by TTX treatment (Figure S4A). In contrast to Bsn, $R_{A/D}$ values for the postsynaptic matrix proteins Pds95 and Gephyrin remained unchanged by the TTX treatment (Figures S4C and S4D), in agreement with recent studies showing only subtle local effects of activity modulation on postsynaptic scaffolding (MacGillivray et al., 2013; Specht et al., 2013; Tang et al., 2016).

A washout (48 hr) of TTX led to an increase in $R_{A/D}$ (Figure 3E), indicating that the effect was reversible. The effect of TTX was evident after 24 hr, but not 3 hr (Figure S4E; data not shown), consistent with a typical timeline for homeostatic synaptic plasticity (Pozo and Goda, 2010). $R_{A/D}$ was not significantly altered by TTX when a spectrally non-overlapping pair of fluorophores was used (Figure S3G), indicating that the observed change in $R_{A/D}$ could be solely attributed to a change in FRET efficiency, rather than a change in the binding efficiency of the antibody.

To directly compare the ratiometric assay with the STORM measurement, we employed our data from the serial dilution experiment (Figure S3A) to plot the relationship between $R_{A/D}$ and the inter-fluorophore distance. Ultrastructural investigations of AZ have shown that the AZ architecture is essentially flat, with the AZ plane immediately underlying the presynaptic membrane by the synaptic cleft (Harris and Weinberg, 2012; Meyer et al., 2014; Murthy et al., 2001; Südhof, 2012; Tang et al., 2016). According to the constraints of the two-dimensional approximation of the AZ structure, the distance between the fluorophores (i.e., labeled Bsn molecules) within the AZ matrix should scale proportionally to the square root of the dilution factor (Figure S3H). In agreement with this, the decrease in $R_{A/D}$ plotted against the estimated increase in distance was well fitted by a single exponential decay curve. The decrease in Bsn $R_{A/D}$ measured following the TTX treatment (4.5%) matched the increase in Bsn-Bsn NND measured using STORM (33.7%) (Figure S3H) directly confirming the validity of the ratiometric approach for visualization of nanoscale clustering.

Having established that blockade of activity results in a decrease in AZ clustering, we tested the converse, i.e., whether enhanced activity would result in an increase in AZ clustering. To this end, we took advantage of an optogenetic stimulation paradigm that allows for controlled induction of neuronal activity in both space and time. Neurons expressing Channelrhodopsin2-YFP (ChR2-YFP) were stimulated with two previously characterized “burst” and “sparse” stimulus patterns (Grubb and Burne, 2010). To quantify the effect of activation in cis, we selected the synapses formed by the ChR2-expressing axons onto the untransfected dendrites. Conversely, to quantify the effect of activation in trans, $R_{A/D}$ was measured in the synapses formed by untransfected axons onto the ChR2-expressing dendrites (Figure 3J). Only “burst” stimulation in trans resulted in an increase in $R_{A/D}$ (Figures 3K and 3L), while $R_{A/D}$ in distal synapses was unaffected by the stimulation (Figure 3M). Specific patterns of neuronal activity therefore drive transsynaptic AZ clustering in a localized manner.

**NMDAR Activity and Cannabinoid Signaling Regulate AZ Matrix Clustering**

A major consequence of postsynaptic depolarization is an increase in postsynaptic Ca$^{2+}$ influx, which in hippocampal neurons is primarily mediated by L-type VGCCs and NMDA-type glutamate receptors (NMDARs) (Bloodgood and Sabatini, 2007). To establish their contributions to the regulation of AZ clustering, we incubated neurons in either the L-type VGCCs blocker nifedipine or the NMDAR antagonist (2R)-amino-5-phosphonovaleric acid (APV). APV, but not nifedipine, resulted in a significant decrease in $R_{A/D}$, showing that NMDAR activation was required.
for AZ matrix clustering (Figures 3F and 3G). This effect was recapitulated by blockade of the AMPA/kainate-type glutamate receptors (AMPAR/KAR) with 2,3-dihydroxy-6-nitro-7-sulphamoyl-benzof(1)quinoxaline-2,3-dione (NBQX), further confirming the role of postsynaptic depolarization in AZ clustering (Figure 3H). Neither APV nor NBQX treatment affected the levels of Bsn (Figure 3I). The timescale of these effects (Figure S4E; data not shown) was consistent with the timescale of slow synaptic scaling (Pozo and Goda, 2010) rather than that of a rapid brain-derived neurotrophic factor (BDNF)-dependent homeostatic regulation of presynaptic activity (Jakawich et al., 2010).

To further investigate the signaling mechanisms linking postsynaptic activity with AZ matrix clustering, we tested for the involvement of endocannabinoid signaling, which is a major activity-dependent retrograde pathway operating in hippocampal neurons (Castillo et al., 2012). Inhibition of endocannabinoid synthesis by tetrahydrocannabinol (THL) resulted in a decrease in R_A/D, as did bidirectional manipulation of cannabinoid signaling by a cannabinoid receptor 1 (CB1) agonist arachidonyl-2'-chloroethylamide (ACEA) and an inverse agonist AM251 (Figures S4F–S4J). These data implicate NMDAR activation and endocannabinoid release as two mechanisms involved in transsynaptic regulation of AZ matrix clustering.

**Inactivity-Induced Presynaptic Recruitment of Multiple AZ Proteins**

Given the discrepancy between our findings and the effects of activity blockade on AZ composition in cortical neurons (Lazarevic et al., 2011), we investigated the effect of activity blockade on presynaptic recruitment of other AZ proteins in the hippocampal synapse. In agreement with our STORM data, there was a significant increase in CaV2.1 channel (P/Q-type VGCCs) recruitment (Figure 4A). In contrast, synaptic levels of RIM were unchanged (Figure 4D), as were those of another adaptor AZ protein Munc13-2 (data not shown). In yet another contrast with cortical neurons, synaptic recruitment of a Bsn-related AZ scaffolding protein Piccolo (Polo) was also increased (Figure 4E), as were the levels of two other presynaptic VGCCs, R-, and N-type (Figures 4B and 4C). Taken together, these data are consistent with the notion that the decreased clustering of the AZ matrix may allow for increased recruitment of presynaptic machinery in hippocampal neurons.

To obtain a functional measure of the TTX effect on presynaptic Ca^{2+} signaling, we measured AP-evoked calcium influx in presynaptic terminals with the genetically encoded presynaptic calcium indicator SyGCaMP6f. In agreement with the structural findings, we observed a strong increase in calcium influx following 48-hr incubation in TTX (Figures 4F and 4G). We then used a specific channel blocker 1H-Agatoxin-IIVA (Aga) to quantify the contribution of the P/Q-type VGCCs to presynaptic Ca^{2+} rise in untreated and TTX-treated cultures. Inhibition of the SyGCaMP6f signal by Aga was significantly increased by TTX blockade (44% versus 32%, p = 0.0032) (Figures 4H–4K). Thus, activity blockade results in a preferential recruitment of P/Q-type VGCCs to the synapse.

**Synapse-Specific Correlation between AZ Matrix Clustering and Presynaptic Function**

To explore the functional relevance of AZ clustering regulation, we hypothesized that AZ clustering may correlate with recruitment of other presynaptic proteins involved in neurotransmitter release. To test for this, we measured enrichment of RIM and CaV_{2.1} in individual AZs and plotted it against local R_A/D. RIM enrichment was negatively correlated with R_A/D (Figures 5A and 5B), showing that the AZs with looser clustering preferentially recruited RIM, in agreement with our STORM data (Figures 1 and 2). In contrast, there was no correlation between R_A/D and the levels of a ubiquitous pre/postsynaptic scaffolding protein CASK (Figure S5B) (Hsueh et al., 1998; Sudhof, 2012). There was also no correlation between R_A/D and levels of CaV_{2.1}, possibly reflecting the existence of an extrasynaptic CaV_{2.1} pool likely to obfuscate a presynaptic specific correlation when performed at this level of resolution (Figure S5A); indeed, in our cells accumulation of the CaV_{2.1} label in the dendritic shaft and the cell body was often observed, and ultrastructural evidence supports the presence of the extrasynaptic CaV_{2.1} pool (Indriati et al., 2011).
Activity Blockade Leads to Recruitment of Multiple AZ Proteins in Hippocampal Neurons and Upregulation of Presynaptic Ca²⁺ Influx through P/Q-type VGCCs

(A) TTX treatment (2 μM, 48 hr) results in AZ enrichment of Ca₂⁺. n = 3.

(B) TTX treatment results in AZ enrichment of Ca₂⁺. Arrows indicate individual AZs; asterisk denotes a Ca₂⁺ punctum not associated with AZ. n = 3.

(legend continued on next page)
et al., 2013). Presynaptic enrichment of Ca$_{2+}$,1 at the AZ by TTX treatment, however, led to emergence of an inverse correlation with Bsn R$_{AD}$ (Figures 5C and 5D), indicating that Ca$_{2+}$,1 was preferentially recruited to the AZs with less Bsn clustering, in agreement with the STORM data (Figures 1 and 2). The levels of all of the above strongly correlated with local Bsn levels, indicating that the bigger synapses contained more of synaptic proteins (Figures S5C–S5F).

To investigate the link between AZ matrix clustering and presynaptic function, we took advantage of the two well-established assays for synaptic vesicle cycling based on (1) uptake of an antibody against the extracellular/luminal domain of the synaptic vesicles protein Synaptotagmin 1 (Scheiffele et al., 2000) and (2) live imaging of a fluorescent GFP-based pH-sensitive synaptic vesicles probe SypHy (Miesenböck et al., 1998). AZ-specific R$_{AD}$ exhibited a negative correlation with the uptake of anti-Syt1 antibody under conditions favoring either spontaneous (20 min at 37°C in presence of 2 μM TTX) or evoked (4 min at 37°C in presence of 50 mM KCl) presynaptic release (Figures S6A and S6B).

Additionally, we combined live SypHy imaging of presynaptic vesicle cycling with post hoc ratiometric imaging of AZ clustering to directly correlate the functional readout from the individual synaptic boutons with the structure of the individual AZs (Figure S6C). Imaging of presynaptic function with SypHy also yielded a negative correlation between R$_{AD}$ and the size of the rapidly releasable pool, assessed functionally from the amplitude of the response to a stimulus of 40 APs delivered at 20 Hz frequency (Figures S5E and S5F). The extent of local AZ matrix clustering is therefore inversely correlated with recruitment of presynaptic release machinery and synaptic vesicle cycling.

### Presynaptic Actin Dynamics Regulate AZ Clustering

To further characterize the mechanism underlying activity-dependent AZ clustering dynamics, we focused on actin dynamics that have been previously implicated in regulation of presynaptic plasticity (Cingolani and Goda, 2008; Morales et al., 2000; Sankaranarayanan et al., 2003). Actin dynamics have been previously suggested to act as a restrictive influence on presynaptic release through curbing of synaptic vesicles cycling (Morales et al., 2000). Thus, actin dynamics represent a promising candidate mechanism for linking neuronal activity and synaptic structure.

We first assessed the effect of long-term activity blockade on actin levels. TTX treatment globally reduced the F-actin levels in the entire neuron, suggesting that actin polymerization was regulated by the network activity (data not shown). Specifically, the levels of F-actin present in Bsn-positive puncta were reduced, consistent with inactivity-induced depolymerization of synaptic F-actin (Figures 6A and 6B). In agreement with this, prolonged pharmacological actin depolymerization by 5 μM LatA for 2 hr decreased Bsn R$_{AD}$; in contrast, block of actin depolymerization with Jasplakinolide (JaspI) had no effect on Bsn R$_{AD}$ (Figure 6C).

On the shorter timescale, treatment with 20 μM LatA decreased Bsn R$_{AD}$ within 5 min, consistent with the previously reported rapid induction of presynaptic plasticity by this actin depolymerization (Morales et al., 2000); at the same time, the area of the AZ remained the same, suggesting that the structural rearrangement was restricted to the nanoscale (Figure 6D). We propose that actin dynamics are a possible candidate for regulating AZ composition through activity-dependent remodeling of the AZ scaffolding.

### DISCUSSION

In this study, we have combined super-resolution, ratiometric, and functional imaging to establish the link between activity, AZ matrix organization, recruitment of presynaptic release machinery and synapse function at an archetypal CNS synapse. Crucially, we show that the AZ structure belies a surprising potential for reversible bidirectional reorganization at the nanoscale level, controlled by local postsynaptic activity that, in turn, fine-tunes presynaptic function. Taken together, our results show that local neuronal activity dynamically controls AZ organization through actin dynamics to modulate presynaptic structure and function.

### AZ Architecture on the Nanoscale and Its Regulation by Activity

Our STORM data show that the non-random organization of the AZ matrix characterized by Bsn clustering shows little overlap with the VGCCs and RIM-positive domains, with spatial segregation on the scale of 25–50 nm. These observations echo the previously reported differences in the distribution of the structural AZ components and VGCCs at the Drosophila neuromuscular junction (Ehmann et al., 2014; Fouquet et al., 2009).
suggesting that the existence of distinct structural domains may be a core feature of the AZ structure.

Our data also reveal an activity-dependent remodeling of the AZ at the nanoscale. Chronic blockade of network activity results in a decrease in the area of the Bsn clusters and a corresponding increase in Bsn NND, without any major changes in the overall AZ morphology or Bsn levels. These results agree with the previously published data in hippocampal neurons (Glebov et al., 2016; Schanzenbächer et al., 2016) but are at odds with another study in cortical neurons (Lazarevic et al., 2011). The reason for this discrepancy is not clear but may be due to the intrinsic differences between cell types. The activity-induced unclustering of the AZ matrix did not affect the distance between the Bsn domains and the neighboring VGCCs/RIM-positive areas, indicating sustained spatial segregation at the AZ following activity-dependent plasticity. This feature is in line with the idea that Bsn domains act to limit the spatial extent of their neighbors.

**Signaling Pathways Shaping Presynaptic Structure and Function**

Our optogenetics data suggest the presence of a putative local mechanism linking postsynaptic activity with presynaptic structure...
and function. These data are further supported by the AZ unclustering observed following chronic NMDAR and AMPAR/KAR blockade and suggest the involvement of postsynaptic Ca²⁺ signaling. Further still, calcium influx through L-type VGCCs was not required, indicating that the mode of calcium influx is important and likely involves local synaptic routes. Indeed, the requirement for NMDAR activation fits with the canonical role of NMDARs as coincidence receptors of transsynaptic depolarization and neurotransmitter release (Hunt and Castillo, 2012). Taken together, these data establish another role for NMDARs in regulating presynaptic function.

The precise details of the downstream events activated by NMDAR activation will require further investigation. Although our pharmacological data are consistent with the role of cannabinoid signaling in regulating AZ clustering and neurotransmitter release, the actual regulatory mechanism is likely to be complex, given that both upregulation and blockade of CB1 resulted in an unclustering of the AZ matrix. Signaling through CB1 receptors is complex and includes feedbacks that downregulate receptor availability following sustained activity (Dudok et al., 2015), which could help explain the responses observed here. Furthermore, manipulation of cannabinoid signaling in the hippocampus has been shown to modulate NMDA receptor function (Hampson et al., 2011), suggesting cross-talk between excitatory neurotransmission and cannabinoid signaling.

**A Model for Integrating Synaptic Activity, Structure, and Function**

How does the dynamic organization of the AZ matrix control presynaptic function? On the basis of our data, a model can be proposed whereby the clustering of the AZ matrix limits the recruitment of the presynaptic machinery, through competition for space (Figure 7). This notion is supported by the following observations: (1) enduring activity-independent spatial segregation between AZ matrix and VGCCs/release sites (Figures 1 and 2); (2) opposite effects of activity blockade on AZ matrix clustering and recruitment of multiple AZ components (Figures 2, 3, 4, and 5); (3) AZ-specific negative correlation between matrix clustering and presynaptic machinery recruitment/synaptic function (Figures 1, 4, and 5); and (4) activity-dependent regulation of synaptic F-actin levels and rapid induction of AZ matrix unclustering by actin depolymerization (Figure 6).

The limiting effect of the AZ matrix scaffolding on presynaptic function is consistent with the evidence showing distinct subdomains within the *Drosophila* AZ (Ehmann et al., 2014; Fouquet et al., 2009), a proposed spatial segregation at the calyx of Held AZ (Nakamura et al., 2015), estimates of molecular crowding at the synapse (Wilhelm et al., 2014), and a role for actin polymerization as an activity-regulated presynaptic restrictive factor (Morales et al., 2000; Sankaranarayanan et al., 2003).

**Limitations of the Current Study**

The modulatory (rather than mandatory) role for AZ scaffolding agrees well with the apparently non-essential role of Bsn and Pclo in synaptic transmission (Hallermann et al., 2010; Mukherjee et al., 2010), in contrast with the essential role of the other, less numerous components of the AZ (Betz et al., 2001; Kaeser et al., 2009; Schoch et al., 2002). Moreover, the restrictive modality of the AZ matrix appears to act as a functional counterpart to the protein-protein interactions of other AZ proteins that serve to recruit presynaptic machinery to the AZ (Betz et al., 2001; Davydova et al., 2014; Kaeser et al., 2011; Südhof, 2012).

Certain considerations arising from the experimental techniques need to be taken into account when interpreting the results of this study. First, a combination of primary and secondary antibody labeling will affect the ability of STORM imaging to

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**Figure 6. Activity-Dependent Actin Dynamics Regulate AZ Clustering**

(A) Synaptic F-actin levels are regulated by activity. Neurons were treated with TTX for 48 hr and stained for Bsn and AF647-Phalloidin. Second row depicts Phalloidin signal present in Bsn-positive puncta (AZs).

(B) The intensity of Phalloidin staining in Bsn-positive puncta in control and TTX-treated cultures. One region of interest corresponds to one Bsn-positive punctum. n = 3, 310–806 regions of interest/experiment.

(C) The effect of 2-hr incubation with 5 μM actin-depolymerizing drug Latrunculin A (LatA) or 2 μM actin-polymerizing drug Jasplakinolide (Jaspl) on R_{A,D}. n = 4, 15 regions of interest/experiment.

(D) Depolymerization of actin by 20 μM LatA rapidly decreases R_{A,D}. Error bars indicate 10–90 percentile range.
Macromolecular Congestion as a Regulatory Mechanism at the Synapse and Beyond

Limiting presynaptic function by molecular congestion is in contrast to the permissive effect afforded by the clustering of the postsynaptic scaffold (MacGillavry et al., 2013; Specht et al., 2013; Tang et al., 2016), highlighting the fundamental differences in regulatory mechanisms operating on opposite sides of the synapse. From a purely mechanistic point of view, the curbing of presynaptic release by the AZ matrix could be viewed as an example of endogenous macromolecular congestion impacting on cellular function, with the clustering state of the AZ matrix restricting the ingress of presynaptic molecules into the AZ, thus limiting the composition of the functional AZ machinery. Similar principles have been proposed to modulate synaptic vesicle dynamics through congestion by the actin cytoskeleton (Morales et al., 2000) or collisions with organelles (Rothman et al., 2016). It is worth considering that, in a broader scope of cell biology, nanoscale structural plasticity of macromolecular assemblies may play a role in other functionally relevant contexts in the cell, e.g., in receptor signaling (James and Vale, 2012), endosome sorting (Wallrabe et al., 2007), and gene expression (Tan et al., 2013). Further understanding of this emerging regulatory modality will benefit from locally correlating nanoscale structural characterization of these systems with measurable functional outcomes.

EXPERIMENTAL PROCEDURES

Detailed experimental procedures and materials can be found in the Supplemental Information.

For ratiometric imaging, coverslips with neurons were fixed, permeabilized, and labeled for immunocytochemistry using antibodies conjugated to two different fluorophores. For optogenetic stimulation, primary neuronal cultures were sparsely transfected with ChR2-YFP and stimulated for 48 hr. For quantification of synapse-specific correlation, coverslips were processed three-color immunocytochemistry, and recruitment of presynaptic machinery to the individual Bsn-positive puncta was correlated with the local R0/D values. For live imaging of presynaptic function, neurons were sparsely transfected with the
vesicle cycling sensor CMV::SypHy or Ca²⁺ sensor SyGCaMP6F and subjected to field stimulation while imaging. Images were analyzed using MATLAB codes (MathWorks). For correlative live-fixed imaging of presynaptic structure and function, live images of SypHy responses were aligned with the fixed ratio-metric images using a MATLAB routine (Figure S6). For STORM imaging, samples were fixed, permeabilized, stained for the proteins of interest, and imaged using either a commercially available N-STORM Nikon system or a custom-built setup as described before (Winterflood et al., 2015). Imaging was performed in objective-type near-total internal reflection fluorescence (TIRF) mode. An image-correlation-based drift correction was employed. All data analysis was performed in ImageJ and MATLAB. Statistical analysis was carried out using GraphPad Prism 6.0. Sample distribution was assessed using D’Agostino and Pearson’s omnibus normality test; to assess the significance of differences between datasets, Mann-Whitney test was used unless noted otherwise. Error bars indicate 10–90 percentile range. **p < 0.01, ***p < 0.005.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures and six figures and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2017.02.064.

AUTHOR CONTRIBUTIONS

O.O.G. designed and oversaw the project, performed the experiments, analyzed the data, and wrote the manuscript with input from other authors. R.E.J. carried out the electrophysiological stimulation experiments, analyzed the data, and contributed to the writing of the manuscript. C.M.W. carried out the dual-color STORM imaging, analyzed the data, and participated in the writing of the manuscript. D.M.O. performed the Ripley’s K-function clustering analysis and participated in the writing of the manuscript. E.A.B. analyzed the confocal microscopy data. P.D. oversaw the cannabinoid signaling experiments and participated in the writing of the manuscript. H.E. oversaw the dual-color STORM imaging experiments and participated in the writing of the manuscript. J.B. developed the registration algorithm, oversaw the project, and co-wrote the manuscript.

ACKNOWLEDGMENTS

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REFERENCES


Supplemental Information

Nanoscale Structural Plasticity of the Active Zone Matrix Modulates Presynaptic Function

Oleg O. Glebov, Rachel E. Jackson, Christian M. Winterflood, Dylan M. Owen, Ellen A. Barker, Patrick Doherty, Helge Ewers, and Juan Burrone
Figure S1. Supporting data for Figs. 1&2: STORM imaging of the AZ structure. (A), Schematic of the optical setup for dual color single-molecule localization microscopy by spectral-demixing dSTORM. F, emission filter; LP-DM, long-pass dichroic mirror (690 nm); M, mirrors; L, lenses; EM-CCD, Electron-multiplying charge-coupled device camera. (B) Emission spectra of AF647 (green) and CF680 (red), transmission of the dichroic (black) and transmission of the band-pass emission filter (gray box), and the 643 nm laser (blue). (C) Representative distributions of the normalized intensity ratios $r=(I_s−I_l)/(I_s+I_l)$, where $I_s$ and $I_l$ are the fluorescence intensities of the short and long wavelength channels respectively. The green and red vertical lines delimit the assignment of the cut-off values. (D-F) Further effect of the TTX treatment on the AZ nanoscale structure; the number of Bsn and Ca2.1 localization events (LEs) per ROI (D), the number of Bsn clusters per 3x3µm ROI (E) and the density of Bsn LEs within the cluster (F). N=3.
Figure S2. Supporting data for Fig. 1: Binning-based correlation analysis of spatial distribution of \(\text{Bsn} \) and \(\text{Ca}_{2.1}\). (A) Schematics of outcomes for differently distributed dually labeled AZ samples under 25nm/pixel, 50nm/pixel and 100nm/pixel binning conditions. As the binning capacity and the pixel size increases, correlations emerge in closely associated spatial arrangements. (B) An example of a binning procedure on a sample labeled for \(\text{Bsn}\) and \(\text{Ca}_{2.1}\). Note the emergence of pixels containing both \(\text{Bsn}\) and \(\text{Ca}_{2.1}\) (right, \(\text{Bsn AND Ca}_{2.1}\)) in the larger bins; these will read out as positive correlation. (C) \(\text{Bsn}\) and \(\text{Ca}_{2.1}\) or RIM-containing pixels are spatially excluded under the 25nm binning regime as evidenced by the colocalization analysis. Segregation: significant negative correlation; Random: no significant correlation; Co-clustering, significant positive correlation. (D) Same as (C), but following TTX treatment. \(* p < 0.05, \ast \ast \ast p < 0.001, \) Fisher’s exact test.
Figure S3. Supporting data for Fig. 3: Control experiments for the ratiometric imaging of protein clustering. (A) Control 1: serial dilution ratiometric staining of Bsn results in an increased distance between fluorophores and therefore decreased R_{A/D}. Neurons were stained with an anti-Bsn monoclonal mouse antibody and the mixture of donor- and acceptor-conjugated secondary anti-mouse antibodies. N=3. ***p<0.001, 1-way ANOVA. Error bars for all plots indicate 10-90 percentile range. (B) The same results were obtained for Transferrin receptor (TfR) N=3. ***p<0.001, 1-way ANOVA. (C) Control 2: partial depolymerization of F-actin results in an increased distance between actin filaments and therefore affects the R_{A/D} measured through binding of an F-actin probe Phalloidin. Left, neurons were treated with 0.5 μM Latrunculin A for 1h before fixation, permeabilization and staining with the mixture of donor- and acceptor-conjugated phalloidin. Note the apparently unchanged intensity of Phalloidin labeling in LatA-treated neurons(Glebov et al., 2015). Scale bar, 20 μm. Right, quantification. ***p<0.001, Student t-test. (D) Control 3: Acute induction of clustering of a cell surface protein results in a decrease in the donor-acceptor distance, manifesting itself in an increase in R_{A/D}. Left, U2OS cells expressing GPI-GFP were incubated with a rabbit anti-GFP antibody for 10min at RT, fixed, permeabilized, stained with a mouse anti-GFP antibody and a ratiometric mix of anti-mouse secondary antibodies. Scale bar, 5 μm. Right, quantification. N=3. ***p<0.001, Student t-test. (E) TTX treatment did not affect the area of the Bsn-positive puncta. (F) Three different anti-Bsn antibodies demonstrate ratiometrically evident TTX-induced unclustering. Top, the schematic of the Bsn polypeptide chain charting the putative epitopes used by the three antibodies (to scale). Bottom left, unclustering of the Bsn matrix quantified by the ratiometric imaging involving the rabbit anti-Bsn monoclonal antibody D63B6. Bottom right, unclustering of the Bsn matrix quantified by the ratiometric imaging involving the mouse anti-Bsn monoclonal antibody 219E1. N=3. P<0.001, Mann-Whitney U test. (G) TTX treatment does not significantly change R_{A/D} in the AF488-AF647 labeled cells (cf. Fig. 3C). Left, comparison of the overlap for the AF568 emission spectrum with AF647 excitation spectrum and AF488 emission spectrum with AF647 excitation spectrum. Note the almost complete lack of overlap in the latter case. (H) Direct comparison of the change in Bsn clustering measured using STORM and ratiometric analysis. Left, in a two-dimensional system, dilution of the labeling results in an increase in the distance proportional to the power of ½. Right, plot of the serial dilution data for Bsn, showing the single exponential fit and the 95% confidence intervals. The increase in distance as measured by STORM imaging (red line) corresponds to the decrease in R_{A/D} measured ratiometrically (green line).
Figure S4. Supporting data for Fig. 3. (A) FRET ratio as measured by the sensitized emission method (Glebov and Nichols, 2004b) is decreased by TTX treatment. N=3, 15 ROIs/condition. (B) Bsn levels were not affected by APV and NBQX treatment. (C-D) \( R_{AD} \) values for postsynaptic matrix proteins Psd95 (C) and Gephyrin (D) were not affected by activity blockade. N=4, 15 ROIs/condition. (E) 24h treatment with TTX or APV or NBQX was sufficient to induce reduction of Bsn \( R_{AD} \). N=3 experiments, 15 ROI/condition. (F) 24h treatment with THL reduces \( R_{AD} \). N=3 experiments, 15 ROIs/condition. (G) Effect of 24h treatment with cannabinoid receptor agonist 10μM ACEA. N=4. (H) Effect of 24h treatment with cannabinoid receptor inverse agonist 4μM AM251. N=4. (I) Effect of ACEA and AM251 on Bsn levels. N=4, 15 ROI/condition. *p<0.05, ***p<0.001, Student t-test. Error bars indicate 10-90 percentile range.
Figure S5. Supporting data for Fig. 5. Synapse-specific correlation between AZ clustering and presynaptic recruitment. (A) Lack of correlation between levels of CaV2.1 and R_A/D (compare with Fig. 5D). (B) Lack of correlation between levels of ubiquitously adapter protein CASK and R_A/D in untreated cultures. (C-F) Levels of the following proteins were plotted against the levels of Bsn. (C) RIM. (D) CASK. (E) CaV2.1. (F) CaV2.1 (48h TTX). Datasets from Fig. 5 were used. r, Spearman’s rank correlation coefficient. All correlations were significant (P<0.0001).
Figure S6. Supporting data for Fig. 5. Correlating AZ matrix clustering with presynaptic function. (A), Evoked vesicle cycling negatively correlates with AZ matrix density. Evoked vesicle cycling was induced by depolarization with 50mM K⁺ for 4min. (B), Spontaneous vesicle cycling negatively correlates with AZ matrix density. Spontaneous vesicle cycling was isolated by applying 2µM TTX for 20min. (C), Schematics of the correlated live-fixed imaging of presynaptic structure and function. Neurons expressing SypHy are imaged live under stimulation by 40APs at 20Hz. From this, ∆F responses of individual synapses are measured. Neurons are then fixed and stained for ratiometric imaging of Bsn (AF568/AF633) and for GFP to amplify the SypHy signal. All fixed image channels undergo affine transformation to align with the live image. The registered fixed GFP image is thresholded to generate a mask, which is applied to registered AF568 and AF633 images, before $R_{A/D}$ values are calculated for each synapse.
**Full Methods**

**Cell culture**

Dissociated hippocampal neuronal cultures were prepared from E18 rat embryos, plated onto poly-L-lysine-coated glass coverslips and maintained according to the standard mixed culture protocol. All experiments involving neurons were carried out at 16-21 days in vitro. COS7 cells were also grown according to a standard culture protocol. For confocal imaging, cells were plated onto 13mm round glass coverslips (thickness 1.0) placed in 35mm Petri dishes (4/dish). For STORM imaging, cells were plated onto 18mm round glass coverslips (Carl Zeiss, Germany), thickness 1.5. For live imaging of SypHy cells were plated on 35mm Grid500 μ-Dishes (Ibidi, Germany).

**Reagents**

Cell culture media was from Invitrogen. Poly-L-lysine was from Sigma. The following primary antibodies were used:

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Secondary antibodies were from Jackson Immunoresearch (USA). AlexaFluor (AF)568, AF647-conjugated Phalloidin and FM1-43fx were from Molecular Probes. APV, NBQX, TTX were from Tocris (UK). AM251, ACEA, Latrunculin A, Jasplakinolide were from Sigma-Aldrich (UK).

Ratiometric imaging assay for nanoscale clustering

**Rationale (see Fig. 2A):** A protein of interest (red circles) is labeled with a monoclonal primary antibody (grey), which is then labeled with a mixture of two secondary antibodies respectively conjugated to a FRET donor and acceptor dye. The amount of FRET between the fluorophores is dependent on the average distance between them, which in turn reflects the average distance between the molecules of the protein of interest; namely, a more or less clustered distribution will result in a smaller or larger average distance and therefore more or less FRET as visualized by the acceptor/donor ratio $R_{A/D}$.

**Protocol:** After treatment, coverslips were fixed with 4% PFA in PBS for 15-20min at room temperature (RT) and permeabilized in 0.2% Triton-X100 in PBS supplemented with 5% horse serum for 10min. Subsequent incubations were carried out in the permeabilization buffer. Coverslips were incubated with an appropriate primary antibody for 60min at RT, washed 4 times in PBS and incubated with a 1:1 mixture of donor- and acceptor-conjugated secondary antibodies at a concentration of 0.3µg/ml each for 60min at RT. AF568 and AF633/AF647/DyLight649 were used as donor and acceptor respectively. Coverslips were then mounted in mounting medium (Southern), allowed to dry for 30min at RT and imaged on a Zeiss LSM710 microscope equipped with a standard set of lasers through a 63x oil objective. Excitation wavelengths were 488, 543 and 633nm. Bandpass filters were set at 500-550 (AF488), 560-615 (AF568) and 650-720nm (AF633, AF647, DyLight649). Image acquisition was typically carried out at the 12-bit rate. Settings were optimized to ensure appropriate dynamic range, low background and sufficient signal/noise ratio. To quantify $R_{A/D}$, 20µm-long regions of interest (ROI) were selected alongside proximal dendrites of neurons manifesting pyramidal morphology, and the ratio between acceptor and donor intensity was quantified after appropriate background subtraction. 3-5 ROI/image were chosen for analysis. To quantify synapse-specific $R_{A/D}$, puncta of fluorescence were individually processed in the same manner.

Sensitized emission FRET imaging

This was carried out essentially as described before (Glebov and Nichols, 2004), with AF568 and AF633 as a donor-acceptor pair. Briefly, FRET values in stretches of dendrites were calculated in samples labeled with both fluorophores, corrected for spectral bleedthrough, and divided by intensity in either donor or acceptor channel to yield FRET ratios.

Optogenetic stimulation

Primary neuronal cultures were sparsely transfected with ChR2-YFP, a gift from K. Deisseroth (http://www.optogenetics.org), at 7DIV using Effectene (QIAGEN, UK). At 15-17DIV cultures were prepared 2h before photostimulation by supplementing the culture medium with additional antioxidants (3.2µM glutathione (Fisher Scientific, UK), 77nM superoxide dismutase, 10nM catalase, 100µM Trolox, 110µM Vitamin C (all from Sigma-
Aldrich, UK). 12-well culture plates were placed directly on top of collimator-topped blue LEDs (Royal Blue Luxeon K2, Philips LumiLEDs; 455 ± 10 nm; ~1 mW mm⁻² at the coverslip surface), powered by a DC/DC LED driver (Recom) and controlled by a digital I/O device (USB-6501, National Instruments) and custom-written software. Photostimulation was carried out for 48 hours using two different paradigms: a ‘1Hz burst’ stimulation pattern consisting of 5 flashes at 20Hz delivered every 5s or ‘1Hz sparse’ stimulation pattern consisting of 1 flash every second. In both cases, flashes were delivered at 40% LED intensity for 20ms, ensuring that at least one spike occurred per flash. Control plates were loosely covered in tin foil to avoid stimulation from neighboring culture dishes and placed on non-functional LEDs.

Quantification of synapse-specific correlation
Coverslips were processed as for ratiometric imaging. AF405-conjugated anti-rabbit secondary antibody was used to visualize RIM, CASK and Ca²⁺.1. To maximize the dynamic range, image acquisition was carried out at the 16-bit rate. To reveal AZ morphology, images were thresholded in ImageJ using the “Moments” algorithm. Individual AZs were selected using the “Analyze particles” plugin. To exclude the overlapping AZs from the sample, the particle size selection criteria were set to 0.1-2µm². Intensities in all three channels were then quantified in a particle-specific manner, yielding synapse-specific values for the level of the protein of interest as well as the donor and acceptor intensities for Bsn. After appropriate background subtraction, Rₐ/D was quantified for each synapse and RIM (or CASK, or Ca²⁺.1) intensity was plotted against Rₐ/D. Before pooling, the data was normalized to maximal values in fixed-cell imaging experiments or median value in live-fixed correlative imaging experiments.

Anti-Synaptotagmin-1 antibody uptake assay
Antibody uptake assay was performed essentially as described before. Neurons were labeled with 1/100 mouse monoclonal antibody (isotype IgG3) against the luminal/extracellular domain of Synaptotagmin-1 (Syt-1) in culture medium at 37°C. To label the spontaneously cycling synaptic pool, labeling was carried out for 20’ in the presence of 2µM TTX. Coverslips were acid-washed to remove the surface label, fixed, permeabilized and stained for Syt-1 and Rₐ/D; Syt-1 and Bsn signals were visualized using the appropriate isotype-specific secondary antibodies (AF488-anti-IgG3, AF568-anti-IgG2a and AF647-anti-IgG2a respectively).

Live imaging of presynaptic vesicle cycling
Neurons were sparsely transfected at 7DIV with CMV::SypHy (a gift from Leon Lagnado (Addgene plasmid # 24478)) for imaging presynaptic vesicle release using Effectene (QIAGEN, UK), and imaged at 17-21DIV. For imaging and stimulation the neuronal growth medium was replaced with HEPES buffered saline (HBS; 139mM NaCl, 25mM KCl, 10mM HEPES, 10mM D-Glucose, 2mM CaCl₂, 1.3mM MgCl₂; pH7.3, 290 mOsm) with 2,3-Dioxo-6-nitro-1,2,3,4-tetrahydrobenzo[fl]quinoxaline-7-sulfonamide (NBQX), 0.025mM amino-5-phosphonovaleric acid (APV) and 6-Imino-3-(4-methoxyphenyl)-1(6H)-pyridazinebutanoic acid hydrobromide (Gabazine) (all Tocris, UK). A field stimulation insert with platinum electrodes (RC-37WS, Warner Instruments, USA) was placed inside the dish. Neurons
were imaged on an inverted Olympus IX71 microscope, equipped with a 60x/1.42 NA oil objective. SypHy was excited using a 470nm wavelength LED (CoolLED, UK) and imaged using the appropriate filters (Excitation: 470+/−20nm; Dichroic: 495 nm long-pass; Emission: 525+/−25nm; Chroma Filters, USA). Time-lapse images were acquired at the frequency of approximately 13Hz using an Evolve 512 EMCCD camera (Photometrics, USA) controlled by Slidebook software (Intelligent Imaging Innovations, USA). Slidebook was also used to trigger a frame-locked delivery of a 1ms, 80V pulse 40 times at 50ms intervals (i.e. 20Hz) for SypHy, where each pulse approximates a single action potential (AP)(Zhao et al., 2011). Several regions in each dish were imaged, within a maximum imaging period of 1 hour.

Images were analyzed using custom written Matlab codes (Mathworks), in which background-subtracted fluorescence intensity values were calculated from the mean of a square 8x8 pixel ROI selected for each presynaptic bouton identified by SypHy fluorescence. ΔF values were calculated as the change in signal intensity from the mean of the baseline (all frames preceding the stimulus), and the peak response was defined as the maximum ΔF within 70 frames of the stimulus. Only synapses in which the peak ΔF response was greater than 3 standard deviations of the baseline were analyzed. ΔF values were normalized to the mean peak response of all synapses in each image before pooling.

**Live imaging of presynaptic calcium influx and agatoxin blockade**

A synapsin::SyGCaMP6F plasmid was constructed by replacing GCaMP3 in SyGCaMP3 (Nikolaou et al., 2012) with GCaMP6F (a gift from Douglas Kim, (Addgene plasmid #40755)) and replacing the promoter. Transfection with this construct and imaging of presynaptic calcium influx were carried out as for imaging of vesicle cycling using SypHy, with the exception that the stimulation applied was 10APs at 20Hz, repeated three times one minute apart. After this, half of the imaging media was removed and replaced with the same volume of media containing 200nM ω-agatoxin IVA (Alomone, Israel) and mixed by pipetting up and down, yielding a final concentration of 100nM. After 5 minutes of incubation in agatoxin the three 10AP 20Hz stimuli were repeated. Analysis was also carried out as above. Responses were averaged across the three trials and ΔF values were normalized to the baseline fluorescence of each ROI to give ΔF/F values.

**Correlative live-fixed imaging of presynaptic structure and function**

The schematic of the approach is presented in Figure S6. After live imaging of SypHy responses, cells were fixed and stained for ratiometric imaging of Bsn as above, with the addition of chicken anti-GFP primary antibody (Abcam, UK) and anti-chicken AF488 conjugated secondary antibody to amplify the SypHy signal. Dishes were mounted with Ibidi mounting medium (Ibidi, Germany). Regions that had previously been imaged live were relocated on an inverted Nikon Eclipse Ti spectral confocal microscope equipped with a 60x/1.40 NA oil objective and NIS Elements software. Excitation wavelengths were 488, 561 and 636nm. Emission bandpass filters were set at 500-550nm (AF488), 570-620nm (AF568) and 662-737nm (AF633). Optical sections were taken at
0.1µm steps to create z-stacks, from which maximum projections were generated in ImageJ. To calculate $R_{A/D}$ values for all synapses in the image, thresholding and particle analysis was carried out as for synapse-specific quantification. From this, the median $R_{A/D}$ value of each image was calculated for later use in normalization. Fixed images were also analyzed using custom written Matlab codes (Mathworks). Corresponding live SypHy images and fixed GFP images were first registered by manual selection of landmarks in both images followed by affine transformation of the fixed image. ROIs selected on live SypHy images were then scaled and overlaid on the registered fixed GFP image for manual confirmation of the position. The registered GFP image was then thresholded using the ‘Moments’ algorithm in ImageJ to generate a mask, which was applied to registered AF568 and AF633 images to ensure that neighboring synapses were not included when ROIs were applied to these channels. Any ROIs falling outside of the GFP positive mask were also excluded. Intensity and $R_{A/D}$ values for each ROI were calculated from the masked images. $R_{A/D}$ values were normalized to the mean $R_{A/D}$ of all synapses in the image before pooling.

**Single color STORM imaging**

Samples were processed in the same manner as for ratiometric imaging, except that the only secondary antibody used was the AF647-conjugated anti-mouse at 1µg/ml. Samples were then incubated in the STORM imaging buffer with MEA (for recipe see http://www.nikoninstruments.com/en_GB/Products/Microscope-Systems/Inverted-Microscopes/N-STORM-Super-Resolution/(brochure)) in the following manner. The 22nm square coverslips were lifted from their dish, excess buffer was removed by blotting with paper and the coverslips were upturned onto 30µl of the imaging buffer placed on a glass slide. Coverslips were secured on the slide using Vaseline, and excess imaging buffer was blotted out with paper. With glass-bottomed dishes, 300µl imaging buffer was added directly onto the dishes. In both cases, imaging was performed immediately after mounting. Imaging was performed on an N-STORM Nikon Ti-E TIRF inverted microscope equipped with an Andor iXon DU897 EMCCD camera and a Perfect Focus system, running on the NIS Elements software. Imaging was carried out at the 12-bit rate through a 100x oil objective with a 647nm laser. The ROI was set at 128x128 pixels, with the pixel size of 160nm. The ROI was pre-bleached at 100% laser power until individual blinking events could be clearly distinguished, and subsequently imaged in the near total internal reflection fluorescence (TIRF) mode for 2000-5000 frames at a rate of 50frames/s. The resulting data was exported into the TIFF format. The positions of individual molecules were determined using the QuickPALM plugin (Henriques et al., 2010) with the default settings on the ImageJ platform.

**Dual color STORM imaging**

Dual color imaging was performed on a custom-built setup as described before (Winterflood et al., 2015). In brief, a 473 nm laser (100 mW, Laserglow Technologies) was used for activation and a 643 nm laser (150 mW, Toptica Photonics) for imaging. Emission light was filtered by two bandpass emission filters (700/75 nm, Chroma) and a longpass dichroic beam splitter (690 nm, AHF Analysetechnik) was used to split the emission light. The imaging buffer consisted of 0.1 M MEA/0.2 M Tris, pH 8.0 with 5 % (w/v) glucose, 0.25 mg/ml glucose-oxidase and 20 µg/ml catalase. Imaging was performed in objective-type near-TIRF mode. A minimum of 20000 frames with an exposure time of 20-33ms was
recorded. An image-correlation based drift-correction was employed. All data analysis was performed in MATLAB (Mathworks). To differentiate between AF647 and CF680 single molecule fluorescence profiles, the normalized intensity ratio was calculated for all localization pairs for the color-assignment using $r=(I_l-I_s)/(I_l+I_s)$, where $I_l$ and $I_s$ are the fluorescence intensities determined by maximum-likelihood fitting for the long and short wavelength channels respectively. The cut-off criteria (green and red lines in Fig. S1C) were then selected manually for each image to minimize the conflation of the fluorophores' identities. NNLEs were measured using a knnsearch function in Matlab. Data was then processed using the off-gap method to remove spurious localizations resulting from photoblinking or fluorophore reactivation (Annibale et al., 2011; Williamson et al., 2011). Areas that unambiguously conformed to the previously reported morphological criteria for AZ in a "face-on" (i.e. flat) configuration (Dani et al., 2010; Schikorski and Stevens, 1997) were manually selected for further analysis. Each experiment was repeated thrice.

**Correlation analysis of spatial distribution in STORM data**

Selected AZ (15/condition, 3 experiments) were processed in Matlab into three bins at 20, 50 and 100nm/pixel respectively, resulting images were exported in TIFF format and further processed in ImageJ using the Colocalization Test plugin. To determine the non-randomness of the distribution, the observed correlation coefficient was then compared using the Costes method (Costes et al., 2004) to the mean value from 100 randomized iterations. The outcomes were then classed into three categories: if the observed correlation coefficient value was significantly lower or higher than the randomized mean, the distributions were considered non-randomly exclusive or inclusive respectively, while the value not significantly different from the randomized mean was considered indicative of a random distribution. Fisher’s exact test was subsequently used to assess the statistical significance of the differences in proportions of AZ belonging to these categories.

**Quantitative clustering analysis**

The list of x and y coordinates of individual molecules was analyzed for clustering using Ripley’s K-function and Getis and Franklin’s local point pattern analysis. For this process, 3x3um representative square regions were chosen for analysis. In Ripley’s K-function, concentric circles are drawn around each point and the number of other points encircled is then counted. This value is then normalized based on the total molecular density such that the K-function at a circle radius $r$ is calculated as:

$$K(r) = A \sum_{i=1}^{n} \sum_{j=1}^{n} \frac{\delta_{ij}}{n^2}$$

where $\delta_{ij} = \begin{cases} 1 & \text{if } d_{ij} < r \\ 0 & \text{else} \end{cases}$

Where A is the analyzed area, n is the total number of points within that ROI and $d_{ij}$ is the distance between two points i and j. The K-function is then linearized such that it scales with circle radius rather than area, giving the L-function:

$$L(r) = \sqrt{\frac{K(r)}{\pi}}$$
A completely spatially random (CSR) distribution of points will have $L(r) = r$ at all $r$. We therefore subtract $r$ and plot $L(r) - r$ versus $r$. In this case a CSR distribution is a straight line at $L(r) - r = 0$. Positive values of $L(r) - r$ indicate the spatial scale of clustering: the greater the value of $L(r) - r$ the more clustered the distribution.

In order to generate the cluster heat maps and extract cluster parameters, Getis and Franklin’s local point pattern analysis was used. Here, the value of $L(r)$ is calculated for each point in the ROI individually for a single specified spatial scale (circle radius). A scale of 50 nm was selected. The equation therefore becomes:

$$L(50) = \sqrt{\frac{A \sum_{i=1}^{n} \left( \frac{\delta_{ij}}{n} \right)}{\pi}}$$

where $\delta_{ij} = \begin{cases} 1 & \text{if } d_{ij} < 50 \\ 0 & \text{else} \end{cases}$

Values of $L(r)$ at each molecule location were then interpolated onto a 5nm resolution grid to generate the cluster maps. To extract cluster parameters, points and areas of this map were designated as being within a cluster if the value of $L(r)$ is above a binary threshold ($L(r) > XX$). This method has previously been demonstrated for the analysis of clustering in single-molecule dSTORM data sets (Williamson et al., 2011).

**Statistics**

Statistical analysis was carried out using GraphPad Prism5.0. Sample distribution was assessed using D’Agostino and Pearson’s omnibus normality test; to assess the significance of differences between datasets, Mann-Whitney test was used unless noted otherwise. Error bars indicate 10-90 percentile range unless noted otherwise. ***P<0.001, **P<0.01, *P<0.05.

**Supplementary References**


