Piccolo modulation of Synapsin1a dynamics regulates synaptic vesicle exocytosis

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Abstract

active zones are specialized regions of the presynaptic plasma membrane designed for the efficient and repetitive release of neurotransmitter via synaptic vesicle (SV) exocytosis. Piccolo is a high molecular weight component of the active zone that is hypothesized to participate both in active zone formation and the scaffolding of key molecules involved in SV recycling. In this study, we use interference RNAs to eliminate Piccolo expression from cultured hippocampal neurons to assess its involvement in synapse formation and function. Our data show that Piccolo is not required for glutamatergic synapse formation but does influence presynaptic function by negatively regulating SV exocytosis. Mechanistically, this regulation appears to be calmodulin kinase II-dependent and mediated through the modulation of Synapsin1a dynamics. This function is not shared by the highly homologous protein Bassoon, which indicates that Piccolo has a unique role in coupling the mobilization of SVs in the reserve pool to events within the active zone.

Introduction

Presynaptic boutons are sophisticated compartments designed for the rapid, regulated release of neurotransmitter via synaptic vesicles (SVs). SVs release neurotransmitter at specialized sites called active zones (AZs; Schoch and Gundelfinger, 2006), which appear by electron microscopy as domains containing docked SVs intertwined with tufts of electron-dense material. These electron-dense tufts are thought to represent the cytoskeletal matrix assembled at the AZ (CAZ), which is comprised of a collection of multidomain scaffold proteins including ELKS (ERC/CAST), Liprin1α, RIMs, RIMBP5s, Bassoon, and Piccolo. The CAZ has hypothesized roles in structurally defining the AZ, maintaining it in register with postsynaptic structures, regulating its size, and recruiting key molecules involved in SV exocytosis such as Munc13 and voltage-gated calcium channels (Garner et al., 2000; Schoch and Gundelfinger, 2006; Fejtova and Gundelfinger, 2006). With the notable exceptions of RIMs and Munc13, our understanding of the roles played by individual CAZ proteins is very limited. In particular, the functions of the two largest CAZ proteins, Piccolo and Bassoon, have remained elusive because of their enormous sizes (>400 kD). Both are expressed very early during neuronal differentiation (Cases-Langhoff et al., 1996; Zhai et al., 2000) and are among the first proteins to arrive at newly forming synapses (Friedman et al., 2000; Zhai et al., 2000; Shapira et al., 2003), which implicates them in nascent synapse formation (Ziv and Garner, 2004; Waites et al., 2005). These multidomain proteins are composed of two N-terminal zinc finger motifs, three coiled-coiled regions and, in the case of Piccolo, a PDZ and two C2 domains (Fig. 1 A; tom Dieck et al., 1998, 2005; Altrock et al., 2003). The loss of Piccolo knockout mice, in which photoreceptor ribbon synapses are grossly malformed (e.g., ribbons detached from the AZ), support a structural role for this protein (tom Dieck et al., 1998, 2005; Altrock et al., 2003). The loss of

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Abbreviations used in this paper: ANOVA, analysis of variance; AZ, active zone; CaMKII, CaM-dependent kinase II; CAZ, cytoskeletal matrix assembled at the active zone; DIV, days in vitro; MAP2, microtubule-associated protein 2; Pr, probability of release; PSD, postsynaptic density; RRP, readily releasable pool; shRNA, short hairpin RNA; SV, synaptic vesicle; TRP, total recycling pool; VAMP2, vesicle-associated membrane protein 2.

The online version of this paper contains supplemental material.

http://doi.org/10.1083/jcb.200711167

Supplemental material can be found at:
Bassoon also reduces excitatory postsynaptic currents and increases the percentage of silent synapses in cultured hippocampal neurons; however, no overt structural defects are seen at these central nervous system synapses (Altrock et al., 2003). This milder phenotype could be caused by a more ancillary role for Bassoon at conventional synapses or by functional redundancy of another CAZ protein such as Piccolo. Addressing these issues using conventional knockout strategies has been hampered by the very large sizes of the genes encoding Piccolo and Bassoon (>350 kb) and the existence of alternatively spliced transcripts and uncharacterized 5’ promoter regions (Cases-Langhoff et al., 1996; tom Dieck et al., 1998; Winter et al., 1999; Fenster and Garner, 2002; Altrock et al., 2003). As a case in point, the Bassoon knockout mouse is not a genetic null but rather a deletion of two large central exons from the Bsn gene, leaving the N and C termini intact. Careful analysis of these mice has demonstrated that these domains, including the N-terminal Zinc finger motifs and C-terminal coiled-coil domain, are still expressed and synaptically localized, which suggests that the true phenotype of Bassoon loss could be masked by the continued expression of these fragments.

In the present study, we have overcome these limitations by using interference RNA to completely eliminate the expression of Piccolo in cultured hippocampal neurons. Our data reveal that although Piccolo is not essential for the formation of glutamatergic synapses, it does play a functional role in SV recycling. Specifically, synapses lacking Piccolo exhibit enhanced SV exocytosis rates, apparently caused by alterations in the activity-dependent dynamics of Synapsin1a at presynaptic boutons. These defects are not detected at synapses lacking Bassoon, which indicates that Piccolo has a specific role in coupling events at the AZ with the regulated recruitment of SVs from the reserve pool.

### Results

#### Design of short hairpin RNAs (shRNAs) that selectively and efficiently knock down Piccolo

To eliminate the expression of Piccolo in developing neurons, we generated several shRNAs against sequences situated in the N terminus of Piccolo (Exon 1; Fig. 1 A; Fenster and Garner, 2002). This region has been shown to encode segments of the highest molecular weight Piccolo isoforms (~560 kD; Fig. 1 A; Fenster et al., 2000; Fenster and Garner, 2002) shown previously to localize to presynaptic AZs (Cases-Langhoff et al., 1996; Fenster et al., 2000; Zhai et al., 2000). After developing and testing two such shRNAs – Pclo6 and Pclo28 – in HEK cells, we assessed their efficacy in neurons using plasmid-based transfection (Fig. S1, available at http://www.jcb.org/cgi/content/full/jcb.200711167/DC1). Pclo28 was slightly more effective than Pclo6 at eliminating Piccolo expression, and was thus used for all further experiments presented in this paper. However, key experiments were repeated with Pclo6 to verify that the observed phenotypes were not caused by off-target effects (Fig. S2).

Given the low efficiency and potential overexpression associated with plasmid-based transfection, we next evaluated whether Pclo28 introduced via lentivirus would similarly eliminate Piccolo in cultured neurons. This was accomplished by subcloning both an EGFP-Synapsin1a reporter gene (for labeling presynaptic boutons expressing Pclo28; Fig. S1) and the H1-driven shRNA into the FUGW vector (Fig. 1 B; Lois et al., 2002) to create LV/EGFP-Synapsin or LV/EGFP-Synapsin/Pclo28. As an initial measure of knockdown efficacy, lysates from 14-d-in-vitro (DIV) hippocampal neurons superinfected with lentivirus, in order to ensure nearly 100% infectivity, were Western blotted and probed with antibodies against Piccolo and several other neuronal proteins (Fig. 1 C). We observed a dramatic and selective loss of Piccolo in lysates from cells infected with Pclo28 and no effect on the levels of β-tubulin, Synaptophsin, Bassoon, or EGFP-Synapsin1a (Fig. 1 C).

We next assessed whether lentiviral expression of Pclo28 reduced synaptic Piccolo. Hippocampal neurons infected with LV/EGFP-Synapsin or LV/EGFP-Synapsin/Pclo28 were immunostained after 14 DIV with antibodies against Piccolo and Bassoon. In control cells, EGFP-Synapsin exhibited a punctate pattern along microtubule-associated protein 2 (MAP2)-positive dendritic profiles (Fig. 1 D), with ≥90% of these puncta immunopositive for Piccolo (Fig. 1, D and F) or Bassoon (Fig. 1, E and G). In contrast, hippocampal cultures infected with LV/EGFP-Synapsin1a/Pclo28 exhibited a dramatic loss of Piccolo immunoreactivity at EGFP-Synapsin1a clusters, with <3% of clusters along dendritic profiles containing Piccolo (Fig. 1 F). Importantly, this shRNA had no effect on Bassoon protein levels or clustering at EGFP-Synapsin1a sites (Fig. 1, C, E, and G). Collectively, these data indicate that Pclo28 efficiently and selectively eliminates Piccolo from synapses.

#### Loss of Piccolo does not affect synapse formation or morphology

The appearance of axonal EGFP-Synapsin1a and Bassoon clusters in the absence of presynaptic Piccolo expression indicates that Piccolo is not necessary for synapse formation. To evaluate this hypothesis, we immunostained cultured neurons infected with either LV/EGFP-Synapsin1a or LV/EGFP-Synapsin1a/Pclo28 with antibodies against several key presynaptic (Synaptophsin, RIM1α, and Munc13-1) and postsynaptic (PSD-95 and NR1) proteins. As shown in Fig. 2, each exhibited a punctate pattern that reliably colocalized with EGFP-Synapsin1a clusters along dendritic profiles, regardless of whether Piccolo was present (Fig. 2 A) or absent (Fig. 2 B). These data indicate that the loss of Piccolo does not disrupt the synaptic targeting of multiple key pre- and postsynaptic proteins.

To assess whether the loss of Piccolo affected synapse ultrastructure, we developed a strategy for visualizing boutons expressing Pclo28 by EM. This was achieved by replacing EGFP-Synapsin1a in the pZOff vector with vesicle-associated membrane protein 2 (VAMP2) tagged at its C terminus with HRP (VAMP2-HRP; Fig. 3 A). VAMP2 is an essential transmembrane component of SVs (Elferink et al., 1989; Sudhof et al., 1989), and in VAMP2-HRP, the C-terminal HRP is situated within the vesicle lumen. Upon reaction with hydrogen peroxide and DAB, HRP forms an electron-dense precipitate, enabling visualization of vesicles within boutons of neurons expressing...
Gross morphological differences were detected between unlabeled synapses, those expressing only V AMP2-HRP, and those expressing V AMP2-HRP/Pclo28 (Fig. 3, E and F). Furthermore, when synapses were carefully quantified for bouton area, AZ/PSD length, docked SVs/AZ length, and SV density (number of vesicles per bouton area), no morphological differences were found (Table I). These data strongly suggest that Piccolo is not essential for the structural assembly of excitatory asymmetrical synapses.

Synapses lacking Piccolo exhibit faster rates of SV exocytosis

Although synapses still form in the absence of Piccolo, its large size and multiple binding partners suggest that it may be functionally important for SV recycling. We thus used the styryl FM dyes (Cochilla et al., 1999) to analyze presynaptic function. We first examined whether boutons lacking Piccolo were presynaptically active. This was accomplished by labeling the total recycling pool (TRP) of vesicles with FM4-64 (90 mM pZOff/V AMP2-HRP). To verify that V AMP2-HRP did not affect Piccolo down-regulation, cultures expressing pZOff/V AMP2-HRP or pZOff/V AMP2-HRP/Pclo28 were immunostained with antibodies against Piccolo and HRP at 6 DIV (Fig. 3 B). Under these conditions, Piccolo immunoreactivity was detected in the axons of untransfected neurons or those expressing V AMP2-HRP alone but not in axons of neurons expressing Pclo28. These data indicate that Pclo28 shRNA was not hampered by coexpression of V AMP2-HRP.

To visualize synapses, hippocampal neurons electroporated at the time of plating with pZOff/V AMP2-HRP or pZOff/V AMP2-HRP/Pclo28 were fixed and processed for EM at 14 DIV. In both cases, presynaptic boutons expressing V AMP2-HRP were readily identified based on the presence of electron-dense SVs (Fig. 3 D). These labeled SVs were easily distinguishable from both unlabeled SVs and the 80-nm dense core vesicles hypothesized to carry Piccolo and Bassoon to nascent synapses (Figs. 3 C and S3, available at http://www.jcb.org/cgi/content/full/jcb.200711167/DC1; Zhai et al., 2001). Qualitatively, no gross morphological differences were detected between unlabeled synapses, those expressing only V AMP2-HRP, and those expressing V AMP2-HRP/Pclo28 (Fig. 3, E and F). Furthermore, when synapses were carefully quantified for bouton area, AZ/PSD length, docked SVs/ AZ length, and SV density (number of vesicles per bouton area), no morphological differences were found (Table I). These data strongly suggest that Piccolo is not essential for the structural assembly of excitatory asymmetrical synapses.
KCl for 60 s; Pyle et al., 2000) in neuronal cultures infected with either LV/EGFP-Synapsin1a or LV/EGFP-Synapsin1a/Pclo28. At boutons with (EGFPSyn) or without (EGFPSyn-Pclo28) Piccolo, >80% of EGFP-Synapsin1a clusters colocalized with FM4-64 puncta (Fig. 4, A and B). These data indicated that synapses lacking Piccolo were indeed presynaptically functional and no more likely to be silent than control boutons. We next compared the total FM fluorescence intensity at boutons containing or lacking Piccolo to determine the relative sizes of the TRP. No difference was observed in the mean intensity of FM4-64 fluorescence (Fig. 4 C), which indicates that TRP size was unaffected by the absence of Piccolo. To evaluate whether the loss of Piccolo led to changes in SV exocytosis, we compared the destaining kinetics of the TRP using both 10- and 5-Hz electrical stimulation (Fig. 4, D and E). Intriguingly, boutons lacking Piccolo destained more quickly than those expressing only EGFP-Synapsin1a (two-way analysis of variance [ANOVA]; P < 0.0001 for both conditions;
Figure 3. **Loss of Piccolo does not affect synapse ultrastructure.** (A) Schematic diagram of the pZOff/VAMP2-HRP vector used to express VAMP2-HRP with or without Pclo28 shRNA. (B) Hippocampal neurons electroporated at the time of plating with pZOff/VAMP2-HRP or pZOff/VAMP2-HRP/Pclo28 and immunostained at 6 DIV with HRP (green) and Piccolo (red) antibodies. Note the lack of Piccolo immunoreactivity in Pclo28-expressing axons. (C) Transmission EM micrograph of a VAMP2-HRP–positive bouton containing both clear-centered (unlabeled SV) and dark-centered SVs (labeled SV). HRP-labeled vesicles are easily distinguished from 80-nm DCVs and other labeled structures. (D) Synaptic boutons from VAMP2-HRP–expressing (labeled bouton) and untransfected (unlabeled bouton) neurons. Arrowheads denote synaptic junctions, identified based on electron-dense PSDs. (E) An excitatory synapse formed between a pZOff/VAMP2-HRP/Pclo28 (Pclo28)-transfected presynaptic bouton and a postsynaptic spine. (F) An excitatory synapse formed between a pZOff/VAMP2-HRP (control) transfected presynaptic bouton and a postsynaptic spine.
Table I. Quantitative ultrastructural analysis of synapses with or without Pclo28

| Parameter                        | Control         | Pclo28          |
|----------------------------------|-----------------|-----------------|
| AZ length (nm)                   | 0.258 ± 0.142   | 0.229 ± 0.102   |
| Docked SVs/AV length (n/μm)      | 13.87 ± 6.26    | 14.29 ± 5.30    |
| SV density (n/μm²)               | 163.8 ± 58.36   | 169.0 ± 49.02   |
| Percentage of SVs labeled        | 20.4 ± 12.5%    | 24.5 ± 14.8%    |

All data are from boutons with at least two clearly labeled synaptic vesicles opposed to a distinct synaptic junction. All parameters were measured by an observer blind to phenotype. n = 64 boutons with 66 active zones for control and 53 boutons with 55 active zones for Pclo28. Data are shown ± standard deviation. t tests revealed no significant differences, with no P < 0.05.

Fig. 4, D and E), which indicates that Piccolo is a negative regulator of SV exocytosis.

Piccolo could regulate SV exocytosis by influencing vesicle docking and/or fusion with the AZ plasma membrane or by controlling SV translocation from the reserve to readily releasable pool (RRP). To explore possible changes in SV docking and/or fusion, we evaluated whether the size of the RRP of SVs or the release probability (Pr) were modified in the absence of Piccolo (Fig. 4, F and G). The former represents the subpopulation of SVs that are docked at the AZ plasma membrane and poised to undergo fusion with the arrival of an action potential, whereas the latter is a measure of the probability that SVs in the RRP will undergo fusion. Two methods were used to estimate RRP size: one using hypertonic sucrose (500 mM, ~800 Osm) and the other using a weak electrical stimulus (2 Hz for 30 s; Pyle et al., 2000). Under both conditions, we found no significant difference in RRP size between wild-type synapses and those lacking Piccolo (t test, P > 0.5; Fig. 4 F).

To determine Pr, we measured the destaining kinetics of boutons under conditions that stimulate RRP release, as described previously (Ryan et al., 1993; Pyle et al., 2000; Mozhayeva et al., 2002; Sankaranarayanan et al., 2003). Again, we found no significant difference in release probability between wild-type boutons and those lacking Piccolo (t test, P > 0.5; Fig. 4 G). Collectively, these data indicated that the increased SV exocytosis rates observed at 5 and 10 Hz were not caused by increased RRP size or Pr.

Piccolo modulates the dispersion kinetics of Synapsin1a

Based on the RRP experiments, we concluded that Piccolo’s negative regulation of SV exocytosis was unlikely to occur at the level of SV priming or fusion, as described for RIM1α and Munc13 (Augustin et al., 1999; Schoch et al., 2002; Weimer and Richmond, 2005; Gracheva et al., 2006; Weimer et al., 2006), but rather at an earlier step, such as translocation of SVs from the reserve pool to the RRP. To date, the only protein that has been implicated in regulating SV translocation is the SV-associated phosphoprotein Synapsin (De Camilli et al., 1990; Greengard et al., 1993; Ryan et al., 1996; Chi et al., 2003). Mechanistically, Synapsin is hypothesized to mediate the clustering and retention of SVs within boutons via its ability to cross-bridge and tether them to the presynaptic actin/spectrin cytoskeleton (Greengard et al., 1993; Ceccaldi et al., 1995). Activity-dependent phosphorylation/dephosphorylation of Synapsin by CaM-dependent kinase II (CaMKII), protein kinase A, MAPK, and protein phosphatases 2A and 2B appears to regulate the exocytosis kinetics of SVs, presumably by regulating Synapsin binding to SVs and/or the actin cytoskeleton (Greengard et al., 1993; Ryan et al., 1993; Jovanovic et al., 2000, 2001; Pyle et al., 2000; Mozhayeva et al., 2002; Chi et al., 2003; Sankaranarayanan et al., 2003). Importantly, Synapsin has also been shown to undergo an activity-dependent dispersion away from presynaptic boutons (Chi et al., 2001), and the rate of dispersion is linked to its phosphorylation state (Chi et al., 2001, 2003). However, the precise relationship between Synapsin phosphorylation, dispersion, and SV exocytosis appears complex and has not been fully characterized.

To explore whether the loss of Piccolo impacts SV translocation via alterations in the properties of Synapsin, we evaluated whether the association and/or dispersion kinetics of Synapsin1a within presynaptic boutons were altered. Here, we took advantage of the fact that our shRNA expression vector contained EGFP-Synapsin1a. Initially, we used antibodies against Synapsin to verify that the lentiviral system did not lead to a gross overexpression of Synapsin at individual boutons. Surprisingly, we found no increase in Synapsin immunostaining at boutons expressing EGFP-Synapsin1a (Fig. 5 A), which indicates that endogenous Synapsin expression is down-regulated in the presence of EGFP-Synapsin to maintain similar total levels per bouton. Next, we quantified and compared the synaptic levels of EGFP-Synapsin1a in boutons of control neurons and those expressing Pclo28. We found no discernable differences in fluorescence intensity when cultures were fixed with 4% paraformaldehyde (Fig. 5 B). However, when cultures were fixed with methanol, which extracts soluble proteins, we observed a significant decrease (t test, P < 0.0001) in the intensity of EGFP-Synapsin1a at boutons lacking Piccolo (Fig. 5 B). This suggested that EGFP-Synapsin1a was not as tightly associated with SVs and/or actin-related structures in the absence of Piccolo but rather was shifted into a more soluble fraction. This concept was supported by FRAP experiments designed to monitor the steady-state exchange kinetics of EGFP-Synapsin1a fluorescence at individual boutons (Fig. 5, C and D). Again, EGFP-Synapsin1a fluorescence recovered more quickly at boutons lacking Piccolo (τfast = 1.2 min and τslow = 11.9 min vs. τfast = 2.2 min and τslow = 33.7 min for control boutons; two-way ANOVA, P < 0.0001; Fig. 5 D), which indicates that mechanisms regulating the association of Synapsin1a with SVs and/or the actin/spectrin cytoskeleton were altered in the absence of Piccolo.

To assess whether the faster exchange kinetics of EGFP-Synapsin1a in the absence of Piccolo were caused by decreased presynaptic stability and/or association of SVs with synapses, we also examined the turnover rates of SV2, an SV integral membrane protein, in the presence and absence of Piccolo. Here, an N-terminally EGFP-tagged SV2 was subcloned into our lentiviral vectors in place of EGFP-Synapsin1a. Like EGFP-Synapsin1a, EGFP-SV2 reliably labeled presynaptic boutons, exhibiting a high degree of colocalization with FM4-64 (Fig. S4, available at
Figure 4. Synapses lacking Piccolo have enhanced rates of SV exocytosis. (A) Images of 14 DIV hippocampal neurons infected with LV/EGFP-Synapsin1a (EGFPSyn, green) or LV/EGFP-Synapsin1a/Pclo28 (EGFPSynPclo28, green) and loaded with FM4-64 (FM, red) with 90 mM K+. Note the high degree of match between EGFP-Synapsin1a and FM4-64 in both merged images. Bar, 10 μm. (B) Bar graphs quantifying the percent colocalization between EGFP-Synapsin and FM4-64 puncta in control cultures (EGFPSyn) or those lacking Piccolo (EGFPSynPclo28; n > 800 puncta per condition, two experiments). (C) Bar graph of FM4-64 fluorescence intensity at EGFP-Synapsin1a puncta comparing the relative sizes of the TRP of SVs at control synapses (EGFPSyn) and those lacking Piccolo (EGFPSynPclo28). FM intensity values at EGFPSyn-expressing boutons were normalized against those from neighboring uninfected cells to enable cross-coverslip comparison. Mean normalized FM intensity from control boutons was set to 100; that from Pclo28 boutons was ratioed against this value (n > 800 puncta, two experiments). (D and E) Destaining kinetics of the TRP at 10 Hz (D) and 5 Hz (E) comparing synapses with (EGFPSyn) and without (EGFPSynPclo28) Piccolo (n = 5 experiments per condition). (F) Bar graph comparing the size of the RRP of SVs in boutons with (EGFPSyn) or without (EGFPSynPclo28) Piccolo, as determined by either the application of 500 mM sucrose (left) or 2-Hz, 30-s stimulation (right). No significant differences were found (t test, P > 0.5). Sucrose experiments were performed twice for each condition and stimulation experiments three times. (G) Destaining kinetics of the RRP during 2-Hz, 30-s stimulation for control (EGFPSyn) boutons and those lacking Piccolo (EGFPSynPclo28; n = 5 experiments per condition). Error bars indicate SEM.
Figure 5. **Steady-state and activity-dependent dynamics of EGFP-Synapsin1a are altered at synapses lacking Piccolo.** (A, left) Images of neurons infected with EGFP-Synapsin1a (green) and immunostained with Synapsin1a antibodies (red). (A, right) Bar graph comparing the intensity of Synapsin immunoreactivity at boutons with or without EGFP-Synapsin1a. Total Synapsin levels are similar for infected and uninfected neurons. n > 100 puncta per condition. Bar, 10 μm. (B) EGFP-Synapsin1a fluorescence intensity at presynaptic boutons with (EGFPsyn) or without (EGFPsynPclo28) Piccolo after methanol versus 4% paraformaldehyde fixation (n > 800 puncta, seven fields of view per condition). (C) Time-lapse images of EGFP-Synapsin1a puncta. Single fluorescent puncta (indicated by arrows) were photobleached and their recovery was monitored over time. (C) Time-lapse images of EGFP-Synapsin1a puncta at boutons with (EGFPsyn, n = 20) or without (EGFPsynPclo28, n = 17) Piccolo. (E) Images of EGFP-SV2 puncta at boutons with (EGFPSV2) or without (EGFPSV2Pclo28) Piccolo, photobleached as in C. Arrowheads indicate the positions of EGFP puncta that are bleached during FRAP experiments.
increases in the dispersion kinetics of Synapsin1a. These data indicate that general presynaptic stability and SV retention were not significantly altered in the absence of Piccolo.

To assess whether the differences in steady-state exchange kinetics translated into activity-dependent differences in the dispersion kinetics of EGFP-Synapsin1a, we used an electrical stimulation protocol (10 Hz for 90 s) shown to promote the dispersion of Synapsin1a (Chi et al., 2001, 2003). Here, changes in the fluorescence intensity of EGFP-Synapsin1a at individual boutons were monitored both during stimulation and for the subsequent 10-min recovery period. In contrast to wild-type boutons, we observed both a more complete dispersion of EGFP-Synapsin1a and a slower rate of recovery in boutons lacking Piccolo (two-way ANOVA, P < 0.0001; Fig. 5, G and H). Intriguingly, we also observed remarkable, previously unreported heterogeneity in the extent of EGFP-Synapsin1a dispersion at individual boutons under both conditions. To evaluate whether the extent of EGFP-Synapsin1a dispersion was coupled to the rate of SV exocytosis, we simultaneously monitored the loss of EGFP-Synapsin1a and FM5-95 fluorescence from boutons during 5-Hz stimulation for 3 min. Our analysis revealed a tight correlation between the two events. At boutons with minimal EGFP-Synapsin dispersion (<10% for all time points), very little FM5-95 destaining occurred (Fig. 6, A and B). In contrast, synapses with more complete EGFP-Synapsin dispersion (>40% after 30 s of stimulation) exhibited dramatic FM5-95 destaining (Fig. 6, A and B). A statistical analysis of the final fluorescence intensity values for EGFP-Synapsin and FM after the 3-min stimulation revealed a strong correlation between the extent of EGFP-Synapsin dispersion and that of FM5-95 destaining at both control boutons and those lacking Piccolo (Fig. 6 B). A statistical analysis of the initial fluorescence intensity values for EGFP-Synapsin and FM after the 3-min stimulation revealed a strong correlation between the extent of EGFP-Synapsin dispersion and that of FM5-95 destaining at both control boutons and those lacking Piccolo (control: ρ = 0.58, P < 0.0001; Pclo28: ρ = 0.59, P < 0.0001; Fig. 6, C and D).

Although this tight correlation was observed for all synapses, the mean degree of Synapsin dispersion and FM destaining was greater at boutons lacking Piccolo, significantly shifting these correlation values to the lower right (Fig. 6, C and D). These data strongly indicate that the accelerated SV exocytosis rates observed at synapses lacking Piccolo are coupled to concomitant increases in the dispersion kinetics of Synapsin1a.

Changes in SV exocytosis and Synapsin dynamics are not observed at synapses lacking Bassoon

Piccolo shares a high degree of structural similarity with Bassoon (Fenster et al., 2000), another CAZ protein, and it is often assumed that these two proteins are functionally redundant (Fejtova and Gundelfinger, 2006; Schoch and Gundelfinger, 2006). We were thus interested in exploring whether the phenotypes seen at boutons lacking Piccolo were also observed in boutons lacking Bassoon. To this end, we generated a set of shRNAs against Bassoon and, after initial screening in the pZOff vector (unpublished data), subcloned one (Bsn16) into the LV/EGFP-Synapsin1a vector under the control of the H1 promoter. This shRNA was found to efficiently and specifically reduce the expression of Bassoon in lysates of cultures infected with LV/EGFP-Synapsin1a/Bsn16 on the day of plating and were harvested after 14 DIV (Fig. 7 A). Similarly, at the synaptic level, we observed a dramatic decrease (>95%) in the percent colocalization of Bassoon and EGFP-Synapsin1a in axons along dendritic profiles (Fig. 7, B and D). This decrease was not observed for Piccolo (Fig. 7, C and E), which indicates that the Bsn16 shRNA is specific for Bassoon.

We next examined whether boutons lacking Bassoon also exhibited changes in the dispersion and reclustering rates of EGFP-Synapsin1a. As shown in Fig. 7, synapses lacking Bassoon exhibited no detectable defects in EGFP-Synapsin dispersion or reclustering (Fig. 7 F) and no change in the destaining kinetics of FM5-95 (Fig. 7 G). Together, these observations strongly argue that the phenotypes observed for Piccolo are real and not artifacts of long-term shRNA expression. Moreover, they indicate that sequence elements unique to Piccolo play a fundamental role in negatively regulating SV exocytosis, apparently by influencing the activity-dependent dispersion of Synapsin1a.

Loss of Piccolo enhances the CaMKII sensitivity of Synapsin1a

Because the association of Synapsin1a with SVs and the actin cytoskeleton in nerve terminals appears to be regulated by CaMKII phosphorylation (Schiebler et al., 1986; Benfenati et al., 1992; Greengard et al., 1993; Stefani et al., 1997; Chi et al., 2001, 2003), we next asked whether the altered dynamics of EGFP-Synapsin1a in the absence of Piccolo could be caused by changes in its CaMKII-dependent phosphorylation. This was examined by blocking CaMKII activity with 10 μM KN62 before and during a 10-Hz, 90-s stimulation. Surprisingly, KN62 had no significant effect on the dispersion kinetics or total amount of dispersion observed for EGFP-Synapsin in wild-type boutons (two-way ANOVA, P = 0.15; Fig. 8, A and C). However, it caused a dramatic reduction in EGFP-Synapsin dispersion at Pclo28 boutons, “rescuing” dispersion to levels seen at wild-type boutons (two-way ANOVA, P < 0.0001; Fig. 8, B and C). To determine whether this decrease in dispersion was accompanied by a decrease in the SV exocytosis rate, we also assessed the impact of KN62 on the destaining kinetics of FM5-95 at Pclo28 boutons. Again, KN62 attenuated the Pclo28 phenotype, slowing the accelerated exocytosis of SVs seen in the absence of Piccolo (two-way ANOVA, P < 0.0001; Fig. 8 D). These results indicate that the effects of Piccolo on presynaptic function are
mediated by CaMKII. Moreover, they provide further evidence that the Synapsin dispersion and SV exocytosis phenotypes are mechanistically linked.

**Synapsin1a is hypophosphorylated in the absence of Piccolo**

The ability of KN62 to rescue the Pclo28 phenotypes suggested that Synapsin might be hyperphosphorylated by CaMKII in the absence of Piccolo. To test this idea, we probed Western blots of hippocampal lysates containing (EGFPSyn) or lacking (Pclo28) Piccolo with two phospho-Synapsin1a antibodies that recognize sites phosphorylated by CaMKII or MAPK (Fig. 9 A; Chi et al., 2003). In these experiments, lysates from 14-DIV cultures were analyzed either directly (untreated) or after high-K+ treatment (90 mM KCl for 2 min) to stimulate Synapsin phosphorylation. As expected, the levels of both endogenous and EGFP-Synapsin1a phosphorylation at sites 3, 4, and 5 increased in all lysates after high-K+ treatment (Fig. 9 B). However, in lysates from Pclo28 neurons, the levels of phosphorylation at these sites under both basal and high-K+ conditions were dramatically reduced (Fig. 9 B). This unexpected finding suggested that the kinase/phosphatase balance at boutons lacking Piccolo was altered and that perhaps these boutons exhibited enhanced phosphatase activity. To test this hypothesis, we screened a panel of phospho-specific antibodies against several pre- and postsynaptic proteins, including GluR1, Synaptotagmin, Rabphilin, Munc18, and Dynamin. Intriguingly, the phosphorylation levels of these proteins as assessed by Western blotting were not altered in the absence of Piccolo (Fig. 9 C), which indicates that the effect could be specific to Synapsin.

**Discussion**

In the present study, we have used interference RNAs to disrupt expression of the AZ protein Piccolo in developing hippocampal neurons. This knockdown was specific for Piccolo and had no overt effects on the expression or synaptic localization of other pre- and postsynaptic proteins, including GluR1, Synaptotagmin, Rabphilin, Munc18, and Dynamin. Intriguingly, the phosphorylation levels of these proteins as assessed by Western blotting were not altered in the absence of Piccolo (Fig. 9 C), which indicates that the effect could be specific to Synapsin.
releasing FM dyes but exhibited enhanced rates of SV exocytosis. Mechanistically, this phenotype was not caused by changes in the size or release probability of the RRP of SVs but rather to an increased dispersion of Synapsin1a from presynaptic boutons. This dispersion defect could be rescued by inhibiting CaMKII activity but was not caused by hyperphosphorylation of known Synapsin1a phospho sites by CaMKII. Importantly, synapses lacking Bassoon did not share these features. Together, these data indicate that Piccolo negatively regulates SV exocytosis by modulating Synapsin dynamics, thus potentially coupling the mobilization of SVs in the reserve pool to events at the AZ.

Roles of CAZ proteins at presynaptic boutons

So far, loss of function studies on CAZ proteins have primarily demonstrated defects in SV docking, priming, and fusion, as seen for Munc18, Munc13, and Rim1α (Augustin et al., 1999; Verhage et al., 2000; Schoch et al., 2002; Weimer and Richmond, 2005; Schoch and Gundelfinger, 2006). At present, it is unclear whether Piccolo and Bassoon are involved in these processes. Their high structural similarity and near-complete overlap of binding partners (Wang et al., 1999; Fenster et al., 2000, 2003; Kim et al., 2003; tom Dieck et al., 2005; Fejtova and Gundelfinger, 2006), and thus likely functional redundancy, will make this issue difficult to resolve until the analysis of synapses lacking both proteins is possible.

However, studies of synapses lacking either Piccolo or Bassoon are beginning to reveal unique roles for each. For example, studies of photoreceptor ribbon synapses demonstrate a crucial functional role for Bassoon in the attachment of ribbons to the arciform density and in neurotransmission at these synapses (Altrock et al., 2003; Dick et al., 2003; tom Dieck et al., 2005).
Furthermore, the increase in number of presynaptically silent synapses in Bassoon-deficient hippocampal neurons (Altrock et al., 2003) argues for a potential role of Bassoon in key aspects of vesicle release, perhaps similar to Munc13, Munc18, and Rim1a (Weimer and Richmond, 2005; Schoch and Gundelfinger, 2006). In contrast, we found no evidence that loss of Piccolo had any affect on the fraction of presynaptically silent synapses or on the size or release probability of the RRP of SVs. These data indicate that Piccolo does not directly participate in SV docking, priming, or fusion. Instead, our data reveal that Piccolo but not Bassoon influences the translocation of SVs from the reserve pool to the RRP by regulating the dynamic properties of Synapsin1a. These findings suggest that Piccolo may perform an integrative function within presynaptic boutons, coupling SV docking and fusion to the mobilization/recruitment of SVs from the reserve pool to the AZ.

Changes in the rates of SV exocytosis can be linked to altered Synapsin dynamics

Synapsins are a well-characterized family of presynaptic phosphoproteins thought to regulate the translocation of SVs from the reserve to the RRP (De Camilli et al., 1990; Greengard et al., 1993; Hilfiker et al., 1999). This concept is supported by knock-out studies showing that Synapsins are essential for maintaining the size of the reserve pool of SVs (Li et al., 1995; Ryan et al., 1996) and by dynamic imaging studies, including this one, demonstrating that they regulate SV exocytosis rates at a range of stimulus frequencies (Chi et al., 2001, 2003). Importantly, the dynamic properties of Synapsin, particularly its activity-dependent dissociation from SVs and the actin cytoskeleton, are regulated by phosphorylation (Torri Tarelli et al., 1992; Hilfiker et al., 1999).

Although Synapsin is phosphorylated by multiple kinases, several studies have implicated CaMKII as its most prominent regulator (Schiebler et al., 1986; Benfenati et al., 1992; Ceccaldi et al., 1995; Stefani et al., 1997). However, these studies used in vitro binding assays, and none assessed the direct contribution of CaMKII to either the dispersion kinetics of Synapsin or to SV exocytosis in intact neurons. Two studies that did examine these functions used serine-to-alanine phospho mutants to demonstrate that the two identified CaMKII sites were important for regulating the rates of Synapsin dispersion and SV exocytosis (Chi et al., 2001, 2003). However, neither was performed at physiological temperatures and neither used pharmacological tools to assess the role of CaMKII in these processes.

In this study, we have carefully analyzed both the relationship between Synapsin dispersion and SV exocytosis, and the regulation of these events by CaMKII in intact neurons at physiological temperature. Regarding the former, our analysis of thousands of boutons revealed a remarkable and previously unreported heterogeneity in the degree of EGFP-Synapsin dispersion and FM5-95 destaining per bouton (from >50 to 0%). From this data, we observed a tight correlation between Synapsin dispersion and SV exocytosis. Though other studies have suggested such a relationship, ours is the first to conclusively demonstrate it. We have also closely examined the role of CaMKII in regulating Synapsin dispersion. Surprisingly, blocking
CaMKII activity with KN62 did not alter Synapsin dispersion in wild-type neurons, which suggests that other kinases may mediate this activity. This concept is supported by several previous studies indicating that protein kinase A and MAPK are important regulators of Synapsin, modulating its association with SVs or actin, respectively, and, in the case of MAPK, mediating SV exocytosis via Synapsin phosphorylation (Jovanovic et al., 1996; Hosaka et al., 1999; Jovanovic et al., 2000, 2001). Intriguingly, removing Piccolo from synapses caused Synapsin dispersion to be regulated by CaMKII. Furthermore, under these conditions, the enhanced rate of SV exocytosis was also CaMKII dependent, as both phenotypes could be rescued by inhibiting CaMKII with KN62. These data support our general conclusion that Piccolo negatively regulates SV exocytosis, apparently via a CaMKII-mediated mechanism.

We have also examined the phosphorylation state of Synapsin in the absence of Piccolo. We found that under basal conditions, Synapsin was hypophosphorylated at both CaMKII and MAPK sites. Although the levels of phosphorylation at both sites increased during stimulation, absolute levels of phosphorylation were always below those found in wild-type neurons. This finding was unexpected, as we had predicted based on multiple previous studies and our own experiments with KN62 that hyperphosphorylation of Synapsin would mediate its enhanced dispersion. However, total phosphorylation levels of Synapsin may not be predictive of its dispersion kinetics, as two other imaging studies have demonstrated that Synapsin dynamics are always below those found in wild-type neurons, which suggests that other kinases may mediate this activity. This concept is supported by several previous studies indicating that protein kinase A and MAPK are important regulators of Synapsin, modulating its association with SVs or actin, respectively, and, in the case of MAPK, mediating SV exocytosis via Synapsin phosphorylation (Jovanovic et al., 1996; Hosaka et al., 1999; Jovanovic et al., 2000, 2001). Intriguingly, removing Piccolo from synapses caused Synapsin dispersion to be regulated by CaMKII. Furthermore, under these conditions, the enhanced rate of SV exocytosis was also CaMKII dependent, as both phenotypes could be rescued by inhibiting CaMKII with KN62. These data support our general conclusion that Piccolo negatively regulates SV exocytosis, apparently via a CaMKII-mediated mechanism.

In conclusion, we have found that Piccolo is not essential for excitatory synapse formation but is a negative regulator of SV exocytosis. These results indicate that one of Piccolo’s functions at the AZ is to regulate the translocation of SVs from the reserve to the RRP through modulation of Synapsin dynamics. Furthermore, our results demonstrate that this role is not shared by Bassoon, providing one of the first indications that these proteins have nonoverlapping functions. Rather puzzling is the absence of other presynaptic phenotypes, especially given the large size of Piccolo and its numerous binding partners. One likely explanation is that many of these are masked by redundant features in Bassoon, issues that will hopefully be resolved by the characterization of synapses lacking both Piccolo and Bassoon.

Materials and methods

Reagents

Antibodies against Piccolo (rabbit), Bassoon (mouse), and MAP2 (rabbit and mouse) were used as described previously (Zhai et al., 2000). HRP antibody was obtained from Advanced ImmunoChemical, Inc., GFP antibody from Roche, neomycin antibody from GeneTex, Inc., α-tubulin antibodies from Sigma-Aldrich, RIM1α and Munc13 from Synaptic Systems GmbH, PSD-95 from Affinity BioReagents, and NR1 from Millipore. Synaptophysin and Synapsin1 antibodies were obtained from Assay Designs and all phospho antibodies were obtained from PhosphoSolutions. Alexa 488 and 568 and Marina blue secondary antibodies were obtained from Invitrogen. Unless otherwise indicated, all other chemicals were obtained from Sigma-Aldrich.

Generation of pZOff vector

The pZOff vector was built on the backbone of the pEGFP-C1 vector (Clontech Laboratories, Inc.). Modifications include eliminating the multiple cloning site by digesting with BglII and BamHI and religating the vector. SalI, PvuI, and

![Diagram of Synapsin1a phospho-sites](image-url)
BamHI restriction sites were subsequently inserted between the MluI and DraIII sites using oligos (5′-CGGCGGTCAAGCTGATCCGCAGATTCATC-3′ and 5′-GGATCCGGATCCGGCGGTCCAG-3′) that destroy both of these sites. The H1 promoter taken from the pSuper plasmid (Oligoengine) was inserted into with Sall and BamHI sites of the modified pEGFP vector, creating pZOff.

Design of shRNAs
shRNAs were designed corresponding to the 21-mer target sequences using Ambion criteria specifying oligo duplexes with 5′-AA overhangs. The target sequence of Pclo28 is 5′-AAGTGCTGTCTCCTCTGTTGT-3′ (nucleotides 64-660 of Rattus norvegicus Piccolo from GenBank/EMBL/DDBJ accession no. NM_020098). The target sequence of Piccolo antisense, 5′-GAGGCGGTCAAGCTGATCCGCAGATTCATC-3′; and scrambled antisense, 5′-AGCITTTCTAAAGGATGTCATCTGCTTGTTGCTTAGCTCAATTTTGTAAGG-3′, 100 pmol of the respective strands were annealed and ligated using phage T4 polynucleotide kinase (New England Biolabs, Inc.) in a ligation reaction. The annealed cDNA duplexes were then phosphorylated using phage T4 polynucleotide kinase (New England Biolabs, Inc.) and ligated into the XbaI site of the EGFP-Synapsin1a plasmid. 100 pmol of the respective strands were annealed in a ligation reaction using a Thermocycler (Bio-Rad Laboratories). The annealed cDNA duplexes were then phosphorylated, ligation, and introduced into HEK293T cells using Lipofectamine 2000 (Invitrogen) with 10 μg of DNA per 10-cm plate. Homogenates from cells expressing Pclo28 or Bsn16 shRNAs were prepared 48 h after transfection. In brief, cells were harvested into homogenization buffer (ice-cold 1× PBS supplemented with protease inhibitors [complete protease inhibitor tablet, EDTA free; Roche]), and then placed directly into loading buffer. Protein levels were standardized empirically using a neomycin antibody and several loading concentration factors. After separation by SDS-PAGE, proteins were transferred to nitrocellulose membranes (GE Healthcare) and probed with primary and secondary antibodies in Western blotting solution (5% nonfat dry milk and 0.05% NP-40 in Tris-buffered saline). Protein bands were visualized by HRP chemiluminescence (PerkinElmer). For Western blots of hippocampal cultures, neurons from lentivirus-infected coverslips were harvested directly into loading buffer and the same procedure was followed. Here, protein levels were standardized using tubulin or GFP antibodies.

EM

The ultrastructural analysis of glutamatergic asymmetrical synapses was performed on dissociated cultures of hippocampal neurons transfected by electroporation at the time of plating with pZOff-VAMP2-4HR±Pclo28 shRNA. Samples were fixed with 2.5% glutaraldehyde in 0.15 M cacodylate buffer for 1 h and processed as described previously (Micheva and Smith, 2005) with a few modifications. In brief, after fixation, neurons were incubated with 1 mg/ml DAB (Sigma-Alrich) in 50 mM Tris, pH 7.5, for 10 min. 0.01% H2O2 (Sigma-Alrich) was then added to the DAB solution for 30 min at room temperature to stimulate HRP-mediated DAB precipitation. After extensive washing, neurons were prepared for EM by a microwave irradiation protocol (described in detail previously [Micheva et al., 2003]). After preparation, ultrathin sections (60 nm) were cut on an ultramicrotome (Ultracut UCT; leica) and placed on copper grids. Samples were poststained with 5% uranyl acetate in 5% aqueous methanol for 1 min and 10% aqueous lead citrate stain. Samples were imaged with a transmission electron microscope (JEM-1230; JEOL Ltd.) at 80 kV accelerated voltage using a charge-coupled device camera (791; Gatan). All sample processing and EM was performed in the Cell Sciences Imaging Facility at Stanford University.

In a blinded fashion, control VAMP2-4HR- and VAMP2-4HR-Pclo28 samples were imaged and synapses were quantitatively evaluated. Only

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synapses with a clearly discernable PSD and at least two unambiguously HRP-labeled SVs were included in the analysis. The AZ was unambiguously defined as the length of presynaptic membrane precisely opposed to the PSD. Docked SVs were defined as all SVs that had an edge within 25 nm of the AZ. All transmission EM quantitative analysis was performed using Image J.

Imaging experiments
All live imaging experiments were performed on a custom-built [by S. Smith, Stanford University, Stanford, CA; and N. Ziv, Technion Faculty of Medicine, Haifa, Israel] confocal microscopy [Axiovert 100TV; Carl Zeiss, Inc.] equipped with a 40× 1.3 NA Plan Neofluar objective [Carl Zeiss, Inc.] and 488 nm and 514 nm lasers [Sapphire 488-20CDRH and Compass 215M-20; Coherent], using OpenView software [written by N. Ziv]. Neuronal coverslips were mounted in a custom-built chamber designed for perfusion and electrical stimulation, heated to 37 °C by a forced-air blower, and perfused with Tyrode's saline solution (25 mM Hepes, 119 mM NaCl, 2.5 mM KCl, 30 mM glucose, 2 mM CaCl₂, 2 mM MgCl₂, 50 μM CNQX, and 10 μM APV, pH 7.4).

FM loading/destaining
Functional presynaptic boutons were labeled with FM4-64 or FM5-95 dye (Invitrogen) by incubation in high-K⁺ Tyrodes solution [90 mM KCl and 31.5 mM NaCl] containing ~1 μg/ml FM dye for 60 s followed by normal Tyrodes + FM dye for 30 s. Neurons were then washed for ~5 min before imaging. Destaining was performed by electrical stimulation at the frequency (10, 5, or 2 Hz) and time interval (180, 90, and 30 s) specified for each experiment or by high-K⁺ Tyrodes solution for 60 s.

Image analysis and quantification were performed with OpenView software and Excel [Microsoft]. GraphPad Prism [GraphPad Software] was used for statistical analysis. To measure the percentage of FM colocalization with EGFP-Synapsin clusters, the number of FM-containing clusters was divided by the total number of EGFP-Synapsin clusters and multiplied by 100. To calculate relative FM intensities at boutons containing or lacking Piccolo, FM intensity values for infected neurons were normalized against those from neighboring uninfected neurons in the same field of view, enabling cross-cover slip comparisons. For FM destaining experiments at 5 and 10 Hz, the total number of action potentials elicited and images acquired were kept constant. Thus, at 10-Hz stimulation, images were acquired every 5 s for a total of 90 s, whereas at 5 Hz they were acquired every 10 s for a total of 180 s. In all cases, intensity values for a given FM punctum at each time point were expressed as a percentage of its initial fluorescence intensity before destaining using the following equation: (current FM intensity at time point / initial FM intensity) × 100. For each condition [EGFPSyn and EGFGSynPclo28], curves from untreated coverslips of cultures infected with LV/EGFP-Synapsin or LV/EGFPSyn-Pclo28 were averaged and plotted as described in the previous section.

Calculation of RRP size and Pr
Boutons were loaded using high-K⁺ Tyrodes to label the TRP (TRP label). RRP release was induced with either 500 mM sucrose or low-frequency electrical stimulation (2 Hz at 30 s; RRP release). Total FM destaining was induced with high-K⁺ stimulation for 60 s [background]. To calculate RRP size, puncta intensity values were put into the following equation to express the RRP as a percentage of the TRP: ([TRP label – background] / [RRP release – background] / [TRP label – background]) × 100. Similar values were obtained for both methods, indicating that they can be used interchangeably to measure RRP.

To calculate Pr, the TRP was loaded with FM5-95 using high-K⁺ Tyrodes, and release of the RRP was induced using 2-Hz, 30-s electrical stimulation. During stimulation, images were acquired every 5 s. Intensity values were expressed as a percentage of initial fluorescence intensity, those exhibiting minimal destaining (values >95% for all time points) were eliminated from the analysis, and remaining curves were averaged and plotted as described in the previous section.

FRAP analysis
EGFP-Synapsin or SV2 puncta were bleached to ~20% of their initial fluorescence intensity by a high-intensity laser beam [488 nm wavelength] at high magnification. Images were taken at a rate of 1 per minute to monitor fluorescence recovery for Synapsin, and 1 per minute followed by 1 per 5 minutes for SV2. For each time point, intensity values were expressed as a percentage of starting fluorescence before bleaching. To control for non-specific photobleaching during image acquisition, the intensities of bleached puncta were normalized against those of unbleached puncta for each time point as described previously [Tsuriel et al., 2006], with the following equation: ([Intensity of bleached Synapsin punctum at time t]/[Mean intensity of all unbleached puncta in field of view at time t]) × 100. To control for variability in the extent of bleaching for each puncta, values were further normalized, making zero the default value for bleached puncta and enabling us to pool and average all recovery curves for a given condition [EGFGSyn and EGFGSynPclo28]. Pr values were calculated using a custom macro written in Excel (N. Ziv, Tsuriel et al., 2006).

Synaptic dispersion
Dispersion of EGFP-Synapsin was induced by electrical stimulation [10 Hz for 90 s] as described previously [Chi et al., 2001]. Puncta intensity values were expressed as a percentage of initial fluorescence intensity before stimulation. Curves for all EGFP-Synapsin puncta were combined for a given condition [EGFGSyn and EGFGSynPclo28], averaged, and plotted. For this analysis, puncta that did not exhibit dispersion [percentage of initial fluorescence values >90 for all time points] were excluded.

To compare extent of Synapsin dispersion to that of FM destaining, EGFP-Synapsin intensity was monitored during the 5-Hz destaining experiment [see FM destaining protocol]. To assess the role of CaMKII in Synapsin dispersion, coverslips were perfused with normal Tyrodes solution containing 10 μM KN62 (tocris Bioscience) and incubated in the drug for 20 min before eliciting dispersion. EGFP-Synapsin intensity was monitored either before stimulation and every 5 s during stimulation [for dispersion curves], or before stimulation and at the last time point (t = 90 s; bar graphs).

FM5-95 destaining was also monitored after incubation with 10 μM KN62. For this experiment, boutons were first loaded with FM5-95 using high-K⁺ stimulation as described. After waiting 15 min to allow for some recovery of EGFP-Synapsin after stimulation, cells were incubated for an additional 20 min in 10 μM KN62. SV exocytosis was then elicited by 5-Hz stimulation for 180 s, and both FM intensity and EGFP-Synapsin intensity were monitored over this time period (images taken every 10 s). Destaining curves were then plotted as described and compared with destaining curves from untreated coverslips of cultures infected with LV/EGFP-Synapsin or LV/EGFPSyn-Pclo28.

Online supplemental material
Fig. S1 shows initial testing of Piccolo shRNA (Pclo28) using pZOff plasmid-mediated knockdown in COS7 cells and neurons. Fig. S2 demonstrates that another shRNA against Piccolo (Pclo6) produces the same phenotypes as Pclo28. Fig. S3 shows additional EM micrographs showing that VAMP2-HRP unambiguously labels the synapses of transfected neurons and does not obscure synaptic junctions or other synaptic structures. Fig. S4 shows that EGFP-SV2, like EGFP-Synapsin, is a reliable presynaptic marker. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200711167/DC1.

We are grateful to Noam Ziv, Nicole Calakos, and Rob Malenka for insightful suggestions and Tim Ryan for the EGFP-Synapsin clones. We also thank Kristina Micheva, John Perrino, and Yemane Gedde for their assistance with EM and hippocampal cultures.

This work was supported by grants from the National Institutes of Health (NS39471 and NS353862 to C.C. Garner), National Research Service Awards to C.L. Waites and R. Terry-Lorenzo, and the Bundesministerium für Bildung und Forschung to E.D. Gundelfinger.

Submitted: 30 November 2007
Accepted: 2 May 2008

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