The dynamin-binding domains of Dap160/intersectin affect bulk membrane retrieval in synapses

Ása M. E. Winther1, Wei Jiao1,*+, Olga Vorontsova1, Kathryn A. Rees1, Tong-Wey Koh2,†, Elena Sopova1, Karen L. Schulze2, Hugo J. Bellen2,3 and Oleg Shupliakov1,§

1Department of Neuroscience, DBRM, Karolinska Institutet, von Eulers väg 3, 171 77 Stockholm, Sweden
2Program in Developmental Biology, NRI, Baylor College of Medicine, Houston, TX 77030, USA
3Department of Molecular and Human Genetics and HHMI, NRI, Baylor College of Medicine, Houston, TX 77030, USA
*Present address: Center for Motor Neuron Biology and Disease, Columbia University Medical Center, New York, NY 10032, USA
†Present address: Dept of Molecular, Cellular and Developmental Biology, Yale University, 266 Whitney Avenue, New Haven, CT 06511, USA
§Author for correspondence (oleg.shupliakov@ki.se)

Accepted 6 December 2012
Journal of Cell Science 126, 1021–1031
© 2013, Published by The Company of Biologists Ltd
doi: 10.1242/jcs.118968

Summary
Dynamin-associated protein 160 kDa (Dap160)/intersectin interacts with several synaptic proteins and affects endocytosis and synapse development. The functional role of the different protein interaction domains is not well understood. Here we show that Drosophila Dap160 lacking the dynamin-binding SH3 domains does not affect the development of the neuromuscular junction but plays a key role in synaptic vesicle recycling. dap160 mutants lacking dynamin-interacting domains no longer accumulate dynamin properly at the periactive zone, and it becomes dispersed in the bouton during stimulation. This is accompanied by a reduction in uptake of the dye FM1-43 and an accumulation of large vesicles and membrane invaginations. However, we do not observe an increase in the number of clathrin-coated intermediates. We also note a depression in evoked excitatory junction potentials (EJPs) during high-rate stimulation, accompanied by aberrantly large miniature EJPs. The data reveal the important role of Dap160 in the targeting of dynamin to the periactive zone, where it is required to suppress bulk synaptic vesicle membrane retrieval during high-frequency activity.

Key words: Scaffolding molecules, Protein migration, SH3 domain, Neuromuscular junction, Drosophila

Introduction
Clathrin-mediated endocytosis and bulk endocytosis are the two major pathways for synaptic vesicle (SV) recycling operating at the synaptic periactive zone (PAZ) (Saheki and De Camilli, 2012). The action of endocytic effector proteins executing these membrane trafficking events is coordinated by large scaffolding molecules, such as dynamin-associated protein 160 kDa, Dap160, in Drosophila (the mammalian ortholog is intersectin), and epidermal growth factor receptor pathway substrate clone 15, Eps15. Both proteins act as a molecular scaffold and their loss leads to very similar defects in endocytosis in vivo (Koh et al., 2007). Experiments in non-neuronal cells have shown that they are localized at endocytic sites via binding to F-bar proteins, FCHO1 and 2 (Henne et al., 2010) and act as a platform to recruit endocytic effectors implicated in regulation of the actin cytoskeletal network at the presynaptic membrane. Although significant progress has been made in the identification of the binding partners of the scaffolding proteins, their precise function in synapses is poorly understood (Dittman and Ryan, 2009; Pechstein et al., 2010).

One of the key endocytic effector proteins implicated in interactions with the scaffolding protein complex is the GTPase dynamin, encoded by the gene shibire (shi) in flies (Evergren et al., 2007; Koh et al., 2004; Marie et al., 2004; Roos and Kelly, 1998). Dynamin is part of the protein machinery that mediates fission of newly formed vesicles from the plasma membrane. In temperature sensitive shi2°/° mutants kept at the restrictive temperature, most SVs fuse with the presynaptic membrane, but endocytosis is blocked and endocytic intermediates with dynamin collars and large vacuoles accumulate. Genetic deletion of all three mammalian dynamin genes also results in a block of synaptic vesicle recycling and accumulation of numerous constricted coated pits (Ferguson and De Camilli, 2012).

Interactions with dynamin involve several Src homology (SH3) domain modules of Dap160/intersectin (Roos and Kelly, 1998). In Drosophila nerve terminals this binding has been proposed to be important for aspects of dynamin function (Broadie, 2004; Koh et al., 2004; Marie et al., 2004; Roos and Kelly, 1998). Surprisingly, the complete loss of dap160 causes phenotypes in neuromuscular junctions (NMJs) that are much less severe than the loss of dynamin (Koh et al., 2004; Marie et al., 2004). Low frequency nerve stimulation results in near normal EJPs in dap160 mutants. Only conditions of high-frequency activity, such as 10 minutes of 10 Hz stimulation, revealed impairments in synaptic transmission (Koh et al., 2004). Similarly, assays of synaptic vesicle endocytosis with FM dye loading revealed no short-term defects in dap160 mutants, although after a 10-minute labeling period, dye uptake was significantly reduced (Marie et al., 2004). However, the dap160 null mutants displayed twofold higher frequency of spontaneous activity and larger amplitude of spontaneous events. Up to 50% decrease in levels of several endocytic proteins, including dynamin, endophilin, and synaptojanin was also reported (Koh et al., 2004; Marie et al., 2004). These findings led to the suggestion that Dap160 may coordinate the function of endocytic proteins at the PAZ, but they did not explain how it fulfilled this function.
In the present study we investigate how Dap160 may coordinate dynamin functions at the PAZ. We used the dap160 null background to express mutant Dap160 proteins lacking dynamin-interacting modules in neurons in order to study its function. Our experiments show that Dap160 relocates from the vesicle pool to the periactive zone during synaptic activity and that it concentrates dynamin at the PAZ. The interaction between the two proteins plays an essential role in controlling bulk SV membrane trafficking at the NMJ, but is not critical for clathrin-mediated endocytosis.

**Results**

**Dap160 accumulates in the distal pool of synaptic vesicles at rest and relocates to the PAZ during synaptic activity**

We first investigated the localization of Dap160 and its binding partner, dynamin, at rest and during synaptic activity following exposure to 60 mM K⁺ (high K⁺) for 10 minutes using confocal microscopy (Fig. 1A). Both proteins strongly colocalized under both conditions (Fig. 1B). A redistribution of the proteins within nerve terminals under stimulation was evident in confocal images (Fig. 1A). To examine their distribution within the active zone, we double-stained NMJs with antibodies against Dap160 and the presynaptic T-bar component, Bruchpilot (Brp) (Fig. 1C,D). The Brp labeling accumulated in spots, which indicated the position of the T-bar (Wagh et al., 2006). The size of individual punctae remained unchanged both at rest and upon high K⁺ stimulation (supplementary material Fig. S1A), while the level of colocalization between Dap160-ir and Brp-ir significantly decreased in stimulated synapses (Fig. 1D). The Dap160-ir accumulated in clear circles around Brp spots (Fig. 1C). Such ‘circles’ have been observed previously and are thought to outline synaptic periactive zones (Marie et al., 2004; Roos and

![Fig. 1. Localization of Dap160, dynamin and Bruchpilot in control (w1118) third instar NMJs.](image)
Dap160 coordinates bulk endocytosis in synapses

Dap160 and dynamin localize to an undefined synaptic compartment to the PAZ upon stimulation.

To define this compartment we studied the subcellular localization of Dap160 at rest using stimulated emission depletion (STED) microscopy (supplementary material Fig. S1B–D). Serial STED images through NMJs show that Dap160 surrounds unlabeled areas of the bouton located at or close to the plasma membrane (supplementary material Fig. S1C,D). We then labeled NMJs for synaptotagmin, a SV-associated protein (Estes et al., 1996; Littleton et al., 1993), to determine if Dap160 colocalizes with a SV marker. Synaptotagmin and Dap160 are largely overlapping but do not completely colocalize, and punctae labeled only with synaptotagmin are clearly seen (Fig. 1E). This further suggests that Dap160 at rest localizes in the intravesicular matrix within the SV pool (Pechstein et al., 2010).

We next used electron microscopy (EM) in conjunction with the immunogold technique to further clarify the subcellular localization of Dap160 and dynamin. The gold particles were silver enhanced to improve visualization of the labeling at the EM level. Dap160 and dynamin were found in the pool of SVs more than 100 nm away from the borders of the T-bar pedestal (Fig. 1F–K). The SVs immediately surrounding the presynaptic T-bar were not labeled (Fig. 1G,H,J,K; supplementary material Fig. S2A). To confirm that these vesicles and structures surrounding the T-bar are accessible to labeling with our immunogold approach we stained the NMJs with antibodies surrounding the T-bar are accessible to labeling with our immunogold approach we stained the NMJs with antibodies

We then reconstructed immunogold-particle localization at release sites for Dap160 in 3D from serial ultrathin sections of NMJs at rest and during synaptic activity in control and shits1 mutant (Fig. 2E–G; supplementary material Fig. S2G–I). In resting control synapses only 10% of gold particles were in the PAZ, while 70% were in the PAZ in stimulated controls, and 90% in stimulated shits1 mutants. Taken together, these experiments show that Dap160 and its binding partner dynamin relocate from the distal vesicle pool region to the PAZ during synaptic activity.

Deletion of Dap160 results in the mislocalization of dynamin but not Eps15 from the PAZ

We then assessed dynamin distribution in dap160 null mutants, dap160^{+/+}/Df(2L)bur-K1 (Koh et al., 2004), at rest and during synaptic activity. Immunofluorescence and immunogold EM experiments show that dynamin distribution is not dramatically changed in mutants when compared to control at rest (Fig. 3A,G). Dynamin is associated with the vesicle pool at rest and it is largely colocalized with Eps15 and the endocytic adaptor complex, AP2 (Fig. 3A–C,G,I; supplementary material Fig. S3A,C). At the EM-level, in dap160 null mutants at rest, a decrease in dynamin gold particle density is detected at the PAZ compared to control (Fig. 3K–M; P<0.05), whereas no decrease is observed for Eps15 labeling versus control (supplementary material Fig. S3D).

In stimulated dap160 null mutant NMJs, dynamin is dispersed throughout the terminal and is no longer accumulated in spots of fluorescence at the PAZ, unlike in stimulated control NMJs (Fig. 3D,H). However, Eps15 is still localized to endocytic punctae and colocalizes with AP2 as in control and shits1 NMJs (Fig. 3E,F,J; supplementary material Fig. S3B,C). Immunogold EM experiments confirmed that dynamin is not concentrated at the PAZ and is distributed along the presynaptic membrane in dap160 null mutant NMJ boutons (Fig. 3N,O). Quantification of gold particles showed a relative reduction of dynamin immunolabeling in the PAZ compared to stimulated control (Fig. 3Q). When compared to stimulated shits1, the percentage of endocytic pits at the PAZ associated with dynamin-immunoreactive gold particles was reduced threefold (from 72.7±24.9%; n=19 to 23.7±26.3%; n=15; mean±SD, P<0.0001, Student’s t-test). The relative distribution of gold particles for Eps15 was not significantly different comparing stimulated control, dap160 and shits1 animals (supplementary material Fig. S3E). Hence, deletion of dap160 leads to a specific mistargeting of dynamin from the PAZ during stimulation.

Dynamins are mislocalized from the PAZ in a dap160 mutant lacking the dynamin-interacting SH3 domains

Genetic deletion of dap160 results in a decrease of about 50% of the levels of several endocytic proteins, including dynamin, as well as to the appearance of satellite boutons, indicating a developmental defect (Koh et al., 2004; Marie et al., 2004). To investigate the functional role of dynamin targeting to the periactive zone by Dap160, we generated dap160 mutants lacking the dynamin interacting modules. To accomplish this, we first confirmed the earlier observations of Roos and Kelly (Roos and Kelly, 1998) that only the SH3A and SH3B domains bind dynamin (Fig. 4A). GST pull-down experiments with various SH3 domain combinations of Dap160 showed that SH3B is the key dynamin binding module (Fig. 4A). SH3AB and

Kelly, 1998; Roos and Kelly, 1999). This suggests that both Dap160 and dynamin relocate from an undefined synaptic compartment to the PAZ upon stimulation.
SH3A-D displayed stronger binding than SH3B alone, suggesting cooperativity among the SH3 domain modules. We then created transgenics in the *dap160*<sup>ΔD/Δf(2L)bur-K1</sup> null background and expressed them using the GAL4/UAS system and a neural-specific GAL4 driver (elav-GAL4C155). Two *dap160* transgenes were generated, one lacking the SH3B domain (*D<sub>B</sub>) and the other lacking both SH3A and SH3B domains (*D<sub>AB</sub>). Animals that expressed the full-length *dap160* cDNA under UAS control served as a positive control (Fig. 4B). The full-length and mutated *dap160* transgenes rescued the lethality of *dap160*<sup>ΔD/Δf(2L)bur-K1</sup> mutants to viable adults, which were fertile and able to fly. From here onwards, *ΔB* and *ΔAB* refers to animals that express the respective mutated *dap160* transgenes in the *dap160* null mutant background. We then confirmed that GST-bound dynamin proline-rich domain (PRD) does not immunoprecipitate Dap160 lacking SH3A and SH3B domains from the head extracts of our mutant transgenics, whereas it successfully precipitated Dap160 in wild-type and full-length rescue controls (supplementary material Fig. S4A). A weak Dap160 band was still detectable in case of *ΔB*. A locomotion assay performed at 25˚C did not reveal significant motor defects in *ΔAB* third instar larvae. However, a large decrease in locomotor activity was observed in *dap160* null mutants (Fig. 4G). Under stress conditions at elevated temperature (34 ˚C), at which the temperature-sensitive null mutant is paralyzed, a locomotion defect in *ΔAB* significantly different from controls was observed, suggesting a synaptic defect (Fig. 4H).

Earlier studies have revealed severe developmental defects in the synaptic architecture of *dap160* null mutants (Koh et al., 2004). It is possible that delivery of dynamin to nerve terminals by Dap160 is important for synaptic development. However, the *ΔAB* mutant, which lacks dynamin-interacting modules, did not display any defects in synaptic architecture or changes in number of satellite boutons (Fig. 4C–F). In addition, immunofluorescence did not reveal significant differences in dynamin expression in NMJs in the *ΔAB* mutant compared to control (supplementary material Fig. S4B–E). Thus, interactions of Dap160 involving...
Fig. 3. Mislocalization of dynamin from the periactive zone in the dap160 mutant (A1/DF).
(A–C) Confocal microscopy images showing the localization of (A) dynamin (Dyn), (B) Eps15 and (C) AP2 at rest in control NMJs. (D–F) Following stimulation there is a redistribution of (D) dynamin-ir, (E) Eps15-ir and (F) AP2-ir in control boutons. (G, H) Confocal images of dap160 mutant NMJs showing that Dyn (red) and Eps15 (green) are largely colocalized at rest and have similar distribution (G). During stimulation, dynamin is no longer accumulated in spots, unlike Eps15 (H). (I, J) Confocal images showing AP2-ir (red) and Eps15-ir (green) in dap160 mutant nerve terminals at rest (I) and after stimulation (J). Note colocalization of both proteins in spots at the plasma membrane (arrow). (K, L) Low-magnification electron micrographs illustrating the overall view of dynamin immunolabeling in dap160 mutant NMJs at rest. (M) High-magnification image showing dynamin-ir within the SV pool in dap160 mutant at rest. (N) Mistargeting of dynamin in the mutant NMJ upon stimulation. (O) High-magnification image of a labeled constricted pit (arrow) accumulated at the periactive zone following high K+ stimulation.

Fig. 4. Properties of the ΔAB mutant. (A) GST-fusion proteins containing Dap160-SH3A and/or SH3B precipitate dynamin from w1118 head extracts, whereas GST-SH3C, GST-SH3D and GST alone do not. Note that SH3AB and SH3A-D precipitate larger amounts. Precipitated proteins were subjected to western blotting with anti-dynamin and anti-actin. 1% of the extract used for the pull-down assay was loaded for lane 1 (Input). L1, linker region N-terminally of SH3A; L2 linker region between SH3A and B. (B) dap160 mutant flies expressing different UAS-dap160-rescue constructs (ΔAB, lacking SH3A and B; ΔB, lacking SH3B and full-length rescue) pan-neuronally produce Dap160, as revealed by western blot of head extracts stained with antibodies against Dap160 and actin. (C, D) Confocal images of NMJ4 stained with anti-Dlg (green) to reveal bouton outlines, anti-Brp (red) to reveal presynaptic active zones and anti-Dap160 (blue, insets). Normal NMJ morphology is seen in control larva (C) and in NMJs from ΔAB mutant (D). (E) Satellite boutons are apparent in the dap160 null mutant (Δ1/DF), arrow indicates example of one satellite bouton. (F) Quantification of the number of satellite boutons in the different dap160 rescue lines compared with control and dap160 null mutant. There were no significant differences between the numbers of satellite boutons in control, ΔAB and full-length rescue larvae. (G) Locomotion behavior in dap160 null mutant is significantly different from that of control, ΔAB and full-length rescue at 25°C. (H) At 34°C, a significant reduction (~24%) in grid squares entered was detected in ΔAB compared to control and full-length rescue larvae. Temperature-sensitive dap160 null mutant does not move at 34°C (not shown). Bar graphs indicate mean ± s.e.m. **P<0.01, ***P<0.001 using ANOVA, Tukey’s post test. Scale bars: 2 μm (C–E).
other portions than the SH3A-B module can successfully control the dynamin transport to the nerve terminal during development.

In resting synapses of both \( AB \) and \( AAB \) larvae, there were no significant differences in localization of dynamin. As in control, a strong colocalization with Eps15 was observed (Fig. 5A,C,E). However, upon stimulation with high potassium, dynamin is mislocalized. Confocal images of \( AAB \) larvae clearly show that dynamin immunoreactivity is no longer concentrated in spots at the PAZ, in contrast to control NMJs (Fig. 5B,D,E). Mistargeting of dynamin in stimulated \( AB \) mutant synapses was also evident, but less pronounced (Fig. 5E). To test if targeting of Dap160 to the PAZ during stimulation is perturbed in \( AAB \) mutants, we double-stained stimulated NMJs with antibodies raised against AP2 and the EH domain region of Dap160. Strong colocalization of Dap160\(^{AAB}\) and AP2 in spots of fluorescence was observed indicating that Dap160\(^{AAB}\) relocate along with AP2, as observed in control (Fig. 5F,G). Taken together, these experiments indicate that dynamin-interacting domains of Dap160 are predominantly responsible for targeting and accumulating dynamin at sites of synaptic vesicle recycling in \textit{Drosophila} NMJs upon intense stimulation.

**Synaptic transmission and styryl dye uptake in NMJs in Dap160 mutants lacking dynamin-interacting domains is impaired**

Earlier experiments revealed defects in synaptic transmission in \textit{dap160} null mutants and showed that they can be restored in animals expressing the full-length rescue construct (Koh et al., 2004; Marie et al., 2004). The defects are temperature dependent. The major physiological defects at room temperature (22 °C) included an increase in average amplitude of miniature junction potentials (mEJPs) and a decrease in amplitude of EJPs after prolonged high-frequency stimulation at 10 Hz. To determine whether neurotransmitter release is perturbed in transgenic \( AAB \) animals, we recorded in 1 mM Ca\(^{2+}\) and observed no significant difference in the amplitude and frequency of spontaneous mEJPs (Fig. 6A–C). The amplitude of evoked EJPs recorded at 0.3 Hz was also not significantly altered (Fig. 6D,E).

We then determined whether \textit{dap160}\(^{AAB}\) NMJs can sustain high-frequency neurotransmitter release. Muscles 6/7 were stimulated at 10 Hz for 10 minutes. The recorded data were binned at 60-second intervals and normalized to the first EJP amplitude bin. The \textit{dap160}\(^{AAB}\) synapses show a significant decrease in average EJP amplitude during stimulation when compared to the control (Fig. 6F). A similar decrease in average EJP amplitude during high-frequency stimulation has been observed in our previous studies in null mutants (Koh et al., 2007).

To study the recovery from synaptic depression, NMJs were stimulated for 4 minutes at 10 Hz and then single stimuli delivered at 0.3 Hz to measure the recovery. EJPs amplitudes in \textit{dap160}\(^{AAB}\) rescue mutants recover to the original level in less than 5 minutes and at a much slower rate than control (supplementary material Fig. S5A). Interestingly, large amplitude spontaneous events were observed during the recovery phase following high-frequency stimulation, some of which reached up to 11.2 mV (Fig. 6G,H). These large fusion events disappeared 60 seconds after the stimulation (Fig. 6I). Hence, the data indicate that dynamin binding domains are necessary to maintain normal synaptic transmission. Slow recovery and presence of large spontaneous events following high-frequency stimulation suggest a defect in endocytosis and/or the release machinery in the synapse during and briefly after high-frequency activity.

To further assay the observed defect in synaptic transmission we assessed loading of the styryl dye FM1-43 in control, \( AAB \), \textit{shitle} and full-length \textit{dap160} rescue third instar larvae using a standard protocol (Verstreken et al., 2008). We observed a statistically significant decrease in FM1-43 loading in the \( AAB \) animals as compared to control and full-length rescue.
Journal of Cell Science

(Fig. 6J–M,Q). In addition, an increase in the number of internal fluorescent structures was observed (Fig. 6N–P; supplementary material Fig. S5B–D,H,I), suggesting that some of the dye was trapped in large membrane compartments in the terminal. These FM1-43 labeled membrane compartments disappeared when the larvae were returned to 1 mM Ca\(^{2+}\) solution and again stimulated with high K\(^+\), supporting the proposition that they are related to the synaptic vesicle cycle (supplementary material Fig. S5E–G).

Subcellular organization of internal membrane structures in the Dap160 mutant lacking dynamin-binding domains

We next performed electron microscopy to investigate the origin of the internal structures formed during stimulation. Third instar larvae from control, \(\Delta AB\), \(AB\), and full-length rescue animals were fixed at rest and after 10 minutes exposure to high potassium. At rest, the morphology of \(\Delta AB\) NMJs is very similar to both control and rescue (Fig. 7A,B; supplementary material Fig. S6A) and the total number of vesicles per terminal is not significantly different (\(P>0.05\); Student’s t-test). However, the packing density of SVs in the distal pool was reduced, while the pool proximal to the T-bar remained unchanged (Fig. 7K). No significant increase in the number of docked vesicles, endocytic pits and endosome-like structures were observed (Fig. 7J,M).

Following stimulation, dramatic structural changes occur in \(\Delta AB\) NMJs. The density and number of SVs in proximal and distal pools is strongly reduced (Fig. 7C,D,L). Numerous large vesicles appear, as verified by serial ultrathin sections. In addition, large bulk membrane invaginations attached to the membrane by tubular connections were observed in the PAZ (Fig. 7D–J). Large vesicles connected by narrow membrane necks to the large membrane invaginations were often seen (Fig. 7E–H). Bulk membrane invaginations were also present in stimulated \(AB\) synapses (supplementary material Fig. S6B).

Surprisingly, no significant difference in the number of coated endocytic pits or collared pits was evident among \(\Delta AB\), \(AB\), full-length rescue and control. Coated pits found in synapses from...
ΔAB, ΔB, rescue and control all had a uniform size and did not vary in shape (supplementary material Fig. S6E). This is different from dap160 null mutant NMJs in which a moderate increase in number of coated pits upon stimulation has been reported (Koh et al., 2004). SVs attached to the active zone by electron dense projections were observed in synapses both in control and mutant NMJs (supplementary material Fig. S6C,D). Tilting the dense protrusions in the electron microscope showed that these electron dense projections are not formed by the plasma membrane, suggesting that they represent docked vesicle assemblies as described in tomographic studies of the cryosubstituted active zones in Drosophila (Jiao et al., 2010a) and are therefore not endocytic intermediates. The number of these docked vesicles was not significantly different in stimulated control and ΔAB NMJs (Fig. 7M). In summary, EM analysis revealed that deletion of dynamin binding domains in dap160 leads to severe defects in bulk membrane uptake at the PAZ during high-rate synaptic activity and that the ΔAB mutant does not inhibit the fission of clathrin-coated vesicles from the presynaptic membrane.

Discussion

The GTPase dynamin is the key constriction enzyme implicated in the fission reaction during SV membrane recycling at the PAZ. Immunocytochemical studies have localized dynamin to distinct regions at the plasma membrane and to the necks of clathrin-coated pits, thus confirming that dynamin can be membrane associated (Takei et al., 1995). Henceforth, labeling for dynamin has been used as a marker for the PAZ in synapses at rest (Marie et al., 2004; Wagh et al., 2006). Other studies have noted that dynamin localization changes during synaptic activity suggesting that dynamin is associated with an unknown cytoskeletal or ‘intravesicular matrix’ within the nerve terminal to prevent its
dispersed in the nerve terminal. Large membrane invaginations and endosome-like structures are accumulated.

Fig. 8. Scheme of the mechanism for dynamin targeting to the periactive zone during the synaptic vesicle cycle. In wild-type (wt) animals the endocytic proteins AP2, Eps15, Dap160 and dynamin are redistributed from the synaptic vesicle cluster to the periactive zone during intense synaptic activity. Fused synaptic vesicle membrane is recycled by clathrin-mediated endocytic proteins AP2, Eps15, Dap160 and dynamin are redistributed from the synaptic vesicle cluster to the periactive zone during intense synaptic activity.

The decrease in neurotransmission in ΔAB is likely due to the dramatic reduction in the number of synaptic vesicles in nerve terminals upon high-frequency stimulation. Accumulation of membrane invaginations and endosomes suggest that SVs are not properly reformed from the bulk endocytic structures accumulated in NMJs. How does this occur? One possibility is that the block of SV recycling occurs at a step of bulk endocytosis pathway that leads to the formation of SVs. Recent studies suggest that this step may involve clathrin, but contrary to the classical clathrin-mediated pathway it utilizes different adaptor complexes, such as AP1 and AP3 (Cheung and Cousin, 2012). The other non-exclusive possibility is that the classical clathrin-mediated endocytic pathway becomes inhibited at very early step prior to the clathrin recruitment at the PAZ. Our studies show that AP2, Eps15 and Dap160 recruitment to the PAZ is not affected in the mutant. The localization of clathrin, AP1, and AP3 at rest and during stimulation remains to be investigated.

Clathrin-coated pits do not seem affected in ΔAB rescue mutants and we did not observe a significant increase when compared to control synapses. The latter supports that dynamin-mediated budding of clathrin-coated pits is not inhibited. This further implies that the targeting of dynamin by Dap160 to the PAZ is not essential for the fission step of the classical clathrin-mediated pathway in Drosophila synapses. However, the proper fission of bulk endocytic intermediates is perturbed. The number of uncoated membrane invaginations linked to the presynaptic membrane with thin ‘necks’ and the number of large interconnected endosomal structures were dramatically increased in the mutants lacking dynamin-binding SH3 domains. Interestingly, it has been noted earlier that appearance of bulk membrane structures in Drosophila NMJs occurs in mutants linked to dynamin function or controlling dynamin function upon inactivation of clathrin heavy chain or in clathrin mutants (Kasprowicz et al., 2008). A massive bulk membrane uptake has also been reported in mouse dynamin 1 knockouts (Hayashi et al., 2008). All this supports the hypothesis that dynamin together with clathrin coordinate the balance between clathrin-mediated and bulk endocytosis in synapses.

Large spontaneous EJPs were recorded during the recovery phase following high-rate activity, suggesting that bulk vesicles observed in nerve terminals upon stimulation fuse at the active zone. Disappearance of these large vesicles after cessation of high-frequency stimulation implies that the fused membrane becomes reformed into SVs most probably by clathrin-mediated endocytosis. Fusion of large vesicles will lead to dramatic distortions of quantal release in synapses. Giant fusion events observed in our experiments possibly represent ‘traffic accidents’, which may have occurred because the scaffolding components controlling bulk membrane trafficking at synapses were imbalanced in the ΔAB mutant. This further indicates that the scaffolding function of Dap160/intersectin and proper targeting of dynamin are particularly important for synaptic function during high-rate synaptic activity.
Materials and Methods
Fly strains and generation of dap160 transgene and expression constructs
w^{111} was used as Dap160 wild-type control with the same genetic background as mutants and transgenes and referred to in the text as ‘control’. shibire 

11 (sh11) was obtained from the Bloomington Drosophila Stock Center. dap160^{w111}/

Df(2L)bar-K1 is referred to as dap160 null is described in Koh et al. (Koh et al., 2004). For rescue experiments the panneural driver line C153-Gal4;Df(2L)bar-K1/CyO, twi-GFP were crossed to UAS-dap160^{111}SC; dap160^{w111}/CyO, twi-GFP, or w; dap160^{w111}/CyO, twi-GFP; UAS-dap160^{w111}28 or w, dap160^{w111}/CyO, twi-GFP; UAS-dap160^{w111} and F1 non-GFP larvae were selected for experiments.

To generate the UAS-rescue constructs with SH3A and SH3B (pUAST-

dap160/AB) or with SH3B (pUAST-dap160/AB) deleted domains, genomic 

sequence downstream SH3B (nt 3586–4221, FlyBase ID FBgn0023383) was PCR-

amplified with forward primers SH3AAB1 or SH3B1 and reverse primer SH3AB1 using the pUAST-dap160 cDNA plasmid as template. Resulting PCR 

fragments had 48 nt or 18 nt extensions located 579 nt or 156 nt upstream of 

amplified region for SH3A and SH3B, respectively. Restriction sites for sub-cloning 

are shown in bold letters (supplementary material Table S1). The SpeI–PsalA- 

or MreI–PsalAI-digested PCR fragments were inserted into SpeI–PsalAI or 

MreI–PsalAI sites of the pUAST-dap160. Resulting plasmids, pUAST-dap160/AB 

and pUAST-dap160/AB, have 579 nt or 156 nt deletions comprising SH3B or SH3A, 

respectively.

The SH3 expression constructs L1-A-L2, L2-B, AB, ABCD, C, D, and 

dynamin-PRD in pGEX-6P-2 vector (GE Healthcare) were obtained by PCR using 

as a template Dap160 cDNA clone IP14822 and shibire cDNA clone LD21682 shi, respectively (FlyBase ID Fblenm014466, Drosophila Genome 

Resource Center, Bloomington, IN) (supplementary material Table S2). The ACD 

construct was generated by self-ligation of blunt ended PCR product (Fusion 

polymerase, Finnzyme), amplified from ABCD plasmid. See all primer sequences 

in supplementary material Table S1. The ACD sequence is identical to one was 

defined as the plasma membrane area 500 nm adjacent to the active zone and 

a 100 nm cytosolic space into the lumen from the plasma membrane as shown 

in supplementary material Fig. S2A. Gold particle density ratio PAZ/outside was 

calculated as the ratio of gold particle density in the PAZ to the gold particle 

density in the area adjacent to the active zone.

GST pull-down
The GST or GST-fusion dap160-SH3-L1-A-L2-B/C/AB/ACD/ABCD 

GST-fusion dynamin PRD proteins were expressed in BL21(DE3) cells and purified 

according to standard protocols using glutathione sepharose (GE Healthcare). For 

GST pull-down 25 μg of corresponding protein were bound to glutathione agarose 

beads (GE Healthcare), added to 2 mg of Drosophila head extract in pull down (PD) 

buffer (20 mM HEPES, 100 mM KCl, 2 mM MgCl2, 1 mM PMSF, 0.5% Triton X- 

100) supplemented with Protein Inhibitor Cocktail (Sigma) in a total volume of 

0.7 ml. The extracts were incubated on a rotation wheel for 4 hours at 4˚C, then 

washed four times with PD buffer. Protein complexes were eluted with 2× SDS-

PAGE sample buffer (Invitrogen) and analyzed by resolving them on a 4–12% SDS-

PAGE gel (Invitrogen) and immuno-blotted using specific antibodies.

Antibodies
Polycrystal rabbit and guinea pig Dap160 antisera previously characterized in (Koh et al., 2007) were used in dilutions: 1:500 (rabbit) and 1:1000 (guinea pig). The Polyclonal rabbit and guinea pig Dap160 antisera previously characterized in (Koh et al., 2007) were used in dilutions: 1:500 (rabbit) and 1:1000 (guinea pig). Anti-Brp and AP2 antibodies were kind gifts from E. Buchner (University of Connecticut). Anti-synaptotagmin 1:1000 (Littleton et al., 1993). The Anti-Brp and AP2 antibodies were kind gifts from E. Buchner (University of Connecticut). Anti-synaptotagmin 1:1000 (Littleton et al., 1993). The

Immunohistochemistry, imaging and quantification of fluorescent image
Labeling of third instar larval fillets was performed as described previously (Verstreken et al., 2008). Secondary antibodies conjugated to Alexa 488, 555 or 647 (Invitrogen) were used at 1:500, secondary antibodies conjugated to Cy5 were used at 1:100 (Invitrogen). Otto dye R1 conjugated secondary antibody was used at dilution 1:100 (Leica). Samples were mounted in Vectashield mounting medium (Vector Laboratories).

STEM and confocal microscopy images for Fig. 1D and supplementary material Fig. S1B,C were collected on SP5 confocal microscope system equipped with a STEED (Leica Microsystems) using 63× oil immersion objective (1.4 NA) and Leica Application Suite Software (Leica Microsystems), which was calibrated to provide resolution up to 70 nm. Other confocal images were captured with 63×

1.4 NA oil immersion objectives using LSM 510 or a LSM 700 (Carl Zeiss), and the accompanying software. Brightness and contrast were adjusted using ImageJ (http://rsb.info.nih.gov/ij/) or Photoshop (Adobe). All measurements were performed in ImageJ or Volocity (PerkinElmer). Volocity was used for evaluating levels of colocalization. Pearson’s correlation coefficient was determined for middle optical sections of individual boutons that were selected as Region of Interest. Measurements based on confocal images were from 15 boutons from at least three different animals for each condition and genotype.

Bouton morphology
To reveal bouton outline, third instar larval fillets were labeled with anti-Dlg. Presynaptic T-bars of the active zone were detected by labeling with anti-Bap. Quantification of satellite bouton number was performed according to Marie et al. (Marie et al., 2004). Small boutons that branch from the major NMJ axis or from the terminal bouton were classified as satellite boutons and scored. Confocal images were obtained as described above from NMJ3, segments 2–A5, at an optical section thickness of 0.5 nm. For each genotype 6–9 NMJs from two (dap160 null) or three (all other genotypes) larvae were analyzed.

Preembedding immunocytochemistry and TEM
Fillets from third instar larvae were prepared in HL3 without Ca2+. For experiments, fillets were incubated in HL3 buffer containing either EDTA for 10 minutes, resting conditions, or stimulated by addition of 60 mM K+ for 10 minutes. The specimens were fixed in 4% paraformaldehyde solution, in 0.1 M phosphate buffer, pH 7.2 and labeled and embedded for EM as earlier described (Jiao et al., 2010b). Serial ultrathin sections were cut with a diamond knife (Diatome), stained with 1% uranyl acetate and lead citrate on grids, and examined with a Tecnai 12 electron microscope (FEI). Images were quantified using NIH ImageJ software, and statistical evaluation was performed using Excel (Microsoft). Relative distribution of gold particles (%) at PAZ or synaptic vesicle pool (SVC) was calculated as the ratio of gold particle density in the PAZ or SVC to the sum of gold particles in the plasma membrane area 500 nm adjacent to the active zone. A 100 nm cytosolic space into the lumen from the plasma membrane as shown in supplementary material Fig. S2A. Gold particle density ratio PAZ/outside was calculated as the ratio of gold particle density in the PAZ to the gold particle density in the area adjacent to the active zone.

3D reconstruction of TEM images
Serial ultrathin sections were photographed using bottom mounted 2k×2k TecnCam F224 CCD Camera (TVIPS). Membrane contours were traced using a digitizer and transferred into Maya 8.0 3D-reconstruction program and rendered as described earlier (Jiao et al., 2010b).

Electrophysiology
Third instar larvae were dissected in Ca2+-free HL3 solution as described previously (Koh et al., 2007). mEJPs were recorded from muscle 6 or 7 in segment A3 with thin-walled micropipettes filled with 3 M KCl in HL3 solution containing 1 mM CaCl2 (Hallermann et al., 2010). A suction electrode backedfill with HL3 solution was used to stimulate motor axons innervating the muscles. Low frequency stimulation was applied at 0.3 Hz, high frequency at 10 Hz. Data were recorded using an Axoclamp 2B amplifier, and ClampFit v10 software running on a PC computer equipped with an A/D interface (Axon Instruments) was used for data collection. Recordings were used in analysis only when the membrane potential was above 60 mV for mEJP measurement and for evoked EJPs when the membrane potential fell by less than 10 mV during the recording period. mEJPs were quantified for frequency and amplitude by taking the average value during 1 min for a minimum of five traces. Evoked EJPs were analyzed by calculating the average amplitude of 10 EJPs per minute bin of 10 Hz stimulation for five or more recordings. Recovery from stimulation was analyzed by measuring the average amplitude of six EJPs evoked at 0.3 Hz following 4 minutes, 10 Hz stimulation for five traces. All recordings were analyzed using Clampfit v10 software and data processed using Graph Pad Prism v5. Recordings were from 5–11 animals per genotype and condition.

FM1-43 dye uptake
FM1-43 dye uptake experiments were performed as described by Verstreken et al. (Verstreken et al., 2008). For each experiment one control and one test larva were dissected on small Sylgard beds in 1× HL3 containing 1 mM CaCl2 and 4 μM FM1-43 (Invitrogen) back to back in a 1.5 ml Eppendorf tube for 10 minutes in a 34˚C waterbath. Excess dye was then washed away repeatedly with Ca2+-free HL3 solution over 15 minutes. For unloading experiments larval fillets was instead washed with HL3 solution containing 1 mM Ca2+ and then again stimulated with high K+ except this time without FM1-43 dye, and then subjected to washing with Ca2+-free HL3 solution. Labeling was captured with a 40X/water immersion objective (NA 1.0) on a LSM 700 (Carl Zeiss). Data acquisition as well as data processing and quantification were performed as described (Verstreken et al., 2008). Three animals per genotype were tested in dye uptake experiments. To test dye unloading two separate experiments for each genotype were carried out.
25 mm² squares and scoring the number of grid squares entered. Larval locomotion was tested at 25°C and 34°C. At elevated temperature the larvae were preheated for 10 minutes on preheated yeast-supplemented apple juice agar plates in humidified chambers before locomotion was tested. For each genotype and condition 24–27 larvae were tested.

Statistics

Statistical analysis of two groups was evaluated using Student’s t-tests. To evaluate the differences between more than two groups one-way analysis of variance (ANOVA) was used, either with Tukey’s post-test, comparing every mean with every other mean, or Bonferroni multiple comparison test, comparing selected pairs of means. Statistical analysis was performed using GraphPad Prism v5.0 or v6.0 (GraphStat Software, San Diego, CA). Non-significant differences are not indicated in figures.

Acknowledgements

We thank E. Buchner, M. González-Gaitán and the University of Iowa Hybridoma Bank for antibodies.

Author contributions

A.M.E.W., O.V., K.A.R., T.W.K., H.J.B. and O.S. designed experiments; A.M.E.W., W.J., O.V., K.A.R., E.S., K.L.S. and O.S. performed experiments; A.M.E.W., W.J., O.V., K.A.R., E.S. and O.S. analyzed data; all authors discussed the data and commented on the manuscript; A.M.E.W. and O.S. wrote the manuscript.

Funding

This work was supported by the Swedish Research Council [grant numbers 13473 and 529-2009-6646/ESF-Euromembrane] and Linneé Center DBRM; the European Union Seventh Framework Programme (SynSys-project) [grant agreement number HEALTH-F2-2009-242167]; and the Wallenberg Foundation and Hjärfnonden (to O.S.). H.J.B. is an investigator of the Howard Hughes Medical Institute.

Supplementary material available online at http://jcs.biologists.org.lookup/suppl:doi:10.1242/jcs.118968/-/DC1

References

(A) The diameter of Bruchpilot-ir spots does not change upon stimulation of control 3rd instar NMJs. Single diffraction-limited Bruchpilot-ir spots were measured in middle optical sections through nerve terminals in three different experiments (n=106 in each case). Bars show mean±SEM. Student’s t-test, $P>0.05$. (B) Serial confocal and (C) stimulated emission depletion (STED) images of NMJs labeled with antibodies against Dap160 at rest. Images marked with asterisk are shown in D at higher magnification. (D) STED images of a NMJ at rest labeled with antibodies against Dap160. Note that the non-labeled area in the image is gradually disappearing inside the bouton (white arrows). Scale bars (B, C) 1 µm; (D) 2 µm.
(A) Cartoon illustrating the definition of quantification zones in NMJs in immunogold experiments. (B, C) Electron micrographs of control NMJs stained with anti-Brp antibodies using the immunogold technique utilized for Dap160 and dynamin localization. Gold particles are localized within the proximal pool of SVs and associated with the T-bar’s extensions. (D-F) Low and high magnification electron micrographs (D, E) from control NMJs labeled with antibodies against cysteine string protein (CSP). (F) Bar graph showing densities of gold particles in the distal and proximal pools of SVs (marked in E) for CSP. Note even distribution of gold particles in both pools of vesicles. Bars show mean ± SEM. Student’s t-test P>0.05. (G – I) Images of middle sections of NMJs stained with Dap160 antibodies from the series used for 3D reconstructions depicted in Figure 2E-G. T-bar; m, mitochondrion; sv, synaptic vesicle. Scale bars: (B, C) 50 nm; (D) 500 nm; (E) 100 nm; (G – I) 200 nm.
Supplemental Figure S3.

(A-B) Confocal images showing that AP2 (red) and Eps15 (green) are largely co-localized in NMJs of shi<sup>⁶¹</sup> at rest (A), and upon high K<sup>+</sup> stimulation. (B) Accumulation of Eps15-ir and AP2-ir in spots at the periphery of the plasma membrane is observed upon high K<sup>+</sup> stimulation (indicated by an arrow in B). (C) Quantification of the co-localization of AP2 and Eps15 at rest and following stimulation in control and shi<sup>⁶¹</sup> NMJs, as measured by Pearson’s correlation coefficient. Note high level of co-localization of both proteins upon stimulation. (D) No difference in the localization of Eps15-ir could be detected in dap160 mutant NMJs at rest as revealed when quantifying gold particles in electron micrographs. (E) Eps15 is accumulated in the periactive zone in control, shi<sup>⁶¹</sup> and dap160 mutant NMJs during stimulation as shown by quantification of Eps15-ir localization in electron micrographs after stimulation. Scale bar 2 µm; Bars show mean ± SEM. Student’s t-test (C, D) and ANOVA, Tukey’s post test (E); *P<0.05, ***P<0.001.
Genetic deletion of SH3A and SH3B domains of Dap160 abolishes interactions between dynamin and Dap160, but does not affect dynamin accumulation at NMJs during development. (A) A dynamin-PRD-GST-fusion protein precipitates Dap160 from Drosophila head extract in control flies and dap160 full-length rescue, but not from dap160 rescue mutants lacking SH3A and SH3B domains (ΔAB). Lane 5 shows GST alone. A very weak Dap160 band could be detected in ΔB mutant (not shown). Precipitated proteins were subjected to Western blotting with anti-Dap160 and anti-actin antibodies. 1% of the extract used for the pull-down assay was loaded for lane 2-3 (input), 0.7% of the extract was loaded in control* lane (lane 1; input). (B) Dynamin labeling intensity in ΔAB NMJs is not significantly different from control. Bars show mean ± SEM, Student’s t-test, P>0.05. (C - E) Confocal images of NMJs stained with antibodies against dynamin (Dyn, red) and Discs large (Dlg, blue; used to reveal NMJ outlines) showing that the level of dynamin-ir is the same as in control, while the level of dynamin is decreased in boutons from the dap160 null (Δ1/Df) mutant as previously shown (Koh et al., 2004). Scale bar: 2 μm.
(A) Recovery of evoked EJP amplitude induced by 0.3 Hz stimulation following 10 Hz stimulation for 10 min, normalized to the largest value in control and ΔAB NMJs. (B) Serial optical sections showing FM1-43 uptake in a ΔAB NMJ, arrows indicate the dye accumulation in endosome-like structures. (C) Cartoon illustrating the method for calculating relative distribution of FM1-43 uptake inside NMJs induced by high K⁺ stimulation as a ratio between the labeling intensity within a 0.7 µm x 0.7 µm square randomly placed at the edge of the bouton (“A”) and the labeling intensity over the whole bouton (“B”). (D) Bar graph showing relative distributions of FM1-43 (“A”/“B” ratio) in ΔAB rescue mutant vs control. (E) Image of boutons from ΔAB rescue mutant loaded with FM1-43 after 10 minutes exposure to high K⁺. (F) Image of boutons unloaded after exposure of the preparation to 1mM Ca²⁺ for 15 min followed by high K⁺ stimulation for 10 min. (G) Quantification of the loss of FM1-43 labeling intensity after unloading in control and ΔAB rescue mutant. (H, I) Images of a control and a ΔAB NMJ shown in Fig. 6N and O at higher magnification. Borders of the boutons are indicated by thick arrows. Student’s t-test *P<0.05; **P<0.01; ***P<0.001; Scale bar: 2 µm.
(A) Electron micrograph of a synapse from a dap160 full-length rescue mutant at rest. (B) Electron microscopic image of a ΔB terminal exposed to high K⁺ displaying large endosome-like profiles accumulated around the T-bar. (C, D) Electron micrographs showing examples of docked vesicles (arrowhead) in the active zone (thick arrow) connected to the presynaptic membrane by an electron-dense connector (shown at higher magnification and indicated with a star in D). (E) A clathrin-coated pit (arrowhead). T, T-bar; m, mitochondrion; sv, synaptic vesicle; el, endosome-like profile. Scale bars: (A, B) 250 nm; (C-E) 50 nm.
**Supplementary Table 1.**

**Primer sequences**

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>ΔSH3ABr</td>
<td>GTGACATGGGTCGGCTGTA</td>
</tr>
<tr>
<td>ΔSH3ABf</td>
<td>ACTAGTGATCGTGCCGCTGGGACGACACGGCTCTTCCAG TapGGAGGCTGATGTGGGCACACAGCGGA</td>
</tr>
<tr>
<td>ΔSH3Bf</td>
<td>GCCGCGGGCGATGTCCACAGGTGCCACAGCGCGA</td>
</tr>
<tr>
<td>DapSH3ABf</td>
<td>ATCGCGGATCCGAGACGGGTACGACTGTCAC</td>
</tr>
<tr>
<td>DapSH3ABr</td>
<td>AGCGTTGCGGCCGCTCACTTTTGGACATAGTTCGAGG</td>
</tr>
<tr>
<td>SH3AImr</td>
<td>AGCGTTGCGGGCGCTCACTCGACATCGGCCGCGG</td>
</tr>
<tr>
<td>linSH3Bf</td>
<td>ATCGCGGATCCCTAGAAAGTCGTAAGG</td>
</tr>
<tr>
<td>SH3Br</td>
<td>AGCGTTGCGGGCCGCTCACTTTTGGACATAGTTCGAAG</td>
</tr>
<tr>
<td>SH3CDr</td>
<td>AGCGTTGCGGCCGCTCACTTTTGGGTCGGACATAGTTCGAG GAGG</td>
</tr>
<tr>
<td>SH3Cf</td>
<td>ATCGCGGATCCATTGCTCAAGTATCGGCC</td>
</tr>
<tr>
<td>SH3Cr</td>
<td>AGCGTTGCGGGCCGCTCAAAACCTTCACGTAAG TGGCC</td>
</tr>
<tr>
<td>SH3Df</td>
<td>ATCGCGGATCCAAGGTTCAGTCTCTCTCATC</td>
</tr>
<tr>
<td>SH3Dr</td>
<td>AGCGTTGCGGGCCGCTCAACATAGTCTGCTAGGGAAG</td>
</tr>
<tr>
<td>SH3ACDr</td>
<td>Pho-CTCGACATCGCCGCGCCG</td>
</tr>
<tr>
<td>SH3ACDf</td>
<td>Pho-GCTGATGTGGCCACACCGG</td>
</tr>
<tr>
<td>GSTprdf</td>
<td>ATCGCGGATCGCTCAATGGCCACCGTGTGTC</td>
</tr>
<tr>
<td>GSTprdR</td>
<td>AGCGTTGCGGCCGCTCACTTTGAAATCGCGA ACTGAAAGG</td>
</tr>
</tbody>
</table>
### Supplementary Table 2.

**Description of constructs**

<table>
<thead>
<tr>
<th>Construct</th>
<th>Forward primer</th>
<th>Reverse primer</th>
<th>Template</th>
</tr>
</thead>
<tbody>
<tr>
<td>pUAST-Dap160ΔAB</td>
<td>ΔSH3ABf</td>
<td>ΔSH3ABr</td>
<td>pUAST-Dap160gen</td>
</tr>
<tr>
<td>pUAST-Dap160ΔB</td>
<td>ΔSH3Bf</td>
<td>ΔSH3ABr</td>
<td>pUAST-Dap160gen</td>
</tr>
<tr>
<td>GST-ABCD</td>
<td>SH3ABf</td>
<td>SH3CDr</td>
<td>Dap160 cDNA clone</td>
</tr>
<tr>
<td>GST-AB</td>
<td>SH3ABf</td>
<td>SH3ABr</td>
<td>GST-ABCD</td>
</tr>
<tr>
<td>GST-C</td>
<td>SH3Cf</td>
<td>SH3Cr</td>
<td>GST-ABCD</td>
</tr>
<tr>
<td>GST-D</td>
<td>SH3Df</td>
<td>SH3Dr</td>
<td>GST-ABCD</td>
</tr>
<tr>
<td>GST-L1-A-L2</td>
<td>SH3ABf</td>
<td>SH3Alin2r</td>
<td>GST-ABCD</td>
</tr>
<tr>
<td>GST-L2-B</td>
<td>linSH3Bf</td>
<td>SH3Br</td>
<td>GST-ABCD</td>
</tr>
<tr>
<td>GST-ACD</td>
<td>SH3ACDf</td>
<td>SH3ACDr</td>
<td>GST-ABCD</td>
</tr>
<tr>
<td>GST-PRD</td>
<td>GSTprdF</td>
<td>GSTprdR</td>
<td>Shibire cDNA clone</td>
</tr>
</tbody>
</table>