Neuronal cholinergic synapses play important roles in both the PNS and CNS. However, the mechanisms that regulate the formation, maturation, and stability of neuronal cholinergic synapses are poorly understood. In this study, we use the readily accessible mouse superior cervical ganglion (SCG) and submandibular ganglion (SMG) to examine the assembly of the postsynaptic complex of neuronal cholinergic synapses. We find that novel splicing forms of PSD93 (postsynaptic density 93) are expressed in SCG. By immunostaining, we show that PSD93 proteins precisely colocalize with neuronal nicotinic acetylcholine receptors (nAChRs) at synapses of the SCG and SMG. Subcellular fractionation demonstrates that PSD93 is enriched in the PSD fraction of SCG, and coimmunoprecipitation shows that PSD93 and neuronal nAChRs form a complex in vivo. Furthermore, two additional components of the well-characterized glutamatergic postsynaptic complex, GKAP/SAPAP (guanylate kinase domain-associated protein/synapse-associated protein-associated protein) and Shank/ProSAP family proteins, are also present at neuronal cholinergic synapses. To assess the function of this protein complex at neuronal cholinergic synapses in vivo, we examined ganglia in mice that lack PSD93. We find that neuronal cholinergic synapses form properly in PSD93 null mice. After denervation, however, synaptic clusters of nAChRs disassemble much faster in mice lacking PSD93 than those in wild-type mice. These results demonstrate that PSD93 is a key component of the postsynaptic scaffold at neuronal cholinergic synapses and plays an important role in synaptic stability. In addition, these results suggest that the mechanism of postsynaptic scaffolding is conserved between neuronal cholinergic and glutamatergic synapses.

Key words: PSD93; PSD95; MAGUKs; cholinergic; nicotinic acetylcholine receptor; synapse; stability; denervation; superior cervical ganglion; submandibular ganglia
ciliary ganglion indicate that the large cytoplasmic loop of the α3 subunit of neuronal nicotinic acetylcholine receptors (nAChRs) is required for the postsynaptic targeting and/or anchoring of the receptor (Williams et al., 1998), suggesting that it may contain a nucleation site for postsynaptic scaffolding. In this report, we provide evidence that the postsynaptic complex at cholinergic synapses of mammalian autonomic ganglia resemble those at glutamatergic synapses. Specifically, in both the sympathetic superior cervical ganglion (SCG) and the parasympathetic submandibular ganglion (SMG), PSD93 proteins localize to neuronal cholinergic synapses and form a complex with neuronal nAChRs in vivo. In addition, the scaffolding proteins GKIAP/SAPAP, which binds directly to PSD93, and Shank/ProSAP, which interacts with GKIAP/SAPAP, are parts of this postsynaptic complex. Furthermore, analysis of mutant mice demonstrates a role for PSD93 in the stabilization of neuronal nAChRs at the postsynaptic membrane. While this manuscript was in preparation, a paper by Conroy et al. (2003) reported that neuronal MAGUKs are present at and regulate the function of cholinergic synapses in parasympathetic chick ciliary neurons. Together, these data suggest that PSD93 family-containing complexes may be a general feature of neuronal cholinergic synapses.

Materials and Methods

**Animals.** Mice with a targeted mutation of the PSD93 gene have been described previously (McCue et al., 2001). For derangement experiments, PSD93−/− mice were crossed with the GFP-J line of transgenic mice, which express green fluorescent protein specifically in neurons; these mice have been described previously (Feng et al., 2000). Mice were anesthetized with Ketamine-XXylazine by intraperitoneal injection and perfused with 4% paraformaldehyde (PFA) in PBS, rinsed in PBS, incubated with 0.1M glycine in PBS for 30 min. The supernatant was either diluted with 2× SDS loading buffer for SDS-PAGE or scraped into 2 ml of PBS for coimmunoprecipitation studies.

**Antibodies.** The mouse monoclonal antibody to the synaptic vesicle protein SV2 was obtained from the Developmental Studies Hybridoma Bank (University of Iowa, Iowa city, IA) (Buckley and Kelly, 1985). The mouse monoclonal anti-synaptophysin antibody was from Synaptic Systems (Gottingen, Germany). The rabbit polyclonal anti-synaptophysin was from Zymed Laboratories (South San Francisco, CA). The mouse monoclonal anti-PDZ antibody was from Upstate Biotechnology (Lake Placid, NY). The rabbit polyclonal PSD93-specific antibody was from Chemicon (Temecula, CA). The rabbit polyclonal anti-GKAP antibody was a generous gift from Dr. Morgan Sheng (Massachusetts Institute of Technology, Cambridge, MA) (Kim et al., 1997). The rabbit polyclonal anti-ProSAP1 was a generous gift from Drs. Tobias Bockers (Freie University, Berlin, Germany) and Ekart Gundelfinger (Leibniz Institute for Neurobiology, Magdeburg, Germany) (Boeckers et al., 1999a). The mouse monoclonal PSD95 antibodies were from Upstate Biotechnology and Transduction Laboratories (Lexington, KY). The rabbit polyclonal anti-SAP102 antibody was a generous gift from Dr. Michael Ehlers (Duke University, Durham, NC). The rabbit polyclonal anti-PSD97 antibody was from Affinity BioReagents (Golden, CO). The rat monoclonal MAB270 (anti-β2) antibody was from Developmental Studies Hybridoma Bank (Lindstrom, 1996). The rabbit polyclonal anti-AChR α3 antibody was generated and affinity purified in the laboratory using a fusion protein from a region of the large cytoplasmic loop that is unique to the α3 subunit. FITC-conjugated goat anti-mouse IgG was from Roche Diagnostics (Indianapolis, IN). All other FITC- or Cy3-conjugated secondary antibodies were from Jackson Immunoresearch (West Grove, PA).

**Antibody characterization.** COS-7 or QT6 cells were transfected with PSD93Δ, PSD95-GFP, SAP97-GFP, or SAP102-myc. Expression of each neuronal MAGUK was verified by either GFP or myc-tag expression or by staining and blotting with the anti-PDZ antibody that recognizes all neuronal MAGUKs. To test antibody specificity, transfected COS-7 cells were fixed for immunostaining, and transfected QT6 cells were lysed for immunoblotting. Each antibody was then tested for their specificity by examining their cross-reactivity with other neuronal MAGUKs using transfected cells. For PSD95, we tested the following antibodies: mouse monoclonal anti-PSD95 from Affinity BioReagents (clone 6G6-IC9, catalog #MA1-045); and clone 7E3-188, catalog #MA1-046, Upstate Biotechnology (clone K28–86.2, catalog #05-494), and Transduction Laboratories (clone 16, catalog #610496), and rabbit polyclonal anti-PSD95 from Chemicon (catalog #1596). Only the mouse monoclonal anti-PSD95 from Upstate Biotechnology and Transduction Laboratories showed specificity. The rabbit polyclonal anti-PSD93 from Chemicon (catalog #ABS168) and rabbit polyclonal anti-SAP97 antibody from Affinity BioReagents (catalog #PA1-044) were specific. For SAP102, a guinea pig anti-SAP102 (Aoki et al., 2001) and a rabbit anti-SAP102 (Lau et al., 1996) were specific.

**Cell culture.** For transfection, COS-7 cells or qaural fibroblast QT6 cells were plated in 35 mm tissue culture dishes with or without glass coverslip and cultured until 50% confluent. Cells were transfected with 1 μg of DNA of each construct per well using Lipofectamine 2000 (Invitrogen, Carlsbad, CA) and allowed to grow for 48 hr after transfection. Cells were either lysed in 1× SDS loading buffer for SDS-PAGE and fixed for immunostaining or scraped into 2 ml of PBS for communoprecipitation studies.

**Immunohistochemistry.** Mice were deeply anesthetized, and submandibular and superior cervical ganglia from mutant or control mice were dissected and embedded in OCT compound (TissueTek, Torrance, CA), frozen in 2-methylbutane chilled in liquid nitrogen, and sectioned at 7 μm on a Leica CM 1850 cryostat.

For antibody staining, sections were rinsed with PBS for 5 min, blocked with 2% BSA and 5% normal goat serum in PBS for 1 hr, and incubated with primary antibody for 3 hr to overnight. Sections stained with the anti-AChR β4 antibody were treated with an antigen retrieval procedure. These sections were first fixed for 5 min in 4% paraformaldehyde (PFA) in PBS, rinsed in PBS, incubated with 0.1M glycine in PBS for 10 min, rinsed with PBS, and incubated for 1 hr at 55°C in 0.1% SDS in PBS. Sections were then rinsed with PBS and blocked in 2% BSA, 5% normal goat serum, and 0.2% Triton X-100 (Sigma, St. Louis, MO) for 1 hr. Primary antibody was added to the sections and incubated for 3 hr to overnight. All sections were incubated with the proper secondary antibodies for 1–3 hr, washed with PBS, and mounted in 90% glycerol with 0.1% p-phenylendiamine.

**PSD preparation.** The PSD fractions of rat SCGs and mouse brains were prepared as described previously (Carlin et al., 1980; Cho et al., 1992; Ehlers, 2003). Briefly, mouse brain homogenates were centrifuged (100,000 × g) to remove nuclei and tissue debris. The supernatant was centrifuged previously described at 100,000 × g to yield a crude membrane (P2). The P2 pellet was washed and recentrifuged to obtain the washed crude synapticosomal fraction (P2’). The P2’ fraction was subjected to hypo-osmotic shock and centrifuged at 25,000 × g to yield the lysed synapticosomal membrane fraction (P3). Synaptic plasma membrane (SPM) fraction was obtained by sucrose gradient centrifugation of resuspended P3. To obtain the PSD11 fraction, SPM was washed and pelleted twice with 0.5% Triton X-100.

**SCG lyase preparation and communoprecipitation.** SCGs were homogenized in a dounce homogenizer (100 strokes) with lysis buffer containing Complete protease inhibitor (Roche Diagnostics), 1% Nonidet P-40 (Sigma), 0.5% deoxycholate (Sigma), and 0.1% SDS (Sigma). The lysate was incubated on ice for 1 hr and then centrifuged at 18,000 × g for 30 min. The supernatant was either diluted with 2× SDS-PAGE loading buffer for Western blot or used in communoprecipitation as described below.

For communoprecipitation, SCG lysates were incubated with 2 μg of either polyclonal rabbit anti-AChR α3 antibody or rabbit anti-synapsin overnight with gentle mixing at 4°C. Twenty-five microliters of protein G-agarose beads (1:1 slurry; Invitrogen) were then added and incubated for 2–4 hr at 4°C. Samples were centrifuged at 1000 × g for 2 min, and the supernatant was removed. Samples were washed six times for 5 min each with washing buffer (PBS with protease inhibitors and 1% Triton X-100; Sigma). Proteins were eluted from agarose beads by addition of SDS loading buffer and boiling for 5 min.

In situ hybridization. Mouse sequences encoding PSD93α (nucleotide 1–319; GenBank accession number NM_022282), PSD93Δ (nucleotide
1–420; GenBank accession number XM_133619), PSD93e (nucleotide 1–420), and PSD93c (nucleotide 113–470) were amplified by PCR and subcloned into pBluescript II SK+ vector (Stratagene, Palo Alto, CA). Plasmids were linearized with restriction enzymes that generated 5’ overhangs. Digoxigenin (DIG)-labeled RNA probes were synthesized with the MAXiScrip in vitro RNA synthesis kit (Ambion, Austin, TX) using T3 and T7 RNA polymerases. DIG-labeled probes were quantified by dot blot and probed with alkaline phosphatase-conjugated anti-DIG antibody (Roche Diagnostics).

Adult ICR mice were killed with halothane in accordance with National Institutes of Health and institutional protocols. SCGs were dissected, embedded in OCT compound (TissueTek), and frozen in 2-methylbutane chilled in liquid nitrogen. Twenty micrometer sections were cut on a cryostat and placed on Superfrost Plus slides (Fisher Scientific, Swannee, GA).

For hybridization, sections were fixed with 4% PFA in PBS for 20 min at 4°C and washed three times for 3 min each with PBS. Sections were then acetylated for 10 min at room temperature in TEA buffer (142 ml of dH2O, 4.65 ml of 5 mM NaCl, 2.01 ml of triethanolamine, 0.975 ml of 8 mM HCl; mix well and add 0.375 ml of acetic anhydride just before use). After being washed three times with PBS, sections were incubated with hybridization buffer (50% formamide, 5X SSC, 5X Denhardt’s solution, 500 μg/ml salmon sperm DNA, and 250 μg/ml yeast tRNA) for 2 hr at room temperature and then with DIG-labeled RNA probes (400 μg/ml in hybridization buffer, heated at 80°C for 10 min, cooled on ice) overnight at 65°C in a chamber humidified with 50% formamide in 5X SSC.

After hybridization, sections were first rinsed two times for 5 min each with 5X SSC at room temperature followed by four times for 1 hr each with 0.2X SSC at 65°C. Sections were then cooled to room temperature and incubated with blocking buffer (10% normal goat serum and 0.2% blocking reagent; Roche Diagnostics) for 1 hr, followed by incubation with alkaline phosphatase-conjugated anti-DIG antibody (1:2000; Roche Diagnostics) overnight at 4°C. Sections were washed (four times for 10 min each) with TBS (150 mM NaCl and 10 mM Tris-HCl, pH 7.5) and incubated with color detection buffer (100 mM NaCl, 50 mM MgCl2, 0.24 mg/ml levamisole, and 100 mM Tris-HCl, pH 9.5) for 5 min. Color reactions were developed in the presence of nitro blue tetrazolium (NBT) (0.35 mg/ml) and 5-bromo-4-chloro-3-indolyl-phosphate (BCIP) (0.175 mg/ml) in color detection buffer. Color reactions were terminated by incubating membranes in TE buffer. A total of 2 × 10^6 clones were screened with the anti-PDZ antibody, and 85 positive clones were recovered.

For library screening with PSD93 cDNA probes, mouse sequence encoding PSD93e (nucleotide 181–1243; GenBank accession number NM_011807) was amplified by PCR and subcloned into pBluescript II SK+ vector (Stratagene). DNA was labeled with DIG using the Dig High Prime DNA Labeling and Detection kit (Roche Diagnostics). Random-primed cDNA library was plated at a density of 1 × 10^5 plaques per 15 cm plate. Phage plates were transferred to nylon membranes, denatured by autoclaving for 3 min, and cross-linked to the membrane by UV cross-linking. Membranes were washed twice with 2X SSC, prehybridized for 2 hr at 45°C in Easy-Hyb solution (Roche Diagnostics), and then incubated with DIG-labeled cDNA probes (50–10 ng/ml) and NBT/BCIP solution overnight at 45°C. After hybridization, membranes were washed twice (10 min each) with 2X SSC and 0.1% SDS at room temperature, followed by washing twice (30 min each) with 0.2X SSC and 0.1% SDS at 65°C. Membranes were rinsed three times with TBS, blocked with 1X blocking solution (Roche Diagnostics) for 2 hr and then incubated with anti-DIG antibody (1:5000 in 1X blocking solution) for 4 hr, followed by washing six times with TBS plus 0.3% Tween 20. Color reactions were developed in the presence of NBT and BCP as described above. A total of 1 × 10^6 clones were screened, resulting in the identification of 45 positive clones.

Reverse transcription-PCR. Poly(A)+ RNA from mouse SCGs and brains were prepared as described previously (Feng et al., 1998a). cDNA was synthesized with Superscript reverse transcriptase (Stratagene) using oligo-dT as a primer. PCR primers were designed to specifically amplify ~100 nucleotide fragments of various splicing forms of PSD93 and a 1.5 kb fragment of EF-1α (elongation factor-1α) as a control (Gruber and Levine, 1997). The following primers were used: PSD93eF (CTGAGCTTCTGACCATCCTAGG) and PSD933R (CGTGGCCGATAGATGCTG); PSD93BF (AGCTGCCTGCTCGTCTGAGCT) and PSD93R; PSD93yF (GTGAAAGGCTATGGCAACATGCTG) and PSD93yR (CGATTGTGACATGTGCTG); PSD93BF (GGAGGAGGCCCTTCTATGCG) and PSD93R; PSD93eF (CCCACTCTGATGTGACAGC) and PSD933R; EF1αF (GAGAAGGACCTTTCTATGCG) and EF1αR (CAGAACCTCTTCTCTCTGGAG). Semiquantitative PCR was performed by a modification of the method of Freeman et al. (1994). Briefly, EF-1α RNA was used to calibrate the relative amount of RNA in SCG and brain samples, and cycle number was adjusted to be within the linear range of amplification. Once optimal conditions were determined, PCR was performed to compare the relative abundance in the SCG and brain of each alternatively spliced form.

Results

PSD93 colocalizes with neuronal nAChRs at mammalian cholinergic synapses

To analyze the molecular mechanisms that regulate the formation, stability, and plasticity of neuronal cholinergic synapses, we chose two autonomic ganglia in the mouse: the sympathetic SCG and the parasympathetic SMG. This system has several advantages. First, the majorities of synapses in both ganglia are cholinergic. This relative homogeneity simplifies the correlation of morphological and electrophysiological differences. Second, synapses within SCG and SMG are accessible and amenable to experimental manipulations in vivo (Kawa and Roper, 1984; Feng et al., 1998a). Third, using mouse ganglia permits molecular genetic approaches (Feng et al., 1998a).

We showed previously that neither rapsyn, the primary scaffolding protein of the cholinergic neuromuscular junction, nor gephyrin, the primary scaffolding protein of glycinergic and GABAergic synapses, is present at neuronal cholinergic synapses in the SCG (Feng et al., 1998a). We began the present study by asking whether MAGUKs are present at these synapses. To this...
end, we first stained SCG sections with a monoclonal antibody (anti-PDZ) that recognizes the conserved PDZ domains of all four neuronal MAGUKs. To mark synapses, sections were costained with an antibody to the synaptic vesicle protein, synaptophysin. SCG neurons have short dendrites (Purves and Lichtman, 1985), and synapses in SCG are concentrated in neuropil-rich regions between neurons and appear as punctate patches (Feng et al., 1998a). As shown in Figure 1a, the anti-PDZ antibody strongly stains SCG synapses. We found previously that neuronal nAChRs cluster at synapses in mouse SCG (Feng et al., 1998a). Double immunostaining shows that most PDZ-rich puncta colocalize with the β4 subunit (Fig. 1b).

In contrast to synapses in the SCG, those of the SMG are axo-somatic (Lichtman, 1977). The large size of these synapses and their localization on cell bodies make them easier to identify and visualize (Purves and Hadley, 1985; Snider, 1987; Gan et al., 2003). To test whether MAGUKs also localize at SMG synapses, we double stained SMG sections with the anti-PDZ antibody and either the anti-β4 antibody (Fig. 1c) or the anti-β2 antibody (data not shown). As in SCG, neuronal nAChRs are clustered at synapses on neuronal cell bodies, as indicated by their colocalization with the presynaptic vesicle proteins synaptophysin and SV2 (data not shown). Similarly, patterns of anti-PDZ antibody staining and anti-β4 staining match precisely (Fig. 1c). Thus, in both sympathetic and parasympathetic ganglia, MAGUKs are present at neuronal cholinergic synapses.

To determine which MAGUKs are present at neuronal cholinergic synapses, we stained SMG sections with antibodies specific for PSD95/SAP90, PSD93/chapsyn-110, SAP97, and SAP102 (Fig. 1d–h). When tested in transfected COS7 cells, all four antibodies stained their respective proteins strongly and specifically (Fig. 1i,j and data not shown) (for details of antibody characterization, see Materials and Methods). In SMG, PSD93 is highly expressed and localized to the synapse, as evidenced by its precise colocalization with synaptophysin (Fig. 1d) and the β2 subunit of nAChRs (Fig. 1e). We also observed weak synaptic staining of PSD95 in SMG (Fig. 1f), but its levels were much lower than those of PSD93, although both antibodies stained transfected cells to a similar extent (data not shown). No synaptic staining of SAP97 or SAP102 was detected (Fig. 1g,h). Similar results were also obtained from SCG preparations (data not shown). Consistent with the immunostaining results, Western blot analysis of SCG lysate showed abundant expression of PSD93 but not PSD95, SAP97, or SAP102 (Fig. 1k). Thus, PSD93 is the predominant MAGUK expressed at neuronal cholinergic synapses of mouse autonomic ganglia.

Characterization of PSD93 splicing forms at neuronal cholinergic synapses

Neuronal MAGUKs show considerable alternative splicing in brain (Lue et al., 1994; Muller et al., 1995, 1996; Brennan et al., 1996; Chetkovich et al., 2002; Rumbaugh et al., 2003). Specifically, alternative splicing at the N termini regulates synaptic trafficking and targeting of PSD95 and PSD93 (Firestein et al., 2000; Chetkovich et al., 2002; Rumbaugh et al., 2003). To facilitate the identification of unique forms at neuronal cholinergic synapses, we constructed a mouse SCG cDNA library and screened it with both the anti-PDZ antibody and a PSD93-specific CDNA probe. The screen with the anti-PDZ antibody also tested whether synaptic staining by the anti-PDZ antibody may represent a novel MAGUK. Among the 85 positive clones isolated by screening with the anti-PDZ antibody, 74 encoded PSD93, five encoded PSD95, and one encoded SAP102; no novel homologues were identified. The relative abundance of PSD93 clones in the SCG cDNA library is consistent with our immunostaining results showing that PSD93 is the predominant MAGUK at neuronal cholinergic synapses.

Sequence analysis of PSD93 clones from both the antibody and cDNA probe screens revealed three novel splicing forms of PSD93. Previous studies of rat brain identified three alternative splicing forms of PSD93 (PSD93α, PSD93β, and PSD93 γ, corresponding to PSD93–5′b, PSD93–5′a, and PSD93–5′c according to Brennan et al., 1996), with the most abundant being PSD93α (Brennan et al., 1996). Surprisingly, the previously identified brain splicing forms were not isolated from the SCG library, with the exception of one clone encoding the PSD93β isoform. Of the three new isoforms (PSD93δ, PSD93ε, and PSD93ζ) that we identified, PSD93δ is most abundant. Both PSD93δ and PSD93ε diverge from PSD93α (the most abundant brain isoform) upstream of alanine 69, whereas PSD93ζ diverges from PSD93α upstream of lysine 15 (Fig. 2a). Compared with PSD93α, the N termini of PSD93δ and PSD93ζ are 35 and 124 amino acids longer, respectively, and PSD93ζ is 42 amino acids shorter. The PSD93δ isoform contains two pairs of N-terminal cysteines for potential palmitoylation, a modification that has been shown to mediate synaptic targeting of PSD95 (Topinka and Bredt, 1998; Craven et al., 1999). Interestingly, the PSD93ζ isoform does not have the cysteine pairs. Rather, it contains an L27 domain, which
has been shown to mediate synaptic targeting of the PSD95β isoform through protein–protein interactions (Chetkovich et al., 2002). Neither a cysteine pair nor an L27 domain is present in the PSD93ζ (Fig. 2a). These data suggest that ganglionic neurons may use alternative splicing to regulate the synaptic targeting of PSD93.

To verify our observations from the SCG library screening, we first used reverse transcription (RT)-PCR to estimate the relative abundance of the splicing variants in the SCG and the brain. The transcripts of PSD93α, PSD93β, and PSD93γ isoforms are readily detected from mouse brain by RT-PCR but are not detected from the SCG using the same conditions. In contrast, the three new isoforms, PSD93δ, PSD93ε, and PSD93ζ, are detected in both the brain and SCG (Fig. 2b). The transcript for PSD93δ is equally abundant in the brain and the SCG, whereas transcripts for PSD93ε and PSD93ζ are more abundant in the brain than in the SCG (Fig. 2b).

We next examined the expression of the various isoforms in SCG by in situ hybridization. We made probes specific for the PSD93α, PSD93δ, PSD93ε, and PSD93ζ isoforms. Little signal was detected in the SCG with the probe for PSD93α (Fig. 2c), although strong signals were detected in the brain with the same probe (data not shown). This is consistent with our results from the library screening and RT-PCR, indicating that PSD93α, although abundant in the brain, is expressed at low levels if at all in the SCG. In contrast, PSD93δ, PSD93ε, and PSD93ζ were all expressed in SCG neurons, with the PSD93δ probe giving the strongest signal (Fig. 2c). Together, these data strongly suggest that ganglionic neurons express unique splicing forms of PSD93.

To characterize PSD93 splicing forms at the protein level, we expressed various splicing forms of PSD93 in quail QT6 fibroblast cells. The size differences of the major brain and SCG splicing forms make it possible to distinguish them on Western blots (Fig. 2d), thus allowing us to predict the isoforms expressed in SCG on the basis of their apparent molecular weights. A single major band at ~120 kDa was detected in SCG lysate when probed with a PSD93-specific antibody that is able to recognize all N-terminal splicing forms. This band comigrates with the PSD93δ isoform transfected in QT6 cells (Fig. 2d). Although conclusive evidence will require the generation of isoform-specific antibodies, these data suggest that PSD93δ is the predominant PSD93 splicing form expressed at ganglionic cholinergic synapses.

PSD93 is enriched in PSD and forms a complex with neuronal nAChRs in vivo

Maguk are core components of the PSD at glutamatergic synapses (Kennedy, 2000). To test whether PSD93 is also a component of the PSD at neuronal cholinergic synapses, we prepared PSD fractions from rat SCG and probed them with PSD93-specific antibodies. As shown in Figure 3a, PSD93 is highly enriched in the PSD fraction prepared from rat SCG. In contrast, PSD95 was not detected in either SCG lysate or SCG PSD fractions (Fig. 3b).

Maguk form a complex with glutamate receptors through either a direct interaction (with NMDA receptors) or an indirect interaction (with AMPA receptors via Stargazin), and these interactions have been implicated in regulating the synaptic targeting and anchoring of glutamate receptors at the postsynaptic membrane (Sheng and Sala, 2001; McGee and Bredt, 2003). To determine whether PSD93 forms a complex with neuronal nAChRs, we performed coimmunoprecipitation from the SCG with a polyclonal antibody against the α3 subunit of the nAChR and probed the immunoprecipitates with the monoclonal anti-
antibodies. The anti-GKAP antibody recognizes all four members present at neuronal cholinergic synapses, we examined the expression (Kennedy, 2000). The supramolecular complex at the postsynaptic membrane contains multiple scaffolding proteins (Garner et al., 2000; Kennedy, 2000; Sheng and Sala, 2001; McGee and Bredt, 2003). In addition to interacting with NMDA receptors and nNOS, MAGUKs also interact with the GKAP/SAPAP family of proteins. GKAP/SAPAP proteins further interact with a variety of proteins, including Shank/ProSAP, another class of multidomain scaffolding proteins (Boeckers et al., 1999a; Naishbitt et al., 1999; Sheng and Kim, 2000). Shank/ProSAP proteins, in turn, bind to the actin-binding protein Cortactin, potentially linking the complex to the cytoskeleton. Furthermore, Shank/ProSAP proteins also bind to Homer, a metabotropic glutamate receptor binding protein (Sheng and Kim, 2000). These interactions increase the complexity and regulatory capability of the postsynaptic complex (Kennedy, 2000). To test whether a similar core group of scaffolding proteins is present at neuronal cholinergic synapses, we examined the expression of GKAP/SAPAP and Shank/ProSAP using specific antibodies. The anti-GKAP antibody recognizes all four members of the family (Kim et al., 1997) (J. M. Welch, D. Wang, and G. Feng, unpublished results). The anti-Shank antibody is specific for Shank1/ProSAP2 (Boeckers et al., 1999a). We found that both GKAP/SAPAP and Shank/ProSAP are expressed in the SMG and are highly concentrated at synapses (Fig. 4a,b). The precise synaptic localization of GKAP/SAPAP and Shank/ProSAP and their ability to interact with the actin cytoskeleton provide a possible mechanism whereby nAChRs might be tethered to the cytoskeleton, thus stabilizing the receptors at the synapse.

**Formation and stability of neuronal cholinergic synapses in the absence of PSD93**

To determine the function of PSD93 at neuronal cholinergic synapses, we examined synapses in the SMG of PSD93-deficient mice. PSD93 knock-out mice are viable, fertile, and outwardly normal (McGee et al., 2001), suggesting that the basic function of the autonomic nervous system is intact. No PSD93 proteins were detected in the SMG of the knockout mice (Fig. 5a). To determine whether neuronal nAChRs are anchored at synapses in the absence of PSD93, we stained SMG sections for the presynaptic vesicle protein synaptophysin and the β2 and β4 subunits of the nAChRs. We found that neuronal nAChRs are clustered at synapses in the SMG of PSD93 mutant mice and that receptor clusters are qualitatively indistinguishable from those of wild-type SMG (Fig. 5c,d), indicating that PSD93 is dispensable for clustering of neuronal nAChRs at ganglionic synapses. Staining with the anti-PDZ antibody showed a low level of residual staining, comparable with the PSD95 staining seen in the wild-type SMG (Fig. 5b). Moreover, staining for individual members of MAGUKs did not reveal any detectable upregulation (data not shown). Although we cannot exclude the possibility that other MAGUKs may play a role at this synapse, our data do not show compensation for the loss of PSD93 by other MAGUKs.

We next examined the localization of other scaffolding proteins in the absence of PSD93. In the PSD93 knock-out mice, the synaptic localization of both GKAP/SAPAP and Shank/ProSAP documented above for wild-type mice (Fig. 4) is maintained (Fig. 5e,f), indicating that interactions with PSD93 are not critical for the synaptic localization of either protein. Although PSD93 is not essential for clustering nAChRs, it might play a role in stabilizing receptor clusters. Denervation of SMG allows study of the stability of neuron–neuron synapses in vivo. It eliminates synaptic adhesion and presynaptic factors that promote receptor clustering, leading to the disassembly of the postsynaptic complex at a rate inversely proportional to its structural integrity.

We found previously that, in the SMG of wild-type mice, presynaptic nerve terminals degenerate 3 d after denervation, whereas neuronal nAChR clusters persist (Feng et al., 1998a). In this study, we used the SMG because the axo-somatic synapses are much easier to visualize and quantify. We first mated PSD93 knock-out mice to a line of Thy1-GFP mice in which the preganglionic axons and terminals, but not postganglionic neurons, are specifically labeled with GFP (Feng et al., 2000). This enabled us to transnet the preganglionic axon bundles innervating the SMG under a fluorescent dissecting microscope with minimal perturbation and tissue damage (Fig. 6a,b). Denervations were performed unilaterally, and both denervated and contralateral non-
denervated (as controls) SMG were dissected 2, 4, and 7 d postoperatively. Full denervation of the SMG was verified at each time point by immunostaining SMG sections for the presynaptic protein synaptophysin. Receptor clusters were assessed by immunostaining SMG sections for the presynaptic 2 subunit of nAChRs using the MAB270 monoclonal antibody (Whiting et al., 1987; Feng et al., 1998a). Because each neuron in the SMG is easily visualized in these sections, we quantified receptor clusters by counting the number of neurons that had receptor clusters as a percentage of the total neurons on the section.

In both wild-type and PSD93 knock-out mice, presynaptic terminals degenerated to near completion 2 d after denervation, as indicated by the lack of synaptophysin staining (Fig. 6c,d). In wild-type SMG, the majority of neurons maintain nAChR clusters after denervation for 2 and 4 d (85 and 67%, respectively). By day 7 after denervation, the fraction of neurons with receptor clusters was reduced to 24% (Fig. 6c). In contrast, in the PSD93−/− SMG, only 23 and 7% of neurons retained nAChR clusters after denervation for 2 and 4 d, respectively (Fig. 6d). Thus, postsynaptic receptor clusters at neuronal cholinergic synapses of PSD93−/− mice disassemble at an approximately threefold to fourfold accelerated rate when denervated (Fig. 6c).

To test whether deletion of PSD93 also accelerates the disassembly of other postsynaptic proteins, we compared synaptic localization of GKAP/SAPAP and Shank/ProSAP in denervated SMG of wild-type and PSD93 knock-out mice. Unlike nAChRs, synaptic clusters of both GKAP/SAPAP and Shank/ProSAP in wild-type SMG persisted for 7 d after denervation (Fig. 7a,c). Furthermore, the quality and quantity of clusters of these two proteins in PSD93−/− SMG were indistinguishable from those of wild-type SMG (Fig. 7b,d). These data indicate that the synaptic localization of GKAP/SAPAP and Shank/ProSAP proteins does not depend on their interactions with PSD93.

Discussion

Neuronal cholinergic synapses play important roles in both the PNS and CNS (Dani and Heinemann, 1996; Role and Berg, 1996). Alterations in nAChR function and expression have been implicated in several neurological disorders, including Alzheimer’s disease, epilepsy, and schizophrenia (Lena and Changeux, 1997; Lindstrom, 1997; Perry et al., 1999). However, the molecular mechanisms regulating the assembly and maintenance of the postsynaptic complex at neuronal cholinergic synapses are still poorly understood. In this report, we provide evidence that PSD93, GKAP/SAPAP, and Shank/ProSAP are all present at the postsynaptic membrane of autonomic neuronal cholinergic synapses. This core complex plays a key role in regulating the stability of neuronal cholinergic synapses, likely by interacting with the actin cytoskeleton. Furthermore, by providing docking sites for various signaling molecules, this core scaffold also provides a link from receptor activation to downstream signaling events.

It has been well established that the multidomain proteins of MAGUKs, GKAP/SAPAP family, and Shank/ProSAP family are key scaffolding molecules at the glutamatergic synapses (Garner et al., 2000; Kennedy, 2000; Sheng and Sala, 2001; McGee and Bredt, 2003). Our studies demonstrate that these key scaffolding proteins are also important components of neuronal cholinergic synapses. A recent study has shown that MAGUKs are also present in chick ciliary ganglion neurons and regulate the function of neuronal cholinergic synapses (Conroy et al., 2003). MAGUKs (Rafael et al., 1998) and GKAP/SAPAP proteins (Welch, Wang, and Feng, unpublished results) have also been found at the neuromuscular junction. These data raise the intriguing possibility that all excitatory synapses use the same group of core scaffolding proteins. On the contrary, none of these scaffolding proteins have been found at GABAergic or glycinergic synapses, suggesting the existence of a completely different postsynaptic complex at inhibitory synapses; this scaffold contains gephyrin, which has not been found at excitatory synapses as a core component (Naisbitt et al., 1997; Feng et al., 1998a,b; Rao et al., 2000) (Welch, Wang, and Feng, unpublished results).

One of the main features of PSD scaffolding proteins is that they all contain multiple protein–protein interacting domains, and each domain can interact with multiple signaling or adaptor proteins (Sheng and Pak, 2000). This has several important implications. First, by recruiting various signaling proteins to the complex, they provide a link from receptor activation to multiple downstream pathways, such as protein phosphorylation (Seabold et al., 2003), transcriptional regulation (Hering and Sheng, 2002), and cytoskeletal interactions (Brennan et al., 1998; Boeckers et al., 1999a; Naisbitt et al., 1999; Passafaro et al., 1999), thus transducing both short-term and long-term effects of synaptic activation. Second, by providing multiple docking sites for signaling and structural proteins, they enable neurons to regulate synaptic structure and function at multiple points for fine tuning.
Third, the availability of multiple protein–protein interaction domains enables different types of synapses or the same type of synapse on different types of neurons to assemble unique signaling cascades, thus rendering different synapses with unique functions. For example, Citron, a PSD95-binding protein, is localized at glutamatergic synapses on GABAergic interneurons but not on pyramidal neurons in the hippocampus (Zhang et al., 1999). Previously, little was known about the postsynaptic signaling complex at neuronal cholinergic synapses. The identification of core scaffolding components provides a starting point to elucidate the molecular architecture of this complex.

The N-terminal sequences play important roles in the synaptic targeting of MAGUKs. For example, dual palmitoylation through a pair of N-terminal cysteines is essential for targeting PSD95 to the postsynaptic membrane (Topinka and Bredt, 1998; Graven et al., 1999). Recent studies also suggest that a variety of mechanisms are used in the synaptic targeting of MAGUKs through the N-terminal sequences, and alternative splicing may regulate this process. For example, PSD95β, an N-terminal splicing variant of PSD95, is not palmitoylated. Instead, its synaptic localization is mediated through the L27 domain unique to this splicing form (Chetkovich et al., 2002). Three N-terminal splicing forms were noted during the initial cloning of PSD93 (Brenman et al., 1996). In this study, we identified three additional N-terminal splicing variants. Among the three new splicing forms identified, one (PSD93δ) contains two pairs of N-terminal cysteines, one (PSD93ζ) has an L27 domain, and the other (PSD93ε) contains neither motif. It has been shown that the first 30 amino acids in PSD93α are required for synaptic targeting but palmitoylation is not (Firestein et al., 2000), suggesting a novel mechanism for targeting PSD93α to the postsynaptic membrane. Interestingly, the N-terminal sequence of PSD93δ, the predominant splicing form at neuronal cholinergic synapses, is distinct from that of PSD93α and bears no homology to any known sequences in the database, suggesting that yet another type of mechanism may be involved in the synaptic targeting of PSD93. Because all three new isoforms are also expressed in the brain, we suggest that different types of neurons in the brain may regulate synaptic function through the expression of different splicing forms of PSD93.

It is not clear how nAChRs interact with PSD93 in the postsynaptic complex. MAGUKs typically bind to the last three amino acids (S/T X V) of the C termini of neurotransmitter receptors through PDZ domain interactions. Interestingly, both the N termini and C termini of all nAChRs are extracellular (Lindstrom, 1996), suggesting a different mode of interaction between PSD93 and nAChRs. Interaction of MAGUKs with neurotransmitter receptors through an intermediate protein is well studied in the case of PSD95 and AMPA receptors. Binding of PSD95 through its PDZ domains to the AMPA receptor-associated protein Stargazin regulates the synaptic targeting of AMPA receptors (Chen et al., 2000; Schnell et al., 2002), and a similar mechanism may be used by cholinergic neurons. Interestingly, two short forms of PSD93, which lack the entire latter half of the protein, including the SH3 and the GK domain, were recently identified from chick ciliary ganglion neurons (Conroy et al., 2003). It was shown that one of the short forms (chick PSD93a) interacts with neuronal nAChRs

Figure 6. Synaptic clusters of neuronal nAChRs are less stable in denervated SMG of PSD93−/− mice. a, b, To facilitate the denervation procedure of SMG, PSD93−/− mice were mated to Thy1-GFP-J mice, which express GFP in preganglionic axons and terminals but not postganglionic neurons. The pictures show an SMG before (a) and immediately after (b) denervation. c, d, Sections of wild-type (c) or PSD93−/− (d) SMG denervated for 0 d (Innervated), 2 d (DEN-2 day), 4 d (DEN-4 day), and 7 d (DEN-7 day) were double stained with anti-synaptophysin (SYN; left panels) and anti-nAChR β2 (right panels). In wild-type SMG, the majority of the nAChR β2 clusters persist 4 d after denervation (c, right panels). In contrast, most of the nAChR β2 clusters disappear 2 d after denervation in PSD93−/− SMG (d, right panels). e, Quantification of percentage of neurons containing nAChR β2 clusters in denervated wild-type (open bars) and PSD93−/− (filled bars) SMG. Numbers on top of the bars indicate the number of denervation experiments performed. Scale bar: c, d, 5 μm.
when cotransfected into HEK293 cells, and both the N-terminal sequences and the PDZ domains are important for this interaction (Conroy et al., 2003). Neither of the two short forms has been reported in mammals, and we did not detect similar short forms from our screening of mouse SCG libraries. It is therefore not clear whether the mammalian PSD93 isoforms also directly interact with nAChRs. Nevertheless, the large cytoplasmic loop of the α3 subunit of neuronal nAChRs has been shown to mediate synaptic targeting and/or anchoring the receptor in chick ciliary ganglia (Williams et al., 1998). Because this loop is also the main cytoplasmic region of the receptor, it is likely that the interaction of PSD93 with nAChRs, direct or indirect, involves this loop.

Despite intensive study, the function of MAGUKs at vertebrate synapses remains poorly understood (McGee and Bredt, 2003). Our data provide the first evidence that PSD93 plays a role in the stabilization of receptor clusters at neuronal cholinergic synapses. Although the mechanism is unclear, PSD93 may act by linking receptor clusters to the cytoskeleton via interactions with GKAP/SAPAP and Shank/ProSAP, which interact with actin through Cortactin. Alternatively, PSD93 may regulate the trafficking and targeting of nAChRs to the postsynaptic membrane. It is well documented that the density of neurotransmitter receptors in the PSD is dynamically regulated by neuronal activity (Carroll and Zukin, 2002; Malinow and Malenka, 2002; Wenthold et al., 2003), and regulation through ubiquitin-proteasome degradation (Ehlers, 2003) and PSD95-mediated trafficking and targeting both play important roles in this process (Chen et al., 2000; Shi et al., 2001; Schnell et al., 2002). Thus, the accelerated disassembly of synaptic nAChR clusters in PSD93 null mice may reflect impaired trafficking and/or targeting of nAChRs to the postsynaptic membrane. Distinguishing these mechanisms in future studies will likely help us to further understand the function of MAGUKs at cholinergic as well as glutamatergic synapses.

References