Molecular dissection of the photoreceptor ribbon synapse: physical interaction of Bassoon and RIBEYE is essential for the assembly of the ribbon complex

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The ribbon complex of retinal photoreceptor synapses represents a specialization of the cytomatrix at the active zone (CAZ) present at conventional synapses. In mice deficient for the CAZ protein Bassoon, ribbons are not anchored to the presynaptic membrane but float freely in the cytoplasm. Exploiting this phenotype, we dissected the molecular structure of the photoreceptor ribbon complex. Identifiable CAZ proteins segregate into two compartments at the ribbon: a ribbon-associated compartment including Piccolo, RIBEYE, CtBP1/BARS, RIM1, and the motor protein KIF3A, and an active zone compartment including RIM2, Munc13-1, a Ca2+ channel α1 subunit, and ERC2/CAST1. A direct interaction between the ribbon-specific protein RIBEYE and Bassoon seems to link the two compartments and is responsible for the physical integrity of the photoreceptor ribbon complex. Finally, we found the RIBEYE homologue CtBP1 at ribbon and conventional synapses, suggesting a novel role for the CtBP/BARS family in the molecular assembly and function of central nervous system synapses.

Introduction

Chemical synapses are highly specialized cell–cell contacts that mediate efficient communication between nerve cells. Ultrastructurally, distinct pre- and postsynaptic regions mark the sites of neurotransmitter release and reception. Presynaptically, the regulated release of neurotransmitter is restricted to the active zone, which is characterized by an electron-dense cytoskeletal meshwork. This cytomatrix at the active zone (CAZ) is exactly aligned with the postsynaptic reception apparatus, an electron-dense cytoskeletal matrix known as the postsynaptic density (Ziff, 1997; Dresbach et al., 2001).

The mature CAZ is defined by a set of multidomain proteins that harbor several protein–protein or protein–lipid interaction domains. The complete protein composition of the CAZ is not known to date, but it includes the proteins Munc13-1 (Brose et al., 1995), RIMs (Wang et al., 1997, 2000), ERC/CAST (Ohtsuka et al., 2002; Wang et al., 2002), Piccolo/Aczonin, and Bassoon (Cases-Langhoff et al., 1996; tom Dieck et al., 1998; Wang et al., 1999). Piccolo and Bassoon are among the first CAZ proteins to appear at nascent synapses in cultured neurons and they are components of an active zone precursor vesicle (Vardinon-Friedman et al., 2000; Zhai et al., 2001; Shapira et al., 2003). Furthermore, they are very large multidomain proteins that are intimately anchored to the cortical actin/spectrin cytoskeleton and are present at both excitatory and inhibitory synapses in the brain (Richter et al., 1999; Wang et al., 1999; Fenster et al., 2000, 2003). This makes Piccolo and Bassoon prime candidates for scaffolding proteins involved in the assembly of functional active zones.

A unique type of chemical synapse, structurally and functionally specialized for the tonic release of neurotransmitter in the dark, is the photoreceptor ribbon synapse. The presynaptic ribbon constitutes an electron-dense band of large surface area that extends from the site of transmitter release into the presynaptic cytoplasm and is covered by hundreds of synaptic vesicles (Rao-Miroztnik et al., 1995). The synaptic ribbon was thought to be a unique structure specialized to ribbon synapses in sensory organs. However, an emerging idea is that all chemical synapses are organized according to a common principle in which structural differences correlate with the kinetics of transmitter release (Zhai and Bellen, 2004). Within this concept, every synapse has dense projections on which vesicles are tethered, and the ribbon is a variation of this common theme. The ribbon is defined

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Abbreviation used in this paper: CAZ, cytomatrix at the active zone.
and organized by a scaffold of proteins that are just beginning to be identified. One of these proteins is RIBEYE. It has self-aggregating properties and is a major constituent of the ribbon (Schmitz et al., 2000). Another protein, the kinesin motor protein KIF3A, is enriched on ribbons, but its function there is still unknown (Muresan et al., 1999). Two other integral components of photoreceptor ribbon synapses are Bassoon and Piccolo (Brandstätter et al., 1999; Dick et al., 2001). In mouse retinæ deficient for functional Bassoon protein, photoreceptor ribbons are not anchored to the presynaptic active zones, but float freely in the cytoplasm and transmitter release is greatly perturbed (Dick et al., 2003). These results demonstrated a critical role of Bassoon in the formation and function of photoreceptor ribbon synapses.

To gain a more detailed picture of the photoreceptor ribbon protein architecture, we exploited the phenotype of the Bassoon mutant mouse. We report the segregation of CAZ proteins into two distinct molecular compartments of the ribbon complex: a ribbon and an active zone compartment. The physiological interaction between Bassoon and RIBEYE seems to be involved in linking the two compartments and the assembly of a functional ribbon complex. Finally, we report that the RIBEYE homologue CtBP1/BARS (COOH-terminal binding protein 1/brefeldin A adenosine diphosphate ribosylated substrate), a protein that has previously been identified as a transcriptional corepressor and Golgi-localized membrane-fissioning protein (Weigert et al., 1999; Turner and Crossley, 2001), is present both at photoreceptor ribbon synapses and at conventional synapses of retina and brain. This finding suggests a novel role for the CtBP family in the molecular assembly and function of central nervous system synapses.

Results

Protein composition of the presynaptic photoreceptor ribbon complex

In a retina deficient of a functional Bassoon protein, the photoreceptor ribbons are not anchored to the presynaptic membrane and float freely in the photoreceptor terminals (Dick et al., 2003). We exploited the phenotype of free-floating ribbons to screen for active zone molecules associated with the ribbon complex. As a marker for photoreceptor ribbon material we chose RIBEYE, the major protein component of photoreceptor and bipolar cell ribbons in the retina (Schmitz et al., 2000). RIBEYE is composed of a unique A-domain specific for ribbons, and a B-domain that is, with the exception of the first 20 amino acids, identical to the transcriptional corepressor CtBP2. Both RIBEYE and CtBP2 are derived from the same gene (Schmitz et al., 2000).

We generated an antiserum against the RIBEYE-specific A-domain. In retina homogenate this antiserum recognizes two protein bands of ~110 and 120 kDa that are also detected with an antibody against the B-domain (Fig. 1 A). In addition, the B-domain antibody recognizes the 50-kD CtBP2 band (Fig. 1 A). In preadsorbed controls, the immunodetection of RIBEYE in Western blots and in mouse retinal sections was abolished (Fig. 1, B and D).

In vertical sections of wild-type mouse retina, the A-domain antiserum produces labeling in the two synaptic layers of the retina, the outer plexiform layer (OPL) and the inner plexiform layer (IPL) (Fig. 1 C). The labeling represents RIBEYE at the glutamatergic ribbon synapses of the photoreceptors in the OPL and the bipolar cells in the IPL. Staining with the B-domain antibody is comparable to that obtained with the A-domain antiserum. In addition, immunoreactivity in the nuclear layers of the retina is observed, which derives from staining of the transcriptional corepressor CtBP2 (unpublished data). Postembedding immunogold EM shows an identical distribution of gold particles for A- and B-domain antibodies over the whole length of the photoreceptor synaptic ribbons (Fig. 1, E and F), a result corroborated by double-labeling immunocytochemistry and confocal laser-scanning microscopy (Fig. 1 G). Of note, the region of the arciform density at the ribbon base is free of immunogold particles in both cases (Fig. 1, E and F; arrowheads). These experiments indicate that both A- and B-domain antibodies recognize the same protein at photoreceptor ribbon synapses, i.e., RIBEYE (see also Schmitz et al., 2000). That RIBEYE represents ribbon material is further demonstrated by the fact that the free-floating photoreceptor ribbons in the Bassoon mutant retina are decorated with RIBEYE immunoreactivity (Fig. 1 H). Both A- and B-domain antibodies were used interchangeably in the study to label photoreceptor ribbons.

With immunocytochemistry and confocal laser-scanning microscopy, we screened in wild-type photoreceptor terminals a number of presynaptic proteins for their localization as compared with RIBEYE and thus for their potential to interact with the photoreceptor ribbon. The proteins synaptotagmin, VGLUT1 (vesicular glutamate transporter 1), SNAP-25 (synaptosomal-associated protein of 25 kD), syntaxin 3, and synaptophysin were present throughout the photoreceptor terminals; they did not specifically colocalize with RIBEYE, and thus were excluded from further analysis (unpublished data). Proteins that colocalized with RIBEYE in wild-type photoreceptor terminals were Piccolo, the kinesin motor protein KIF3A, RIM1, RIM2, Munc13-1, a Ca$^{2+}$ channel α1 subunit, ERC2/CST1 (Fig. 2), and Bassoon (see Fig. 6).

In a next step, we addressed the question of whether these proteins are integral parts of the protein network of the ribbon or are associated rather with the active zone membrane. Therefore, we screened these proteins for their colocalization with RIBEYE in Bassoon mutant photoreceptor terminals with their freely floating ribbons. The right panel of Fig. 2 summarizes the results of this mutant screen. Although Piccolo, KIF3A, and RIM1 colocalized with RIBEYE in the mutant photoreceptor terminals, the proteins RIM2, Munc13-1, the Ca$^{2+}$ channel α1 subunit, and ERC2/CST1 did not. Of note, RIM2 and the Ca$^{2+}$ channel α1 subunit colocalize at hot spots in the mutant photoreceptor terminals, but not RIM1 and RIM2 (unpublished data). It should be pointed out that the appearance of the photoreceptor ribbons labeled for RIBEYE changes from a horseshoe shape in the wild-type photoreceptor terminals (Fig. 2, left) to immunoreactive puncta of variable size in the mutant photoreceptor terminals (Fig. 2, right).
A likely reason could be that, although the size of mutant and wild-type ribbons appears similar in single ultrathin sections, the three-dimensional extension of the mutant ribbons is smaller. The large immunoreactive puncta represent aggregates of several free ribbons in mutant cone photoreceptor terminals, the small immunoreactive puncta are single free ribbons in mutant rod photoreceptor terminals (Fig. 1 H, 2). Furthermore, in the mutant photoreceptor terminals, staining became more diffuse for RIM2, Munc13-1, the Ca\textsuperscript{2+}/H1\textsubscript{1} subunit, and ERC2/CAST1.

To examine if the differential localization of CAZ proteins within the synaptic ribbon complex observed in Bassoon mutants reflects the wild-type situation, postembedding immunogold localization analyses were performed on wild-type retina. Our immunoblot analyses revealed that large amounts of RIBEYE, Piccolo, and ERC2/CAST1 were coimmunoprecipitated specifically with Bassoon (Fig. 4 A). Control IgG did not precipitate these proteins (Fig. 4 A). Only very low amounts of RIM1/2 and Munc13-1 coimmunoprecipitated with Bassoon as observed upon long exposition of the blots shown in Fig. 4 A (unpublished data). The cell adhesion molecule cadherin (Fig. 1 A)

Figure 1. Characterization of the photoreceptor ribbon protein RIBEYE in wild-type and Bassoon mutant mice. (A) Immunoblot of mouse retinal homogenate probed with antibodies against RIBEYE A- and B-domain. Two protein bands (double arrow) of \( \sim 110 \) kD and 120 kD, representing RIBEYE, are recognized by both antibodies. In addition, the anti-RIBEYE B-domain antibody labels the SO4D CIBP2 protein band (arrow). Protein bands above and below the RIBEYE doublet are unspecific and appeared only occasionally. (B) Preincubation of the anti-GST-RIBEYE A-domain antiserum with an excess of the MBP-RIB179-448 fusion protein (+) completely prevents the immunodetection of RIBEYE. −, no RIBEYE antigen included. Incubation of the same blot with the anti-RIBEYE B-domain antibody shows the presence of RIBEYE in both lanes. (C) In a vertical cryostat section through mouse retina, the anti-RIBEYE A-domain antiserum stains the photoreceptor ribbons in the outer plexiform layer (OPL) and the bipolar cell ribbons in the inner plexiform layer (IPL). (D) Preadsorption of the anti-RIBEYE antiserum with the antigen results in a complete loss of RIBEYE staining. (E and F) EM and postembedding immunogold labeling shows that photoreceptor ribbons are decorated with gold particles for RIBEYE A- (E) and B-domain (F). Note the absence of gold particles at the base of the ribbons (arrowheads), the region of the arciform density. (G) Confocal laser-scanning micrograph of a region of the OPL double labeled for RIBEYE A- (green) and B-domain (red) demonstrate the colocalization of the two immunoreactivities at the photoreceptor ribbons. (H) An aggregate of free-floating ribbons in a cone photoreceptor terminal of Bassoon mutant retina decorated with RIBEYE immunoreactivity (preembedding labeling).

ONL, outer nuclear layer; INL, inner nuclear layer; GCL, ganglion cell layer. Bars: 10 \( \mu \)m (C and D); 0.1 \( \mu \)m (E and F); 5 \( \mu \)m (G); and 0.4 \( \mu \)m (H).
Figure 2. Screening the photoreceptor ribbon synaptic complex for ribbon-associated proteins in wild-type (+/+) and Bassoon mutant (−/−) retinae. (Left) Confocal laser-scanning micrographs of photoreceptor ribbons in the +/+ retina double labeled for RIBEYE combined with the following CAZ proteins: Piccolo, KIF3A, RIM1, RIM2, Munc13-1, Ca\(^{2+}\) channel α1 subunit, and ERC2/CAST1. All CAZ proteins colocalize with RIBEYE as seen in the merge of the stainings. (Right) In the −/− retina, Piccolo, KIF3A, and RIM1 colocalize with RIBEYE, but not RIM2, Munc13-1, Ca\(^{2+}\) channel α1 subunit, and ERC2/CAST1. Bars, 5 μm.
4 A), the Ca\(^{2+}\) channel \(\alpha 1\) subunit (unpublished data), and the kinesin motor protein KIF3A (see Fig. 6 F) were not coimmunoprecipitated with Bassoon. There was also no immunoprecipitation of the polyphosphoinositide phosphatase synaptojanin 1 (see Fig. 6 F), which was recently shown to be the protein mutated in the zebrafish nrc mutant that displays a photoreceptor phenotype similar, although not completely identical, to the Bassoon mutant phenotype (Van Epps et al., 2004).

The identification of RIBEYE in the immunoprecipitated Bassoon complexes prompted us to investigate the putative Bassoon–RIBEYE interaction more closely because RIBEYE, in contrast to Piccolo and ERC2/CAST1, is specific for ribbon synapses (Schmitz et al., 2000). Our hypothesis of a Bassoon–RIBEYE interaction was strongly supported by yeast two-hybrid experiments that were performed in parallel in order to identify Bassoon binding partners that specifically interact with the central part of Bassoon that is lacking in Bassoon mutant mice. A screen conducted with the bait RB29 on a rat brain cDNA library revealed the RIBEYE homologue CtBP1 as potential binding partner for Bassoon (Fig. 4 B). As CtBP1 shows strong homologies to RIBEYE, both interactions were studied in more detail. Subclones of RB29 were generated and tested for their interaction with CtBP1 as well as with the A- and B-domain of RIBEYE. As documented in Fig. 4 B, a region situated between the coiled-coil (CC) regions 1 and 2 of Bassoon encoded by clones RB29, RB35, and RB46 (aa 1653–2087) are able to interact with both CtBP1 and the RIBEYE B-domain, but not with the RIBEYE A-domain. No other regions of Bassoon seem to interact with CtBP1 and RIBEYE (Fig. 4 B). These data demonstrate that the interaction of RIBEYE and CtBP1 with Bassoon is mediated by a region that is missing in the mutant protein, and therefore might contribute to the observed phenotype of freely floating photoreceptor ribbons in mutant mice.

**Reconstitution of RIBEYE B-domain/GFP-Bassoon interaction upon heterologous expression**

To confirm the results from the yeast two-hybrid analysis and the immunoprecipitation experiment of an interaction between
a central region of Bassoon and the RIBEYE B-domain, we examined the ability of Bassoon and RIBEYE to interact upon heterologous expression in COS cells.

The RIBEYE B-domain was targeted to the outer mitochondrial membrane by generating a fusion construct with an appropriate mitochondrial targeting sequence and a flag epitope tag (Mito-RIB-B). The construct is successfully localized to mitochondria as indicated by costaining with anti-Flag antibodies and MitoTracker (Fig. 5, A–C). A clear ring-like structure of the anti-Flag immunostaining (green) encircling mitochondria (red) suggests that the RIBEYE construct is successfully targeted to mitochondria and inserted in a correct orientation into the outer mitochondrial membrane (Fig. 5 C, inset).

Next, we analyzed the cellular distribution of GFP-Bassoon constructs in single transfection experiments. Most of the GFP-Bassoon constructs tested were diffusely distributed and not associated with mitochondria as exemplified for the GFP-RB46 construct in Fig. 5, D–F (compare with Fig. 4 B for extension of constructs). Some signal was also observed in the nucleus (Fig. 5 D).

When cells were cotransfected with GFP-RB46 and Mito-RIB-B (Fig. 5, G–J), GFP-RB46 localization clearly overlapped with that of Mito-RIB-B. The cytosol and the nuclei of double-transfected cells were devoid of any GFP-RB46 (Fig. 5, G–J), suggesting that the interaction with RIBEYE is of high affinity. Similar results were obtained with a more extended GFP-Bassoon fusion protein construct, GFP-RB35 (unpublished data).

Parts of Bassoon that showed no binding in the yeast two-hybrid analysis, such as RB40, maintained their diffuse distribution in both the cytosol and the nucleus when cotransfected with Mito-RIB-B. The GFP-RB45 construct, which includes a region of Bassoon that displayed very weak interaction in the yeast two-hybrid analyses, also did not adopt the mitochondrial distribution pattern when cotransfected with Mito-RIB-B (Fig. 5, K–M).

Reverse experiments with the GFP-RIBEYE B-domain, which shows a diffuse distribution when transfected alone, cotransfected into cells expressing the RB35 fused with mitochondrial targeting sequences, confirmed the formation of Bassoon–RIBEYE complexes. In contrast, the Bassoon domains of RB40 and RB45 fused to a mitochondrial targeting sequence failed to recruit the GFP-RIBEYE B-domain (unpublished data).

These data clearly confirm that the domains of Bassoon encoded by RB35 and RB46 interact with RIBEYE in living cells, whereas those encoded by RB40 and RB45 do not.

Interaction of Bassoon and RIBEYE at the base of the ribbon
With immunocytochemistry and light and electron microscopy we examined in detail the expression patterns of Bassoon and RIBEYE at the photoreceptor ribbon synapses. In light microscopy, the immunoactivity of the two proteins overlap in a horseshoe shape, with RIBEYE (Fig. 6 A, red) surrounding Bassoon (Fig. 6 B, green) as seen in the merge of the two stainings (Fig. 6 C). This finding is corroborated by the results from double-labeling postembedding immunogold experiments, in which we combined staining for Bassoon and RIBEYE using gold particles of different size (20 nm for Bassoon, 10 nm for RIBEYE).

![Figure 5. Reconstitution of Bassoon–RIBEYE protein complexes by cotransfection of the Bassoon and RIBEYE binding domains in COS cells.](image)

Fig. 6 D shows two different planes of section through photoreceptor ribbon synapses. One plane is parallel to the extent of the active zone (en face view), the other is orthogonal to it (cross-section view; Fig. 6 D, inset). Consistent with a previous quantitative examination of the localization of Bassoon (Dick et al., 2001), in both cases the larger gold particles for Bassoon are located closest to the active zone (Fig. 6 D, arrowheads). Together, these findings suggest a physical interaction between Bassoon and RIBEYE at the base of the photoreceptor ribbon.
The region of Bassoon that interacts with RIBEYE is missing in the 180-kD Bassoon mutant protein (Bsn/H9004Ex4/5), which is still made in the Bassoon mutant mouse (Altrock et al., 2003), but it is not targeted to the active zone and is diffusely distributed throughout the photoreceptor terminals (Dick et al., 2003). As the interaction between Bassoon and RIBEYE should be absent in mutant mice, we examined if RIBEYE expression and distribution is affected in mutant retinae. As shown in Fig. 6 E, the overall amount of RIBEYE in the Bassoon mutant retina is reduced to about half (52% ± 14%) as compared with wild-type retina (100% ± 19%). To confirm that mutant Bassoon protein does not bind to RIBEYE, we performed immunoprecipitation experiments with an antibody against the NH2-terminal region of wild-type Bassoon (BSN1.6), which recognizes both wild-type and mutant Bassoon (Dick et al., 2001). In contrast, ERC2/CAST1 is found in both precipitates. Kinesin KIF3A and synaptojanin are not detected in either precipitate. Control experiments were performed with preimmune serum (preIS). IP, immunoprecipitation. Bars: 2 μm (A–C), 0.2 μm (D).

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The RIBEYE B-domain homologue CtBP1 is present at ribbon and conventional synapses

As the interaction of Bassoon was originally observed with CtBP1, a homologue of the RIBEYE B-domain, we monitored the distribution of this protein in vertical sections of wild-type mouse retina using a monoclonal mouse anti-CtBP1 antibody. Interestingly, in addition to the expected nuclear staining, strong CtBP1 labeling was observed in the two synaptic layers of the retina (Fig. 7 A). Double-labeling experiments with an antiserum against the CAZ protein Piccolo, which is present at ribbon and conventional synapses in the retina (Fig. 7 B; Dick et al., 2001), show a complete colocalization of CtBP1 and Piccolo in the two plexiform layers of the retina (Fig. 7 C). Postembedding immunogold EM further proved the presence of CtBP1 at ribbon and conventional synapses in the retina (Fig. 7, D and E). Immunoblot analysis revealed no cross-reactivity of the CtBP1 antibody with RIBEYE (Fig. 7 F), demonstrating
that indeed the 50-kD CtBP1 protein is present synaptically. In the Bassoon mutant photoreceptor terminals with their free-floating ribbons, CtBP1 and Piccolo immunoreactivities completely colocalize at the synapses in the outer plexiform layer (OPL) and inner plexiform layer (IPL) of the retina. (D and E) Electron micrographs of a photoreceptor (D) and an amacrine cell synapse (E) immunogold labeled for CtBP1. At the ribbon synapse the gold particles for CtBP1 decorate the ribbon (arrowheads), at the amacrine cell synapse they are located some distance from the active zone (arrowheads) at the edge of the electron-dense CAZ material. (F) Western blots of retina homogenate probed with antibodies against CtBP1 and the RIBEYE A- and CtBP2/RIBEYE B-domain. The antibody against CtBP1 recognizes the 50-kD CtBP1 protein and does not cross-react with RIBEYE. (G–I) Micrographs of cultured hippocampal neurons double labeled for CtBP1 (G) and Piccolo (H). In hippocampal neurons, like in retinal neurons, a fraction of CtBP1 immunoreactivity is localized at synapses where it colocalizes with Piccolo as seen in the merge of the two stainings (I). (J and K) CtBP2 (J) is not present at hippocampal synapses labeled with Piccolo (K). (L) Immunoblots showing that CtBP1 can be coimmunoprecipitated from brain synaptosomal extracts with a monoclonal anti-Bassoon antibody. The boxes in G–K mark the regions that are shown at higher magnification. INL, inner nuclear layer; GCL, ganglion cell layer; Ext., brain synaptosomal extract; IP, immunoprecipitate; S, supernatant. Bars: 20 μm (A–C), 0.2 μm (D and E), and 10 μm (G–K).

Discussion

Photoreceptor cells, the primary neurons of the retina, are non-spiking neurons that use graded changes in membrane potential to transmit sensory information over a wide dynamic range of light intensities. The continuous adjustment of the synaptic output to changes in the incoming signals requires a specialized type of synapse, the ribbon synapse, that sustains high and long-lasting rates of neurotransmitter release (von Gersdorff, 2001; Parsons and Sterling, 2003).

The photoreceptor ribbon complex is subdivided into two molecular compartments

Photoreceptor ribbon synapses exocytose hundreds of vesicles per second, which requires a large pool of readily releasable
vesicles and a fast release mechanism. The current hypothesis of calcium-dependent exocytosis at retinal ribbon synapses suggests that (1) the vesicles tethered to the ribbon are primed; (2) the vesicles that are in close contact with the presynaptic plasma membrane at the ribbon base comprise the small, rapidly releasable pool of vesicles; and (3) the remaining vesicles tethered to the ribbon comprise the large, readily (slower) releasable pool (von Gersdorff et al., 1996; Heidelberger et al., 2002; Parsons and Sterling, 2003).

The ultrastructural appearance of a row of synaptic vesicles tethered to either side of the ribbon and the expression of a kinesin motor protein KIF3A at retinal ribbon synapses (Muresan et al., 1999) led to the conceptually attractive model that the ribbon, like a conveyor belt, moves primed vesicles to the docking/release site at the base of the ribbon. However, this model raises the question of how the whole readily releasable pool can be exocytosed within 1–2 ms, which is much too fast for moving the rows of vesicles to the release site by an active, kinesin-powered movement (Heidelberger et al., 1994). A challenging concept to explain these questions is compound exocytosis, a process by which vesicles tethered to the ribbon fuse with each other and with the vesicles docked at the plasma membrane (Parsons and Sterling, 2003). However, experimental evidence for compound fusion at ribbon or conventional chemical synapses is sparse.

Both scenarios, conveyor belt or compound fusion, postulate that membrane fusion and transmitter release occurs focally at the ribbon base. This is supported by our findings of the preferential clustering of a Ca$^{2+}$ channel $\alpha_1$ subunit and Munc13-1 in addition to RIM2 and ERC2/CAST1 at the presynaptic plasma membrane/arciform density compartment. As Munc13 seems to be an essential component of the priming machinery for synaptic vesicles at conventional synapses (Rosenmund et al., 2003), it may be assumed that the final priming steps take place near the active zone membrane.

In our search for CAZ proteins that potentially interact at the photoreceptor ribbon complex, we were able to dissect a second cytomatrix compartment that is distinct from the plasma membrane/arciform density compartment, the ribbon (Fig. 8). The two compartments are likely to reflect a segregation of CAZ functions at the photoreceptor ribbon synapse. This is most obvious by the differential distribution of RIM1 and RIM2. RIMs, CAZ proteins that function in a late step of exocytosis following the docking of synaptic vesicles, directly and indirectly interact with many proteins implicated in vesicle priming and neurotransmitter release at presynaptic active zones of synapses, including Munc13-1, ERC2/CAST1, and L-type calcium channels (Wang et al., 1997; Schoch et al., 2002; Calakos et al., 2004; Südhof, 2004). Ribbon-associated molecules including RIM1, Piccolo, RIBEYE/CtBP2, and CtBP1 may be involved in tethering synaptic vesicles to the ribbon and in preparing them for exocytosis. In particular, the functions proposed for RIBEYE/CtBP2 and CtBP1 could play a role in this latter process. Both proteins were first characterized as transcriptional corepressors (Turner and Crossley, 2001). In addition, they display homology to NAD$^{+}$-dependent dehydrogenases and bind NAD$^{+}$/NADH (Schmitz et al., 2000; Zhang et al., 2002). Most interestingly, in the context of preparing vesicles for exocytosis at the ribbon, they may function as lysophosphatidic acyl-CoA transferases (LPAAT) and thus modulate the curvature of lipid membranes (Kooijman et al., 2003). Indeed, CtBP1 has been implicated in membrane fission processes at the Golgi complex, where it is involved in a fission reaction generating aligned vesicles from continuous tubular structures (Weigert et al., 1999).

As shown in this paper, CtBP1 is not only a protein of the photoreceptor ribbon synapse, but also of conventional brain synapses. This suggests a general role for CtBP family members in active zone function. Our results add molecular evidence to the hypothesis that the ribbon has a correlate at the conventional synapse (Zhai and Bellen, 2004).

**Bassoon, a physical link between the two CAZ compartments at the photoreceptor ribbon synapse**

The floating ribbon phenotype in Bassoon-deficient retina suggests a functional involvement of Bassoon in anchoring the ribbon to the active zone (Dick et al., 2003). One possible explanation for this phenotype is that due to the lack of functional Bassoon, ribbon assembly is disturbed and/or delayed and therefore an appropriate anchoring is impossible. Data presented here offer a more precise explanation: Bassoon might be involved in linking the two newly identified compartments of the ribbon complex, i.e., the ribbon itself and the presynaptic plasma membrane/arciform density compartment. Immunogold data show that Bassoon is localized right at the border between the two CAZ compartments. At least two molecular interactions will be relevant for the attachment of the ribbon to the presynaptic plasma membrane—one interaction linking Bassoon to the synaptic ribbons, and one linking Bassoon to the plasma membrane.

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**Figure 8. Differential localization of CAZ proteins defines two compartments of the photoreceptor ribbon synaptic complex.** The ribbon-associated complex of CAZ proteins includes RIBEYE/CtBP2, CtBP1, KIF3A, Piccolo, and RIM1; the plasma membrane/arciform density-associated complex includes RIM2, Munc13-1, ERC2/CAST1, and a Ca$^{2+}$ channel $\alpha_1$ subunit. Bassoon localizes at the border between the two compartments.
Here, we show that Bassoon directly binds and functionally interacts with RIBEYE, the protein postulated to serve as a central building block of the synaptic ribbon (Schmitz et al., 2000). The cotransfection experiments in mammalian cells in vivo show a strong interaction between Bassoon and RIBEYE, and demonstrate that the interaction domains found by yeast two-hybrid analysis are sufficient for the interaction of the two proteins. The domain in Bassoon responsible for establishing the connection with RIBEYE is lost in the mutant Bassoon protein, and consequently the interaction between the two proteins. Currently, we do not know what the link is between Bassoon and the arctiform density/plasma membrane compartment. A likely candidate is ERC2/CST1, which has been shown to directly interact with Bassoon (Takao-Rikitsu et al., 2004) and which is present at the photoreceptor ribbon synapse (this paper). Because the interaction sites for ERC2/CST1 and RIBEYE map to different locations on Bassoon (ERC2/CST1, Takao-Rikitsu et al., 2004; RIBEYE, this paper), a simultaneous binding of the two proteins to Bassoon is possible. We propose that Bassoon contributes essentially to the link between the two compartments at the photoreceptor ribbon complex. The loss of the Bassoon–RIBEYE interaction explains the soon mutant retina.

Materials and methods

All animal experiments were performed in compliance with the guidelines for the welfare of experimental animals issued by the Federal Government of Germany, the National Institutes of Health, and the Max Planck Society. Mice were used at an age of 7–9 wk.

Antibodies

The following primary antibodies were used: mouse anti-CtBP1 [postembedding immunoelectron microscopy (post-EM) 1:500; immunohistochemistry (IHC) 1:2,500–1:5,000; Western blot (WB) 1:5,000] and anti-CtBP2 mAbs (post-EM 1:1,000; IHC 1:10,000; WB 1:5,000; BD Biosciences), rabbit anti-pan-Munc13 pAb (WB 1:250; BD Biosciences), mouse anti-RIM mAb (WB 1:1,000; BD Bio- sciences), rabbit anti-RIM1 (post-EM 1:50; IHC 1:500) and anti-RIM2 pAbs (IHC 1:500; Synaptic Systems GmbH), rabbit anti-Synaptotagmin I pAb (WB 1:3,000; Synaptic Systems GmbH), rabbit anti-ERC1/2/3 pAb (WB 1:5,000; Synaptic Systems GmbH), mouse anti-Bassoon mAb (IHC 1:2,500; WB 1:1,000; StressGen Biotechnologies), rabbit anti-Bassoon pAb (post-EM 1:50; IHC 1:100; Sigma- Aldrich), mouse anti-pan cadherin mAb (WB 1:100; a gift of Dr. P. Beesley, Royal Holloway University London, Egham, Surrey, UK), rabbit anti-parvalbu- min pAb (post-EM 1:100; Alomone Labs), mouse anti-kinesin II mAb (post-EM, IHC 1:50; WB 1:2,000; BABCo), guinea pig anti-Piccolo (IHC 1:8,000; WB 1:1,000–1:2,000) and rabbit anti-Piccolo pAbs (IHC 1:400; Dick et al., 2001), and mouse anti-Munc13-1 mAb (IHC 1:800; a gift of Dr. N. Brose, Max Planck Institute for Experimental Medicine, Göttingen, Germany).

Furthermore, rabbit antisera were produced against bacterially expressed GST fusion proteins of Bassoon (aa 73–181; BSN1.6), ERC2/CST1 (aa 73–185), and RIBEYE (aa 179–48).

Retinal tissue preparation and light and electron microscopic immunocytochemistry

A description of the preparation of the retinal tissue for light and electron microscopic immunocytochemistry is given in Dick et al. (2001). For light microscopy, retinas were fixed in 4% PFA in phosphate buffer (0.1 M, pH 7.4) for 15–30 min. For postembedding immuno-EM fixation was in 0.5% glutaraldehyde and 4% PFA in phosphate buffer for 2 h.

Retinal sections were incubated in primary antibodies overnight. For light microscopy, secondary antibodies were coupled to Alexa 594, 488, or 350 (1:500; Molecular Probes, Inc.); for postembedding immuno-EM, secondary antibodies were coupled to 10- or 20-nm gold particles (ICN Biomedicals).

For light microscopic analysis, labeled sections were examined with a confocal laser-scanning microscope (LSM5 Pascal; Carl Zeiss Microlimming, Inc.). Images were adjusted for contrast and brightness using Adobe Photoshop version 5.5, and figures were arranged using Corel Draw 9. For postembedding immunocytochemistry, tissues were examined and photographed with an electron microscope (EM10; Carl Zeiss Microlimming, Inc.) on negative film.

Western blot and immunoprecipitation

For Western blots of retina homogenate, retinae were homogenized, total protein was precipitated with TCA, dissolved in sample buffer, separated on 3.5–8% Tris acetate gels (25 μg/lane), and transferred to PVDF membranes by tank blotting (Attraok et al., 2003). For immunodetection, membranes were blocked and primary antibodies were applied overnight at 4°C. For characterization of the RIBEYE antisera, 5 μl RIBEYE serum was preincubated for 1 h with an excess of purified MBP-RIB (aa 179–448) fraction or with buffer alone and then diluted for overnight incubation of the Western blots. The HRP-coupled secondary antibodies were visualized by chemiluminescent detection (Amersham Biosciences).

Blots were reprobed several times with different antibodies. Evaluation of relative amounts of RIBEYE immunoreactivity was performed by determining gray values in nonsaturated exposures of X-ray films in a given area using Adobe Photoshop software. In four individual animal pairs, the mean value of the +/+ animals was defined as 100%.

For immunoprecipitation experiments, retinae were homogenized in 1 ml extraction buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1% Triton X-100, and 0.5% sodium deoxycholate) per animal. Insoluble material was pelleted. For antibody immobilization, protein G–Agarose beads (25 μl bed volume; Roche) were incubated overnight at 4°C with 10 μg monoclonal Bassoon antibody and as control 10 μg mouse IgG (Sigma-Aldrich), or with 5 μl rabbit anti-BSN1.6 antisera and 5 μl of a preimmune serum as control followed by washes. The extract corresponding to one retina was applied overnight to the antibody-coupled beads. Bound proteins were recovered after extensive washes in homogenization buffer by boiling in sample buffer. Extracted material (including the unbound proteins in the supernatant) were concentrated by TCA precipitation. SDS-PAGE and immunodetection were performed as described above.

For immunoprecipitation experiments from brain material, synapticosomal fractions were prepared as described previously (Phillips et al., 2001) and were extracted with detergent buffer (0.5% sodium deoxycholate, 0.1% Triton X-100, 70 mM NaCl, 25 mM Tris-HCl, pH 8.0, and total protein 0.125 μg/μl). Mouse anti-Bassoon mAb and unspecific mouse IgG (Santa Cruz Biotechnology, Inc.) were chemically coupled to Gamma-Blut Plus Sepharose (Amersham Biosciences) to abolish background derived from antibody proteins in the immunoprecipitation fraction. In detail, 20 μg IgG was incubated for 1 h with 100 μl of beads in PBS/T buffer (8.1 mM Na2HPO4, 1.5 mM KH2PO4, 2.7 mM KCl, 137 mM NaCl, and 0.1% Tween 20, pH 7.4). Beads were washed with PBS/T and with TEA buffer (0.2 M triethanolamine, pH 8.2) before incubation for 45 min at RT in coupling buffer (20 μM dimethyl pimelimidate dihydrochlorid in TEA). Beads were washed with TEA and incubated for 10 min at RT in 20 mM ethanolamine (pH 8.2). After washing with PBS/T, beads were stored in 5% BSA in PBS/T with 0.025% NaN3 at 4°C.

For immunoprecipitation, 10 μl of antibody-coupled beads washed in extraction buffer were incubated for 1 h at 4°C with 500 μl synapto- somal extract. After washing, bound proteins were eluted with sample buffer without reducing agent for 5 min at 60°C. The eluate was incubated for 5 min at 90°C in the presence of 50 mM dithiothreitol as a reducing agent. SDS-PAGE and immunodetection were performed as described above.

DNA constructs

Bassoon fragments RB25 (aa 609–1692), RB29 (aa 1692–3263), RB30 (aa 3263–3938), and RB31 (aa 1–609) were obtained by restriction enzyme digestion of a full-length Bassoon cDNA plasmid (Drechsel et al., 2003); Bassoon fragments RB35 (aa 1653–2348), RB38 (aa 2349– 2469), RB40 (aa 2470–2928), RB42 (aa 2929–2978), RB44 (aa 2979– 3263), RB45 (aa 2088–2348), and RB46 (aa 1653–2087) were gener- ated by amplifying the corresponding regions from a full-length rat Bassoon cDNA construct by PCR. A RIBEYE fragment encoding the A domain (aa 1–563) was obtained by PCR amplification of the corresponding se- quence using a rat genomic DNA as a template. Fragments encoding the B domain of RIBEYE (aa 564–988) or CBP1 (aa 1–430) were amplified.
Molecular Composition of the Ribbon Synapse


The presynaptic active zone protein Bassoon is essential for photoreceptor ribbon synapse formation in the retina. Neuron. 37:775–786.


Turner, J., and M. Crossley. 2001. The CtBP family: enigmatic and enzymatic


