Superfluous Role of Mammalian Septins 3 and 5 in Neuronal Development and Synaptic Transmission

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The septin family of GTPases, first identified for their roles in cell division, are also expressed in postmitotic tissues. SEPT3 (G-septin) and SEPT5 (CDCrel-1) are highly expressed in neurons, enriched in presynaptic terminals, and associated with synaptic vesicles. These characteristics suggest that SEPT3 or SEPT5 might be important for synapse formation, maturation, or synaptic vesicle traffic. Since Sept5+/−/− mice do not show any overt neurological phenotypes, we generated Sept3−/− and Sept3−/− Sept5−/− mice and found that SEPT3 and SEPT5 are not essential for development, fertility, or viability. Changes in the expression of septins were noted in the absence of SEPT3, SEPT5, and both septins. SEPT5 association with other septins in brain tissue was unaffected by the removal of SEPT3. No abnormalities were observed in the gross morphology and synapses of the hippocampus. Similarly, axon development and synapse formation were unaffected in vitro. In cultured hippocampal neurons, the size of the recycling synaptic vesicle pool was unaltered in the absence of SEPT3. Furthermore, synaptic transmission at two different central synapses was not significantly affected in Sept3−/− Sept5−/− mice. These results indicate that SEPT3 and SEPT5 are dispensable for neuronal development as well as for synaptic vesicle fusion and recycling.

The septins are a multigene family of guanine nucleotide-binding, filament-forming proteins. Several septins have well-established roles in cytokinesis in a variety of cells that span a broad variety of species from yeast to mammals. The septin gene family was originally identified in the budding yeast Saccharomyces cerevisiae in a genetic screen for critical determinants of cytokinesis (23). S. cerevisiae has seven septin genes (CDC3, CDC10, CDC11, CDC12, SHS1/SEP7, SPR3, and SPR28 genes). Caenorhabditis elegans has two (UNC-59 and UNC61 genes), Drosophila melanogaster has five (Pnut, Sep1, Sep2, Sep4, and Sep5 genes), and humans have one (SEPT1) to SEPT9 (21, 46). The septins share a conserved GTPase domain that comprises the bulk of their sequence. The GTPase domain has characteristic amino acid motifs found in other P-loop GTPases (21, 46). Septin complexes assemble into filaments in vitro and these complexes vary in composition, with identified complexes comprising SEPT4, SEPT5, and SEPT8 (SEPT4/5/8; SEPT7/9b/11; and SEPT2/6/7/9) (40, 43, 54, 60). Septin complexes assemble into filaments in vitro and these have a 4- to 10-nm diameter (17, 19, 24, 29). Although they do not have inherent curvature, septin filaments are flexible and tend to roll up into rings of 0.5- to 0.7-μm diameter (34). Septin rings are found in a variety of cellular contexts in vivo: at the mother-bud neck junction in yeast (9), at the yeast cell cortex (49), at the annulus of sperm tails (26, 36), as a circumferential band in platelets (39), and in cells where actin filament organization is disrupted with cytochalasin D (34, 69). Septin complexes are found along central actin stress fibers (35, 60).

Septins have a range of functions unrelated to cytokinesis that are only just beginning to emerge, including apoptosis, vesicular transport, cytoskeletal dynamics, cell polarity, and sperm motility (30, 33, 41, 57). Interestingly, septins are abundant in the central nervous system, which is comprised largely of postmitotic neurons. They are associated with many neurological diseases such as Parkinson’s, Alzheimer’s, schizophrenia, and hereditary neuralgic amyotrophy (3, 27, 31, 38, 56), but their neurobiological function is unknown.

We originally set out to test the function of septins in the mammalian central nervous system by genetically disrupting assembly (55). At the N-terminal side of the GTPase domain, most septins contain a stretch of polybasic amino acids, which mediates their association with phosphatidylinositol phospholipids (10, 50, 76).

Subsets of septins have a high affinity for each other and form large, stable heteromeric complexes in stoichiometric ratios (17, 19, 34). Mammalian septin complexes vary in composition, with identified complexes comprising SEPT4, SEPT5, and SEPT8 (SEPT4/5/8; SEPT7/9b/11; and SEPT2/6/7/9) (40, 43, 54, 60). Septin complexes assemble into filaments in vitro and these have a 4- to 10-nm diameter (17, 19, 24, 29). Although they do not have inherent curvature, septin filaments are flexible and tend to roll up into rings of 0.5- to 0.7-μm diameter (34). Septin rings are found in a variety of cellular contexts in vivo: at the mother-bud neck junction in yeast (9), at the yeast cell cortex (49), at the annulus of sperm tails (26, 36), as a circumferential band in platelets (39), and in cells where actin filament organization is disrupted with cytochalasin D (34, 69). Septin complexes are found along central actin stress fibers (35, 60).

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the Sept5 gene in mice (47). There were several good reasons to believe that SEPT5 had a role in neurotransmitter release. First, SEPT5 was isolated as a constituent of a synaptic vesicle protein complex associated with synaptophysin (4). Second, SEPT5 was localized to the presynaptic side of the synapse in close apposition with synaptic vesicles (32). Third, it was shown that SEPT5 directly binds to the synaptic vesicle fusion protein syntaxin 1 (4). Fourth, overexpression of SEPT5 inhibited the secretion of storage granules from an insulin-secreting cell line (4), and its absence enhanced the secretion of storage granules from platelets (11). Despite the overwhelming evidence supporting a role for SEPT5 in synaptic vesicle fusion, no alterations in synaptic transmission were observed in the hippocampus of Sept5<sup>−/−</sup> mice (47). Since septins form heteromeric protein complexes with each other, another septic could have overlapping functions with SEPT5.

SEPT3 was discovered as a substrate of cyclic GMP–dependent protein kinase (72, 74) and is the only septin known to date that is exclusively expressed in postmitotic neurons (73), implying that it has no role in cell division and is likely to function in a neuron-specific process. It is intriguing to think that SEPT3 has a critical role in the synaptic vesicle cycle because it is highly enriched in presynaptic terminals of hippocampal neurons, associates with synaptic vesicles, and colocalizes with synaptophysin and dynamin I (73). To determine whether SEPT3 is the critical septic essential for synaptic vesicle trafficking or whether SEPT3 and SEPT5 function together, we now report on the targeted disruption of the Sept3 gene as well as the generation of mice lacking both Sept3 and Sept5.

**MATERIALS AND METHODS**

**Animal handling.** All procedures involving animals were in compliance with guidelines of the Animal Care Committee at The Hospital for Sick Children (Toronto, ON, Canada).

**Antibodies.** Antibodies against the mammalian septins were made in-house as previously described (2, 60, 69). The Sept3 monoclonal antibody (clone SP20) was a gift from W. G. Honer (University of British Columbia, Vancouver, BC, Canada). Antibodies to synaptophysin and MAP2 were from Chemicon (Temecula, CA). The antibody to neurofilament-H (clone SMI31) was from Covance Research Products (Berkeley, CA). The horseradish peroxidase-conjugated, Cy3-conjugated, and Cy5-conjugated anti-mouse and anti-rabbit secondary antibodies were from Jackson ImmunoResearch Laboratories (Westgrove, PA). AlexaFluor 488-conjugated secondary antibodies were from Invitrogen (Burlington, ON, Canada).

**Sept3 gene targeting vector and generation of Sept3<sup>−/−</sup> mice.** Rat Sept3 (G- septin-n) cDNA was used as a radiolabeled probe to screen the RPCI 129/SvEvTACBR mouse bacterial artificial chromosome (BAC) library by Southern blot analysis. Exons 2 through 8 of the mouse Sept3<sup>−/−</sup> were mapped on a fragment of the 44M13 BAC clone by restriction digest analysis and by DNA sequencing. The long arm of homology from intron 5 to exon 9 was isolated on a 5.5-kb DNA fragment cut with KpnI. The short arm of homology was subcloned into the XhoI restriction site using KpnI-adaptors, and directionality was checked by restriction digest analysis. The subcloning was in the opposite transcriptional orientation to the Neo<sup>+</sup> and thymidine kinase genes. The Sept3 targeting vector was linearized with NotI, and 10 μg of linearized DNA was introduced by electroporation into the mouse embryonic stem (ES) cell line R1. Transfected ES cells were grown on gelatinized plates for several days in selective medium containing G418 and ganciclovir.

Recombination in ES cell clones was confirmed by Southern blot analysis. Proper integration at the 5′ end of the homologous target sequence was determined by mobility shift of an Spht-digested DNA fragment. ES cells undergoing proper homologous recombination were prepared for aggregation. ES cells were grown sparsely on gelatinized plates to produce 8- to 16-cell clumps. These clumps were mixed with embryos at the morula stage (8 cells) isolated from 12 superovulated CD1 females, and both cell populations were allowed to aggregate for 1 day. Chimeric embryos (20 to 30) were reimplanted into pseudopregnant female mice. The chimeric male mice were bred with CD1 female mice to produce the F1 generation. F1 mice heterozygous for the mutation were then bred to get the F2 generation. Germ line transmission of the mutation was verified by Southern blot analysis.

**Genotyping.** PCR analysis was used to routinely genotype mouse tails using primers to regions in the short arm of homology (Ex2A start, GGG CTC CCA GAG AGC AGC AAC AC) and the neomycin gene (Neo, CCC ATG GCC GGC GGC GAT CTG GTC) to yield a 0.74-kb DNA fragment from the recombinant locus. Primers to exons 5 (Ex5, ATT GCC AGC AAA CAA CCT GCC CCT GCT GAC) and exon 6 (Ex6, GCT TCA TGA ACT CAA GAT GCC GTG GTC) were used to yield a 1.4-kb DNA fragment from the wild-type locus.

**Generation of Sept3<sup>−/−</sup> Sept5<sup>−/−</sup> mice.** Compound disruption of both the Sept3 and Sept5 genes was created by interbreeding Sept3<sup>−/−</sup> and Sept5<sup>−/−</sup> mice (47).

**Tissue preparation and Western blot analysis.** Brains from wild-type and Sept3<sup>−/−</sup> mice at various stages of development were homogenized in 20 mM HEPES (pH 7.4), 150 mM NaCl, 320 mM sucrose, and 2 mM EDTA plus protease inhibitors (1 μg/ml leupeptin, 1 μg/ml pepstatin, 20 μg/ml phenylmethylsulfonyl fluoride, and 1 mM EDTA). Aliquots were solubilized in 2% concentrated sodium dodecyl sulfate (SDS) loading buffer (100 mM Tris-HCl [pH 6.8], 4% SDS, 0.2% bromophenol blue, and 20% glycerol) with 5% β-mercaptoethanol when reducing conditions were required. As previously described (20), 20 μg of each sample was electrophoresed through a 10% or 12% SDS-polyacrylamide gel, transferred to polyvinylidene difluoride (PVDF) membrane (Millipore, Billerica, MA), and blocked in 5% milk powder in phosphate-buffered saline (PBS)-Tween. Following primary and secondary antibody incubations, immobilized antibodies were detected with an ECL Western blotting detection reagent (GE Healthcare Biosciences Corp., Piscataway, NJ) and exposure to Kodak film (Rochester, NY).

**Immunoprecipitation.** Mouse brains were flash-frozen with liquid nitrogen, crushed using a mortar and pestle, resuspended in 10 ml HKA buffer (10 mM HEPES-KOH [pH 7.4], 140 mM potassium acetate, 1 mM magnesium chloride and protease inhibitors), and homogenized using a Dounce homogenizer. The homogenized brain tissue was centrifuged at 800 × g for 10 min at 4°C to remove cell debris and nuclei. The resulting supernatant was incubated with 2 μg of each septic antibody in HKA buffer for 1 hour at 4°C and passed through the protein G beads 3 times. Triton X-100 insoluble material was pelleted at 14,000 rpm at 4°C. Protein concentrations were determined by the bicinchoninic assay (Pierce, Rockford, IL), and 2 mg of protein lysate was used for immunoprecipitation. A 10-μl bed volume of protein G beads (Invitrogen Canada Inc, Burlington, ON, Canada) was washed with 1 ml of HKA buffer. Nonspecific binding sites were blocked with 0.1% gelatin and 0.1% bovine serum albumin in HKA buffer for 30 min at room temperature and then incubated with 2 μg of SEPT5 antibody (clone SP20) for 4 h at 4°C. Following immunoprecipitation, the beads were washed three times with 1 ml of HKA buffer containing 1% Triton X-100 and 0.1% gelatin, followed by a wash with the Triton X-100 concentration increased to 2%. Protein complexes were eluted from the beads with 2× SDS loading buffer containing 5% β-mercaptoethanol.

**Imaging mouse brains.** Anesthetized wild-type and Sept3<sup>−/−</sup> mice were transcardially perfused with heparinized saline overvitalized at room temperature. Sections were cut sagittally and processed for hematoxylin and eosin staining by the Pathology Department at The Hospital for Sick Children (Toronto, ON, Canada).
tations of ethanol, samples were embedded in Spurr resin (Canemco Inc., Montreal, QC, Canada). Sections were cut en face, stained with 2.0% uranyl acetate for 15 min and 0.2% lead citrate for 5 min, and then imaged with a Philips CM100 electron microscope.

**Immunostaining hippocampal neurons.** Mouse hippocampal neurons were isolated from hippocampi from 17-day-old fetal mouse brains as previously described (18). After 3 or 10 days in culture, the neurons were fixed with 4% paraformaldehyde in PBS for 20 min, permeabilized with 0.1% Triton X-100 in PBS for 10 min, and blocked with 5% horse serum in PBS for 1 h at room temperature. Primary antibodies were diluted in PBS containing 0.5% horse serum and incubated for 1 h at room temperature or overnight at 4°C. Fluorescence-labeled secondary antibodies were added for 1 h at room temperature and then washed with PBS. Images of fluorescently stained hippocampal neurons mounted onto glass slides with Dako mounting medium (Glostrup, Denmark) were taken with a Zeiss LSM510 Multiphoton laser scanning confocal microscope (Zeiss, Thornwood, NY) and processed with Adobe Photoshop and Adobe Illustrator software. In order to quantify axon outgrowth, digital images of hippocampal neurons were converted to GIF format with Adobe Photoshop software and then analyzed with ImageJ software using the NeuronJ plug-in.

**Synaptic vesicle recycling with FM 1-43.** Primary mouse hippocampal neurons from wild-type or Sept3−/− brains were cultured for 10 days on glass coverslips. The coverslips were secured in an imaging chamber (RC-21BRFS and P-2 Series 20 platform; Warner Instruments, Hamden, CT) and mounted on the stage of a Leica DMIRB microscope (Richard Hill, ON, Canada) using a stage adapter (Series 20 Stage adapter; Warner Instruments, Hamden, CT). Before each experiment, the growth medium was replaced with Tyrode’s buffer (119 mM NaCl, 2.5 mM KCl, 1.25 mM MgCl2, 0.5 mM CaCl2, and 26 mM NaHCO3, pH 7.4). Dye loading (a measure of endocytosis) was done in Tyrode’s buffer containing 47 mM K+ (isotonic replacement of Na+ for K+) and 15 µM FM 1-43 (Invitrogen-Molecular Probes, Burlington, ON, Canada) for 90 s. The cells were washed by perfusion with Ca2+/Mg2+-free Tyrode’s buffer for 10 min. Dye unloading (a measure of exocytosis) was elicited with five 1-min applications of Tyrode’s buffer containing 90 mM K+ (separated by 1 min washes with Ca2+/Mg2+-free Tyrode’s buffer) to ensure the release of the dye-labeled vesicles (an average of three images) were collected on initial dye loading (F0) and after unloading (F∞). The size of the recycling pool was calculated for each synapse as the change in fluorescence intensity: ΔF = F0 − F∞. All FM 1-43 experiments were performed at room temperature. AMPA (α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid) and NMDA (N-methyl-D-aspartate) glutamate receptors were blocked with CNQX (6-cyano-2,3-dihydroxy-7-nitroquinoxaline; 10 µM) and APV (2-amino-5-phosphonovaleric acid; 50 µM), respectively, throughout the course of the experiments to prevent recurrent excitatory and induction of synaptic plasticity. Fluorescence images were taken during a 40× (numerical aperture, 0.75) objective lens by exciting fluorescently labeled neurons at 485 nm using a Chroma fluorescein isothiocyanate filter cube (Rockingham, VT) and collecting the emitted light through a 535-nm emission filter. Light-stimulated calcium influx in intracellular calcium indicators (Fura-2, Fura-red) was measured with a high-field-image using Nomarski optics, was controlled by Metallur software, version 6.3R6 (Universal Imaging Corp., West Chester, PA), through a Labado1-2 shutter controller (Sutter Instruments, Novato, CA). The data were calculated using Microsoft Excel software and plotted using GraphPad Prism, version 4. Statistical measurements between frequency histograms were made using the nonparametric Kolmogorov-Smirnov test (http://www.physics.csbsju.edu/stats/KS-test.html). Measurements between frequency histograms were made using the nonparametric standard error from a population of synapses.

**Hippocampus electrophysiology.** Transverse slices of 3- to 8-week-old mouse hippocampus were cut at a thickness of 250 to 300 µm using a vibrating microtome (Leica VT 1000S, Wetzlar, Germany) in ice-cold cutting solution containing 112 mM NaCl, 2.5 mM KCl, 1.25 mM MgCl2, 2.5 mM CaCl2, 10 mM glucose, 1.25 mM NaH2PO4, 2 mM KCl, 2 mM NaHCO3, 6.7 mM MgCl2, and 0.5 mM CaCl2, at a pH of 7.3 when bubbled with in 95% O2 and 5% CO2. The slices were incubated at 34°C for 30 min in the recording solution followed by an additional 30 min at 34°C in a recording solution containing 125 mM NaCl, 2.5 mM KC1, 10 mM glucose, 1.25 mM NaH2PO4, 2 mM Na-pyruvate, 3 mM m-isonicotin, 0.5 mM ascorbic acid, 26 mM NaHCO3, 1 mM MgCl2, and 2 mM CaCl2 at a pH of 7.3 when bubbled in with 95% O2 and 5% CO2. Subsequently, the temperature of the slices was brought down to room temperature. During whole-cell voltage clamp recordings, slices were superfused with recording solution, and to facilitate EPSC recording 10 µM bicineculline, a γ-aminobutyric acid receptor blocker was included. For synaptic experiments in CA1 pyramidal neurons, the pipette solution contained 97.5 mM NaCl, 33.3 mM glutamic acid (133 mM glutamate), 5 mM MgCl2, and 1 mM CaCl2, at pH 7.3 with bubbled in with 95% O2 and 5% CO2. Synaptic vesicle recycling with FM 1-43.

**Western blot analysis of septin expression from wild-type and mutant mice.** Three brains from P30 wild-type, Sept3−/−, and Sept5−/− mice were individually prepared as described in the Materials and Methods section.
Methods section (see “Tissue preparation and Western blot analysis”). For each gel, 30 μg of protein lysate from each genotype was loaded per lane. Two more sets of gels were created with the other brains (three independent sets). Following electrophoresis and gel transfer, each blot was first probed with a septin antibody (see “Tissue preparation and Western blot analysis”) and then with a glyceraldehyde-3-phosphate dehydrogenase (GAPDH) antibody to control for variability in gel loading. The results on autoradiographic film were digitized with a Dell A920 scanner to facilitate densitometry measurements using an ImageJ software package. Only protein bands that corresponded to known septin isoforms were analyzed. Changes in septin expression in mutant backgrounds were normalized to wild-type levels and are reported as percent expression levels (mutant/wild type × 100%). Similarly, GAPDH protein levels were measured and normalized to the wild-type level. These results were used to calibrate the percent expression level of septin from the corresponding blot. Measurements from all three sets were averaged, and significant differences, indicated by asterisks above the bar graphs, were determined using a Student’s t test. The criterion for statistical significance was chosen to be a P value of <0.05.

Immunostaining vibratome slices. Brain slices (200 μm in thickness) from the auditory cortex of P16 mice were cut and allowed to recover for 1 h in aCSF at 37°C. Subsequently, the brain slices were fixed, permeabilized, blocked, and stained using a Basic Vector Mouse-on-Mouse Immunodetection Kit according to the manufacturer’s instructions (Vector Laboratories, Burlingame, CA). Sept3, Sept5, and DNA were detected using a rabbit anti-SEPT3 antibody, a mouse SP20 antibody, and Hoechst 33342 stain (Invitrogen-Molecular Probes, Burlington, ON, Canada), respectively. See “Immunostaining hippocampal neurons” in Materials and Methods for the acquisition and processing of digital images of fluorescently labeled brain slices.

RESULTS

Generation of Sept3−/− and Sept3−/− Sept5−/− mice. The function of Sept3 was examined by disrupting the Sept3 gene locus in mice. A gene targeting vector was designed to replace a genomic region containing exons 2 to 5, which includes the P-loop consensus motif found in all GTPases (Fig. 1A). Out of the 167 antibiotic-resistant clones, four ES cell clones underwent homologous recombination with the targeting vector (Fig. 1B and C). Proper integration of the short and long arm of homology was confirmed for the ES cell clone 9F1, so it was used to generate chimeric founder mice, and the mutation was bred to homozygosity. Germ line transmission of the mutant allele was initially confirmed by Southern blot analysis (Fig. 1D) and then routinely determined by PCR (Fig. 1E). The absence of Sept3 protein from brains of Sept3−/− mice was confirmed by Western blot analysis (Fig. 1F).

Sept3−/− mice are viable at birth and live as long as their wild-type counterparts. Breeding Sept3−/− mice produced Sept3−/− pups at the expected Mendelian ratio (data not shown). We then determined if Sept3−/− mice were fertile since the lack of Sept4 causes male sterility in mice due to loss of annuli, ring-like structures important for sperm motility, from sperm tails (26, 36). Although no Sept3 protein was detected in testes by Western blot analysis, putative Sept3 transcripts were found in the testes by Northern blot analysis (71, 74), raising the possibility that the protein may have a specialized role in this tissue. However, Sept3−/− male mice are still fertile (data not shown). Sept3−/− female mice were also tested for fertility because the expression of Sept3 in female reproductive organs was not determined previously. Similar to Sept3−/− male mice, Sept3−/− female mice were also fertile (data not shown). These results indicate no primary functional role for Sept3 in germ cells. Sept3−/− mice were subsequently bred with Sept5−/− mice to generate Sept3−/− Sept5−/− mice. Like Sept3−/− mice, Sept3−/− Sept5−/− mice were viable and lived as long as their wild-type counterparts. In addition, both male and female double knockout mice were fertile.

SEPT5 complex in the absence of SEPT3. In Drosophila embryos, Sep2 but not Sep1 failed to accumulate at the cellularization front in the absence of Pnut (1). It was postulated from this observation that septin complex assembly might occur in a hierarchical manner. We therefore determined whether the lack of Sept3 affected the ability of Sept5 to associate with other septins. Sept5 was found to immunoprecipitate with Sept2, Sept4, Sept6, Sept7, Sept8, Sept9, and Sept11 (Fig. 2). Sept1 was not found in this complex but has been observed in another brain septin complex (34). This result does not distinguish between a single complex and multiple complexes containing SEPT5. Nevertheless, the association of other septins with Sept5 was unaffected by the lack of Sept3. Therefore, the absence of Sept3 did not prevent or enhance the interaction of any other septin with Sept5.

Septin expression in the absence of SEPT3 and SEPT5. The lack of one protein can affect the stability of other proteins that are associated with it in a complex. For example, the targeted disruption of amphiphysin I results in complete loss of its partner amphiphysin II by proteolysis (12). Likewise for septin complexes, knockdown of Sept2 by RNA interference causes the level of Sept7 to decrease and vice versa (34). Moreover in Sept5−/− mouse brains, there is a significant change in the expression of Sept2 and Sept7 (47). Therefore, the expression levels of other septins were compared by Western blot analysis between wild-type, Sept3−/−, Sept5−/−, and Sept3−/− Sept5−/− mouse brains at 0, 2, and 4 weeks of age (Fig. 3). Quantitation of septin levels at 4 weeks of age revealed several differences in the mutant backgrounds (Fig. 4). In the absence of Sept3, there was a significant decrease in the expression of Sept2, Sept6, and the largest form of Sept8. In the absence of Sept5, there was a significant increase in the levels of Sept2 and a significant decrease in the levels of Sept7, which is consistent with previous results (47). In addition to Sept7, a significant decrease in the levels of Sept3, Sept6, the largest isoform of Sept8, and Sept11 was also observed. In the absence of both Sept3 and Sept5, there was a significant decrease in the levels of Sept6, Sept7, both forms of Sept8, and Sept11, whereas one isoform of Sept4 increased. Although the expression level of Sept12 was not tested in either Sept3−/− or Sept5−/− mouse brains, its expression level was not detected in the brains of Sept3−/− Sept5−/− mice. This septin is predominantly expressed in the testes and closely related to Sept3 because it lacks a predicted coiled-coil domain. It should be noted that extra bands of unconfirmed identity were observed for Sept1, Sept9, and Sept11. The appearance of these bands may be associated with the mixed genetic background of the single and double knockout mice. Furthermore, the levels of Sept10, Sept13, and Sept14 were not tested in this study due to the lack of specific antibodies for these septins.

Axon outgrowth and synapse formation. Septins are known to associate with the exocyst complex (24), which is a group of proteins implicated in neurite outgrowth and synapse formation. Thus, the roles of Sept3 and Sept5 were examined in the development of isolated neurons. Primary hippocampal neurons were cultured from wild-type and Sept3−/− mouse brains and tested for their ability to polarize, form axons, and make synapses. Primary hippocampal neurons lacking Sept3 exhibited normal growth rates and acquired characteristic
markers of polarity after 3 days in vitro (Fig. 5A). Staining for neurofilament heavy chain and F-actin, which marked emerging axons and growth cones, respectively, revealed normal development of these structures. When the loss of SEPT3 was compounded with the loss of SEPT5, these neurons were still able to polarize, which can be seen by the asymmetric distribution of axonal (neurofilament-H) and dendritic (MAP2) markers in their processes (Fig. 5A). Similarly, no significant

FIG. 1. Gene targeting strategy, ES screening, and genotyping. (A) The mouse Sept3 gene was disrupted by replacing exon 2 to exon 5 with a Neo' selection cassette. The cassette was targeted to the Sept3 locus by appending DNA fragments that were homologous to the Sept3 gene (targeting vector). A large piece of intron 1 and a small piece of exon 2 were used for the short arm of homology, whereas a piece of intron 5 through to the end of intron 8 was used for the large arm of homology. The endogenous and recombinant loci are shown above and below the targeting vector, respectively. For perspective, the location of relevant protein domains in relation to the encoding exons is shown at the top (PRO is the proline-rich domain, G1 is the P-loop motif, and G3 and G4 represent the other GTP binding features). Screening for recombination events was performed by Southern blotting using the 5' and 3' probes or by PCR using the Ex5-Ex6 and Ex2-Neo primer pairs. The sizes of DNA fragments that each probe is expected to detect by Southern blotting or that each primer pair is expected to amplify during PCR are shown on the right. The locations of only the relevant restriction enzyme cut sites are indicated for simplicity. (B) Screening results for proper homologous recombination of the short arm of homology in ES cells. Among the three ES cell clones shown, only clone 9F1 was targeted correctly. (C) Screening results for proper homologous recombination of the long arm of homology in ES cells. Out of six ES cell clones shown, only clones 9F1, 2A2, 11E2, and 12E2 were targeted correctly. (D) Genotype determined by Southern blot analysis of mouse tails from an F2 generation. (E) Genotype determined by PCR analysis of mouse tails from an F2 generation. (F) Western blot analysis of 20 μg of brain tissue from wild-type and Sept3−/− mice using an antibody to SEPT3.
difference in axonal growth rates was observed when neurofila-
ment-H-positive, MAP2-negative processes of stage III (13) wild-type and Sept3−/− Sept5−/− hippocampal neurons were measured after 2 days in vitro (67.7 ± 1.6 μm and 69.2 ± 1.8 μm, respectively [SEM]; P = 0.52; total number of neurons analyzed from three wild-type litters, 357; total number of neurons analyzed from four Sept3−/− Sept5−/− litters, 438).

After 10 days in vitro, primary hippocampal neurons form a large number of synapses onto neighboring dendrites or cell bodies. At this stage, presynaptic terminals are enriched with the synaptic vesicle protein synaptophysin and Sept3 (Fig. 5B). When neurons isolated from Sept3−/− mice were grown for 10 days in culture, they were still able to form synapses (Fig. 5B). The accumulation of synaptophysin-positive rich areas along the processes of these neurons revealed no obvious differences in either the distribution or abundance of synaptic contacts with dendrites or cell bodies. Sept5 is also enriched in presynaptic terminals of hippocampal neurons, where it colocalizes with the synaptic vesicle protein VAMP2 (47). Even without Sept3 and Sept5, hippocampal neurons made synaptic connections with each other after 10 days in vitro (Fig. 5B).

**Hippocampal architecture is normal in the absence of Sept3 and Sept5.** Although axon outgrowth and synapse formation appeared normal in vitro, the correct development of complex brain structures may be more sensitive to the absence of critical proteins such as Sept3 and Sept5. The anatomy of the hippocampus was examined in detail because both Sept3 and Sept5 are expressed in this region of the brain (73) and could be affected by their absence. Gross defects in the hippocampus are easy to identify because the neurons in this area have a distinct topographical organization. Nevertheless, no major abnormalities were detected in the architecture of the hippocampus viewed sagittally from Sept3−/−, Sept5−/−, and Sept3−/− Sept5−/− brains (Fig. 6A).

**Nerve terminal ultrastructure.** Although hematoxylin- and cosin-stained brain sections are suitable for making qualitative assessments about tissue architecture, we turned to transmission electron microscopy, which provides better resolution and contrast to address subcellular structures. Since Sept3 and Sept5 are associated with synaptic vesicles and concentrated in synapses (32, 73), the ultrastructure of nerve terminals from the CA1 region of the hippocampus were compared between wild-type, Sept3−/−, Sept5−/−, and Sept3−/− Sept5−/− mice (Fig. 6B). Presynaptic terminals from mouse hippocampal neurons had no obvious changes in terminal area, synaptic vesicle size, or synaptic vesicle density in the absence of Sept3 or Sept5. Likewise, when the loss of Sept3 was compounded with the loss of Sept5, the morphology of these terminals was similar to that of the wild type.

**Synaptic vesicle protein expression in the absence of Sept3 and Sept5.** Although the architecture of Sept3−/− and Sept3−/− Sept5−/− neurons looks similar to wild-type neurons, there could be differences in their function. Since septins are thought to be important for secretion, we examined the levels of synaptic vesicle fusion proteins in the absence of Sept3 and Sept5 (Fig. 7). In Sept3−/− Sept5−/− brain tissue, the levels of syntaxin 1 and synaptophysin were 68% and 65% of that in wild-type brain tissue, whereas the levels of VAMP2 and SNAP-25 were 98% and 108% of that in wild-type tissue, respectively. These observations are consistent with previous reports showing that Sept5 associates with synaptophysin and syntaxin (4). Loss of Sept5 might disrupt the stability of these two proteins in nerve terminals, thereby impairing the process of synaptic vesicle fusion. Therefore, we turned to FM 1-43 imaging and electrophysiology on single and double knockout neurons to address this question.

**Properties of synaptic transmission in the absence of Sept3 and Sept5.** Genetic mutations of presynaptic proteins can alter certain properties of synaptic transmission in the absence of any obvious morphological defects. In the absence of Sept3, no differences from wild-type mice were observed in several synaptic transmission parameters at the hippocampal CA3-CA1 synapse (47). The parameters used to examine presynaptic physiology were revisited in mice lacking Sept3 and
SEPT5 to determine if a phenotype would manifest with a compound genetic deletion.

Paired-pulse facilitation is a form of short-term plasticity seen at many synapses and is characterized by an increase in neurotransmitter release evoked by a second stimulus during twin-pulse stimulation. When the latency between the first and second pulse is shorter than the time it takes to extrude Ca$^{2+}$ out from the nerve terminal, the residual intracellular Ca$^{2+}$ left over from the first stimulus contributes to a greater Ca$^{2+}$ rise, and hence greater evoked release, during the second stimulus. Thus, changes in paired-pulse facilitation can indicate changes in the sensitivity of the presynaptic release apparatus. Paired-pulse facilitation is taken as the ratio between the postsynaptic response amplitude evoked by the second stimulus and the response amplitude evoked by the first. When the paired-pulse ratio (PRR) was measured with an interpulse interval of 150 ms (Fig. 8A, B, and E), no difference between wild-type and Sept3$^{-/-}$ Sept5$^{-/-}$ mice (Fig. 8C, D, and G), no significant difference was observed between them (38.4 ± 6.1 pA [n = 10 cells] versus 44.0 ± 5.0 pA [n = 10 cells], respectively; P = 0.49). When the frequencies of spontaneous EPSCs were measured from the same mice (Fig. 8C, D, and F), no significant difference was observed (0.44 ± 0.09 Hz for wild-type and 0.47 ± 0.08 Hz for Sept3$^{-/-}$ Sept5$^{-/-}$ mice; n = 10 cells for each genotype; P = 0.84). The analyses of spontaneous EPSCs indicate that in the absence of both SEPT3 and SEPT5, there appears to be no change in the synaptic vesicle release probability or postsynaptic receptor responsiveness.

The synaptic vesicle recycling pool. Primary hippocampal neurons typically contain 100 to 200 synaptic vesicles per terminal. The recycling pool size is about 10 to 20% of the total (22) and is defined as the population of synaptic vesicles that maintain neurotransmitter release during moderate (physiological) stimulation (48). To obtain a quantitative measurement of the functional synaptic vesicle pool size, the fluorescent indicator dye FM 1-43 was used to measure the size of the recycling pool. When added to the bath containing high K$^+$ (dye loading), some FM 1-43 inserted into the plasma membrane is internalized into recycling synaptic vesicles, providing

![FIG. 3. Levels of septin expression in the absence of SEPT3, SEPT5, and both septins. Brains from wild-type, Sept3$^{-/-}$, Sept5$^{-/-}$, and Sept3$^{-/-}$ Sept5$^{-/-}$ mice at P0, P14, and P28 were homogenized and then solubilized in SDS gel sample buffer. In each lane, 20 μg of protein sample was separated by SDS-polyacrylamide gel electrophoresis, transferred onto PVDF membrane, and probed with the indicated septin antibodies. A number of the septins undergo alternative splicing to produce multiple isoforms of different sizes on gels.](http://mcb.asm.org/)
FIG. 4. Quantitation of septin expression in the absence of SEPT3, SEPT5, and both septins. P30 brains from wild-type, Sept3<sup>+/−</sup>, Sept5<sup>+/−</sup>, and Sept3<sup>−/−</sup> Sept5<sup>−/−</sup> mice were homogenized and solubilized in SDS gel sample buffer. In each lane, 30 μg of protein sample was separated by SDS-polyacrylamide gel electrophoresis, transferred onto a PVDF membrane, and probed with the indicated septin antibody followed by the GAPDH antibody. The relative level of each septin after calibration was measured in three independent experiments and averaged. Any significant deviation in septin expression from wild-type levels is indicated by an asterisk.
FIG. 5. Loss of SEPT3 or SEPT3 and SEPT5 does not affect the development of primary hippocampal neurons in vitro. Primary mouse hippocampal neurons from wild-type, Sept3−/−, and Sept3−/− Sept5−/− brains were cultured for 3 days (A) or 10 days (B). Wild-type and Sept3−/− neurons cultured for 3 days were stained for SEPT3, F-actin, and neurofilament-H (green, red, and blue, respectively, in the merged panel).
one index of synaptic vesicle endocytosis (37). Destaining of synaptic vesicles by exocytosis (dye unloading) is triggered by a second K\(^+\) depolarization. Wild-type nerve terminals from 10-day hippocampal neuronal cultures were depolarized by bath application of 47 mM K\(^+\) in the presence of 15 \(\mu\)M FM 1-43 for 90 s (Fig. 9A, steps 1 to 5) to completely label the recycling pool of synaptic vesicles (22). After a 10-min wash in Ca\(^{2+}\)-free buffer, a punctate distribution of dye-loaded nerve terminals was clearly evident (Fig. 9B, middle panel). When this procedure was repeated for Sept3\(^{-/-}\) nerve terminals, the total amount of dye uptake was not different from wild-type terminals (data not shown). This suggests that Sept3 is not essential for synaptic vesicle endocytosis and that its absence does not affect the amount of recycling that occurs in response to high K\(^+\) for 90 s. Next, dye-loaded synaptic vesicles were depolarized with 90 mM K\(^+\) to elicit exocytosis (dye unloading) (Fig. 9A, steps 5 to 7). Following the depolarizing stimulus, wild-type nerve terminals were completely destained (Fig. 9B, right panel), and the same was true for knockout nerve terminals (data not shown). The overall size of the synaptic vesicle recycling pool was quantitatively determined (\(\Delta F = F_{\text{pre}} - F_{\text{post}}\)) for nerve terminals in the presence and absence of Sept3 and displayed as a frequency distribution (Fig. 9C). The distribution of recycling pool sizes was found to be broad but overlapping for both genotypes. Using the data sets from the experiment shown in Fig. 9C, a cumulative frequency curve was constructed for each genotype (Fig. 9D) so that the differences between genotypes could be examined by a Kolmogorov-Smirnov test. This test, which measures the statistical difference between two data sets without making any assumptions about their distributions, confirmed that there was no change in the size of the recycling pool in the absence of Sept3 (\(P = 0.362; n = 532\) terminals from seven wild-type pups and \(n = 555\) terminals from six Sept3\(^{-/-}\) pups). The results demonstrate there was no overall effect on synaptic vesicle fusion and recycling in Sept3\(^{-/-}\) mice.

**SEPT3 and Sept5 are not crucial for the replenishment of the immediately releasable pool of synaptic vesicles.** In order to determine the importance of Sept3 and Sept5 in the replenishment of the readily releasable pool (RRP) of synaptic vesicles, we turned to the calyx of Held nerve terminal which makes a mono-synaptic connection with the MNTB. Like hippocampal boutons, the calyx of Held expresses both Sept3 and Sept5, and these two septins show a strong degree of colocalization throughout the terminal (Fig. 10). The large size of the calyx of Held terminal makes it feasible to do paired whole-cell voltage clamp recordings (Fig. 11A). Furthermore, direct voltage clamping of the presynaptic terminal allows for precise control of the ionic currents triggering release of synaptic vesicles, which can be measured from the current response of the postsynaptic MNTB neuron (6, 18). These characteristics make the calyx of Held-MNTB synapse a unique model in which to examine aspects of synaptic transmission and dissect the components of the RRP.

The pool of synaptic vesicles can be subdivided into two main components: those that are releasable under conditions of typical activity (RRP) and those that refill the RRP, or the reserve pool (44). The RRP can be further subdivided into an immediately releasable pool (IM-RRP) and an intermediate pool (IT-RRP). The IM-RRP pool consists of those synaptic vesicles that are available for release in response to the Ca\(^{2+}\) current (\(I_{\text{Ca}}\)) evoked from a single presynaptic action potential and are most likely those synaptic vesicles in close proximity to voltage-gated Ca\(^{2+}\) channels (VGCCs) and active zones. The IT-RRP pool consists of synaptic vesicles that are recruited into release during sustained elevations in intraternal [Ca\(^{2+}\)], which occurs during repetitive high-frequency activity or sustained presynaptic influxes of Ca\(^{2+}\) (51, 53).

To assay whether loss of Sept3 and Sept5 impacted the ability of the calyx of Held to replenish the IM-RRP, we depolarized voltage-clamped presynaptic terminals from –80 mV to 40 mV for 4 ms, which triggered a large inward Ca\(^{2+}\) current through VGCCs (Fig. 11B and C). During the step, a majority of VGCCs enter an open state; however, they do not flux Ca\(^{2+}\) ions as 40 mV is past the effective reversal potential for Ca\(^{2+}\). Upon repolarization, a steep electrochemical gradient drives Ca\(^{2+}\) ions into the terminal, during the brief period before VGCCs close, producing a large tail current (Fig. 11C). The Ca\(^{2+}\) flux through individual VGCCs creates [Ca\(^{2+}\)] domains which are analogous to those resulting from the inervation of a single action potential into the nerve terminal (7). However, under these conditions, all VGCCs contribute to the \(I_{\text{Ca}}\) rather than the relatively small fraction activated during an action potential, especially at physiological temperatures (75A). The resulting synaptic output contains all of the synaptic vesicles that would be available during any single action potential, or the IM-RRP.

By separating two such depolarizations by an interstep interval (\(\Delta t\)) of variable duration, we determined the portion of the IM-RRP that recovered for a range of \(\Delta t\) (5 ms to 10 s) in both wild-type and Sept3\(^{-/-}\) Sept5\(^{-/-}\) synapses (Fig. 11D). This protocol failed to completely deplete the IM-RRP even at the shortest \(\Delta t\) possible (5 ms). At this short time interval, recovery was 54% ± 4% and 40% ± 8% for wild-type and Sept3\(^{-/-}\) Sept5\(^{-/-}\) synapses, respectively, which were not significantly different (unpaired \(t\) test, \(P > 0.05\)). The recovered portion of the IM-RRP continued to increase with increasing \(\Delta t\) and followed a biphasic time course. Fast recovery time constants were 39 ± 13 ms and 43 ± 7 ms for wild-type and Sept3\(^{-/-}\) Sept5\(^{-/-}\) synapses, respectively, while slow time constant neurons lacking Sept3, processes containing the axon-specific protein neurofilament-H still form growth cones at their tips (white arrow). Sept3\(^{-/-}\) Sept5\(^{-/-}\) neurons cultured for 3 days were stained in the top series of panels for Sept3, F-actin, and neurofilament-H (green, red, and blue, respectively, in the merged panel) or in the bottom series of panels for Sept5, F-actin, and MAP2 (green, red, and blue, respectively, in the merged panel). Like Sept3\(^{-/-}\) neurons, F-actin-rich growth cones at the tips of axons (arrow) are still present in the absence of Sept3 and Sept5. Wild-type and Sept3\(^{-/-}\) neurons cultured for 10 days were stained for Sept3 and synaptophysin. Sept3\(^{-/-}\) Sept5\(^{-/-}\) neurons cultured for 10 days were stained in the left series of panels for Sept3 and synaptophysin or in the right series of panels for Sept5 and VAMP2. A digitally magnified image of the region highlighted by the hatched box is shown directly below. The arrows point to presynaptic terminals enriched with synaptophysin. In neurons lacking Sept3 or Sept3 and Sept5, many punctate regions enriched with synaptophysin or VAMP2 still form (white arrows).
stants were 2,500 ± 600 ms and 2,400 ± 600 ms, respectively. Given that no significant differences were observed between wild-type and Sept3−/− Sept5−/− synapses in the rates of recovery of their IM-RRP (Fig. 11C and D), we concluded that SEPT3 and SEPT5 are not essential for the trafficking of synaptic vesicles to the active zone and their positioning into the immediately available pool.

**SEPT3 and SEPT5 are not crucial for the replenishment of the readily releasable pool of synaptic vesicles.** Although SEPT3 and SEPT5 are not essential for replenishment of the IM-RRP, they may be involved in trafficking synaptic vesicles from the reserve pool of synaptic vesicles to the broader pool of available synaptic vesicles recruited primarily during repetitive activity (IT-RRP and IM-RRP) (67). To investigate whether SEPT3 and SEPT5 are involved in the replenishment of the RRP, we depolarized voltage-clamped presynaptic calyces from −80 mV to 0 mV for 10 ms (Fig. 12A). At 0 mV, the maximal current flux occurs through P/Q-type VGCCs, which are the exclusive type mediating release of synaptic vesicles at this developmental stage in the calyx of Held (28). During

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**FIG. 6.** Hippocampal anatomy in wild-type, Sept3+/−, Sept5+/−, and Sept3−/− Sept5−/− brains. (A) A series of magnified images of the hippocampus from wild-type, Sept3+/−, Sept5+/−, and Sept3−/− Sept5−/− mice. Pictures were taken from sagittal sections of formalin-fixed, P30 brains that were stained with hematoxylin and eosin. The bubbles and large spaces around each hippocampus were a result of artifacts that were introduced during tissue preparation. (B) A series of electron transmission micrographs that were taken of nerve terminals from the CA1 region of the hippocampus (boxed area illustrated in panel A). Two representative examples from each genotype are shown.
FIG. 7. Expression of synaptic vesicle proteins in the absence of SEPT3 and SEPT5. Brains from wild-type and Sept3−/− Sept5−/− mice at P0, P14, and P28 were homogenized and then solubilized in SDS gel sample buffer. In each lane, 20 μg of protein sample was separated by SDS-polyacrylamide gel electrophoresis, transferred onto PVDF membrane, and probed with the indicated antibodies.

to the hippocampus (20). Furthermore, this group also verified that SEPT3 does not play a significant role in neuronal development and synapse formation. Consistent with the lack of gross anatomical changes in presynaptic nerve terminals, no changes in the physiology of synaptic vesicle pool size, synaptic vesicle fusion, and synaptic vesicle recycling were observed in the absence of SEPT3.

SEPT5 is another septin that is predominantly expressed in the nervous system, but it is also expressed in blood cells (75). Like SEPT3, generation of Sept5−/− mice revealed that lack of SEPT5 was not essential for mouse development, fertility, or viability, and no irregularities were found in synaptic transmission. However, secretion of large, dense core granules from platelets was enhanced (11), supporting the idea that SEPT5 is a negative regulator of exocytosis (4). The discrepancy in secretion between nerve terminals and platelets lacking SEPT5 was thought to be due to functional redundancy because SEPT3 is not expressed in platelets (data not shown). Therefore, we tested the hypothesis that either SEPT3 or SEPT5 is sufficient for synaptic transmission by creating Sept3−/− Sept5−/− mice. Once again, though, loss of both septins did not produce any changes beyond what was observed for either single knockout mouse line, ruling out the possibility that SEPT3 and SEPT5 functionally compensate for the loss of the other.

Mouse knockouts for two other septins, Sept4−/− and Sept6−/−, were originally reported not to have severe neurological impairments (26, 36, 45). Targeted deletion of Sept4 resulted in male mice that were sterile, due to a structural defect in the sperm tail annulus which impaired sperm motility. In the brains of Sept4−/− mice, no gross structural abnormalities were detected, but a reduction in the number of Purkinje cells in the cerebellum was observed with no obvious motor deficiencies. Sept6−/− mice are fertile, and they do not have any gross structural abnormalities in the cerebellum or impaired neurological function, as assessed by motor performance on a rotor rod task (45). Generation of compound Sept4−/− Sept6−/− mice did not produce defects beyond those already observed in Sept4−/− and Sept6−/− mice (45). In an exciting new finding, it was shown that Sept4−/− mice exhibit reduced dopaminergic synaptic transmission and are susceptible to the aggregation of α-synuclein (27), which are hallmark symptoms associated with Parkinson’s disease.

In addition to mouse knockouts, knockdown of SEPT7 was examined in hippocampal neurons by two independent groups (61, 70). Both groups found that lowering SEPT7 protein levels resulted in neurons with impaired dendritic branching and increased density of elongated spines/immature protrusions. This phenotype is consistent with the localization of SEPT7 to dendritic branch points and the base of filopodia. In Sept3−/− Sept5−/− mice, the level of SEPT7 is significantly reduced from wild-type levels. However, the extent to which this reduction has an effect on dendrite morphology is likely to be minimal since synaptic transmission in the hippocampus of Sept3−/− Sept5−/− mice appears to be normal. A greater than 55% reduction in SEPT7 levels may be necessary to see a phenotype in mice.

Several reasons might potentially explain the absence of an observable phenotype in Sept3−/−, Sept5−/−, or Sept3−/− Sept5−/− mice. One possibility is that another related septin
FIG. 8. Loss of SEPT3 does not affect paired-pulse facilitation or the amplitude and frequency of spontaneous EPSCs in hippocampal CA1 pyramidal neurons. (A) The top trace represents superimposed AMPA receptor-mediated EPSCs from a wild-type mouse evoked by paired stimuli (150 ms apart). The bottom trace represents an average of 15 consecutive traces. (B) The top trace represents superimposed AMPA receptor-mediated EPSCs from a Sept3−/− Sept5−/− mouse evoked by paired stimuli (150 ms apart). The bottom trace represents an average of 15 consecutive traces. (C) A representative 4-s segment of spontaneous EPSCs recorded from a wild-type mouse. A blow-up of the postsynaptic currents is shown in the inset (inset scale, 10 pA/6 ms). (D) A representative 4-s segment of spontaneous EPSCs recorded from a Sept3−/− Sept5−/− mouse. A blow-up of the postsynaptic currents is shown in the inset (inset scale, 10 pA/6 ms). (E) Pooled data showing the PPR of EPSCs for wild-type and Sept3−/− Sept5−/− mice. (F) Pooled data showing frequency of spontaneous EPSCs for wild-type and Sept3−/− Sept5−/− mice. (G) Pooled data showing average amplitude for spontaneous EPSCs for wild-type and Sept3−/− Sept5−/− mice. Error bars in panels E, F, and G show SEMs.
from a similar class is compensating for the loss of these septins during development or within nerve terminals. SEPT3, SEPT9, and SEPT12 belong to a group of septins that lack a predicted coiled-coil domain in their C-terminal extension (65). However, the levels of three SEPT9 isoforms did not change dramatically in the brains of Sept3⁻/⁻ or Sept3⁻/⁻ Sept5⁻/⁻ mice. This result does not exclude the possibility of significant changes in small, isolated regions of the brain where SEPT3 function is most important. The other member of this class, SEPT12, appears to be specifically expressed in the testes (58) and does not get upregulated in brain tissue in the absence of SEPT3 and SEPT5. In contrast to SEPT3, SEPT5 belongs to a separate class with three other septins (SEPT1, SEPT2, and SEPT4) based on sequence similarity. In the absence of SEPT5, the levels of SEPT1 appeared unaltered whereas SEPT2 levels appeared to be upregulated. SEPT2 is required for septin filament assembly in vitro and depletion of this septin in cell lines leads to an increased incidence of binucleated cells (35, 54). However, immunofluorescence analysis of SEPT2 in primary rat hippocampal neurons isolated from Sept5⁻/⁻ mice did not reveal any upregulation of SEPT2 in these neurons (data not shown). Once again, the changes in SEPT2 levels could be occurring in small, isolated regions of the brain where SEPT5 function is most important. However, it remains to be demonstrated whether septins in the same class are functionally interchangeable. Surprisingly, in the absence of both SEPT3 and SEPT5, very little upregulation was observed in the expression of the other septins, with the exception of one isoform of SEPT4. Instead, the protein levels of several septins were significantly reduced. However, it has not
been clearly demonstrated whether our antibodies can detect all septin isoforms in the brain. Furthermore, the levels of SEPT10, SEPT13, and SEPT14 remain to be examined.

A second explanation for the absence of an observable phenotype in Sept3−/−, Sept5−/−, or Sept3−/− Sept5−/− mice is that functional redundancy could occur at the level of more distantly related septins. Immunoprecipitation of SEPT3 or SEPT5 from brain lysates pulls down many other septins. Since the lack of SEPT3 had little impact on the composition of septin complexes isolated from neurons, the overall function of septin complexes may not be impaired. Functionally redundant septins are present in budding yeast and fruit flies. For example, S. cerevisiae lacking Shs1/Sep7 grow as well as the wild type, suggesting that Cdc3, Cdc10, Cdc11, and Cdc12 are sufficient on their own for cell division (64). Even a complex of Cdc3, Cdc11, and Cdc12, without Cdc10, is sufficient for growth at suboptimal growth temperatures under nutrient-rich conditions (19). In Drosophila, depletion of the septin peanut did not inhibit cytokinesis during embryogenesis until the pupal stage even without maternal contribution of peanut protein (1). The absence of peanut and Sep1 did not result in gross abnormalities in the developing fly central nervous system (14). It is thought that the continued presence of Sep2 is sufficient to allow embryonic cell divisions and neuronal development to proceed normally in the absence of peanut and Sep1 (1).
Interestingly, in *C. elegans* and *D. melanogaster*, there are no septin family members like SEPT3 that lack a predicted C-terminal coiled-coil domain. This suggests that these septins may be involved in highly specialized functions that are not apparent.

In the absence of certain septins, septin filaments may be structurally unstable but still retain sufficient function during neuronal development and synaptic transmission to produce a normal phenotype. While the lack of Cdc10 does not impair cytokinesis at suboptimal growth temperatures, it does affect the position of the remaining septins at the mother-bud neck junction (19). Instead of accumulating evenly at this junction, septins accumulate more on the bud side, as judged by immunofluorescence microscopy. Electron microscope images of budding yeast lacking Cdc10 show no discernible filaments where wild-type septin filaments are normally observed. When purified *S. cerevisiae* Cdc3, Cdc11, and Cdc12 are combined in vitro, they make filaments without Cdc10; however, the Cdc3/Cdc11/Cdc12 (Cdc3/11/12) filaments are short, unstable, and curved and do not make lateral associations (64). This contrasts with the wild-type filaments which are long, rigid, straight, and invariably paired. The mammalian SEPT2/6/7 complex is presumed to be the functional equivalent of the budding yeast Cdc3/11/12 complex. The SEPT2/6/7 complex makes filaments in vitro that are 10 nm in diameter but are curved and unstable like Cdc3/11/12 filaments (34, 54). This suggests the possibility that mammalian septin filaments lacking SEPT3, like yeast septin filaments lacking Cdc10, may retain enough structural integrity to sufficiently function in neurons. Although yeast lacking Cdc11 exhibits more severe growth defects than yeast lacking Cdc10, these mutants are still capable of cell division at 22°C. In addition, the remaining septins are able to form filaments, albeit they are structurally similar to the Cdc10-deficient filaments. Therefore, septin filaments lacking SEPT5 may retain sufficient integrity to function properly in nerve terminals. However, it remains to be demonstrated whether Cdc10 and SEPT3 or Cdc11 and SEPT5 share homologous functions.

It is possible that that septins may not be required for neural development, synapse formation, maturation, or synaptic transmission. Therefore, SEPT3 and SEPT5 may play entirely unrelated roles or else their roles in these functions are regulatory rather than obligatory. In *Schizosaccharomyces pombe*, deletion of all mitotic septins only delays cell division, resulting in a chain-of-cells phenotype (5, 62). In this case it was suggested that septins may function to enhance the efficiency of cytokinesis. Similarly, SEPT3 and SEPT5 in nerve terminals may function to enhance the efficiency of synaptic vesicle fusion or recycling. Such a role would not be detectable as an overall block or enhancement of synaptic vesicle exocytosis or endocytosis unless the appropriate stimulation or conditions were found.

Although the molecular details of septin function in the brain remain elusive, the neural importance of septins is reflected in their association with many neurological disorders. Moreover, the association of septins with the synaptic vesicle trafficking machinery suggests that they are likely to be involved in neuronal growth and synaptic transmission in some way that is yet to be defined. Although the lack of an individual septin such as SEPT3, SEPT4, SEPT5, or SEPT6 does not seem to have a large impact on synaptic transmission, the generation of combination knockouts along with detailed electrophysiological measurements to dissect out septin function in intact brain circuits promises to be an exciting platform for future studies of septin function.

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