Disruption of CCTβ2 Expression Leads to Gonadal Dysfunction

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There are two mammalian genes that encode isoforms of CTP:phosphocholine cytidylyltransferase (CCT), a key rate-controlling step in membrane phospholipid biogenesis. Quantitative determination of the CCT transcripts reveals that CCTα is ubiquitously expressed and is found at the highest levels in the testis and lung, with lower levels in the liver and ovary. CCTβ2 is a very minor isoform in most tissues but is significantly expressed in the brain, lung, and gonads. CCTβ3 is the third isoform recently discovered in mice and is expressed in the same tissues as CCTβ2, with its highest level in testes. We investigated the role(s) of CCTβ2 by generating knockout mice. The brains and lungs of mice lacking CCTβ2 expression did not exhibit any overt defects. On the other hand, a large percentage of the CCTβ2−/− females were sterile and their ovaries exhibited defective ovarian follicle development. The proportion of female CCTβ2−/− mice with defective ovaries increased as the animals aged. The rare litters born from CCTβ2−/− × CCTβ2−/− matings had the normal number of pups. The abnormal ovarian histopathology was characterized by disorganization of the tissue in young adult mice and absence of follicles and ova in older mice, along with interstitial stromal cell hyperplasia which culminated in the emergence of tubulostromial ovarian tumors by 16 months of age. Grossly defective CCTβ2−/− ovaries were associated with high follicle-stimulating (FSH) and luteinizing (LH) hormone levels. Male CCTβ2−/− mice exhibited progressive multifocal testicular degeneration and reduced fertility but had normal FSH and LH levels. Thus, the most notable phenotype of CCTβ2 knockout mice was gonad degeneration and reproductive deficiency. The results indicate that although CCTβ2 is expressed at very low levels compared to the α-isofrom, loss of CCTβ2 expression causes a breakdown in the gonadal response to hormonal stimulation.

Phosphatidylcholine (PtdCho) is a major component of biological membranes in higher eukaryotes and is also secreted by specialized tissues for important extracellular tasks. CTP:phosphocholine cytidylyltransferase (CCT) is a key rate-controlling step in the major biosynthetic pathway leading to PtdCho in most tissues (for reviews, see references 15, 17, and 36). In mammals there are two genes, Pcyt1a (formerly Cptct), located on murine chromosome 16, and Pcyt1b, located on the X chromosome, that encode proteins termed CCTα and CCTβ, respectively (35, 38). The two genes exhibit tissue-specific expression, with CCTα predominating in most tissues and CCTβ being most abundant in brain tissue (32). Two transcripts arise from the Pcyt1a gene that encode the identical CCTα protein. Alternate splicing of the X-linked Pcyt1b gene directs the synthesis of two mRNAs that encode the CCTβ2 and CCTβ3 isoforms in mice (32, 35, 38). CCTβ3 is 28 residues shorter at the amino terminus than CCTβ2 due to transcript initiation at an alternate first exon (32). In humans, intron retention gives rise to a CCTβ1 transcript found in the expressed sequence tag database that is predicted to encode a smaller protein that lacks the carboxy-terminal domain (35).

However, the β1 isofrom is not detected in mice (32), and there is no evidence that the CCTβ1 protein is expressed at significant levels in human tissues.

The CCT proteins are divided into four functional domains. The CCTα amino-terminal region contains a cluster of positively charged amino acids that specify nuclear localization (57–59), whereas the CCTβ proteins lack this signature sequence (32, 36). Thus, CCTα is predominantly a nuclear protein in most cell types (58), with the exception of lung cells (48), and a smaller amount of extranuclear CCTα is found associated with the endoplasmic reticulum (35). In contrast, CCTβ proteins are localized to the endoplasmic reticulum compartment and are absent from the nucleus (35). Both proteins have almost identical catalytic and regulatory helical domains, and accordingly the enzymatic activities of the α and β isofroms depend on their interaction with lipid regulators (35). The catalytic core (residues 72 to 233) is highly conserved among all cytidlyltransferases, from bacteria to mammals (47, 51). A phospholipid sensor domain, or helical domain (residues 256 to 288), is unique to the mammalian proteins and consists of three consecutive 11-residue repeats that form an amphipathic α-helix (30). The helical domain regulates the protein’s reversible association with biological membranes (2, 18, 20, 30, 31) together with its activity (56, 64). The curvature elastic stress hypothesis (4) provides a theoretical framework for understanding both the positive and negative functions of specific membrane lipids in CCT regulation via the helical domain (4, 19). Both the α and β proteins have carboxy-
terminal domains with multiple phosphorylation sites (35). Increased phosphorylation attenuates the activation of CCTα by lipid mediators (3, 65), phosphorylated CCTα is often found dissociated from the cell membrane (55), and the carboxy-terminal domain itself also imparts regulation of enzymatic activity by anionic lipids (37). The function of the CCTβ2 carboxy terminus in controlling activity has not been investigated, but the amino acid sequences of the carboxy-terminal domains of the mouse α, β2, and β3 isoforms are 49% identical (57% similar), suggesting that the proteins are regulated similarly by anionic lipids and phosphorylation. Truncation of both the helical and carboxy-terminal domains gives rise to a dys-regulated CCTα that has higher basal activity in the absence of lipid and is refractory to lipid stimulation (for a review, see reference 36).

The biochemical data thus far has not distinguished a unique, nonredundant role for the CCTβ isoforms in cell or tissue function. A few clues about the regulation of CCTα have emerged from studies with cell culture models, but CCTβ proteins were not detected in these systems. Redistribution of CCTα from a predominantly nuclear location to extranuclear sites is associated with growth factor stimulation of PtdCho synthesis (45). Transient elevation of CCT activity in response to apoptosis is accompanied by increased CCTα availability in the extranuclear compartment (34). While these two studies suggest that CCTα outside the nucleus supports a quantitative increase in or replacement of bulk cellular PtdCho, CCTα within the nucleus is also thought to be functional, because the nuclear phospholipids contains unique PtdCho molecular species (28). Overexpression of CCTβ2 or CCTβ3 protein complement a mutant cell line with conditionally defective endogenous CCT activity (32), as does overexpression of CCTα or even catalytically compromised CCTα mutant proteins (37). Acceleration of PtdCho synthesis by the overexpression of CCTα2 is dissociated from the cell membrane (55), and the carboxy-terminal domain itself also imparts regulation of enzymatic activity by anionic lipids (37). The function of the CCT terminal domain itself also imparts regulation of enzymatic activity by anionic lipids (37). The function of the CCT terminal domain itself also imparts regulation of enzymatic activity by anionic lipids (37). The function of the CCT terminal domain itself also imparts regulation of enzymatic activity by anionic lipids (37).

In conclusion, the Neo resistance gene cassette was inserted into exon 2 of the CCTβ gene and a portion of the 3′ coding sequence from exon 2 was deleted. A negative selection cassette encoding the diphtheria toxin antigen (DTA) was added at the 3′ end of the mouse genomic DNA insert. To accomplish this addition, synthetic oligonucleotides were used to construct a new multiple cloning site that was inserted in pBlue-script KS (+) to yield pMAK6. Next the 1,459-bp HindIII-NcoI fragment of CCTβ2 DNA was subcloned into pMAK6, followed by the 6-kb PSI-BamHI gene fragment. This removed 35 bp of coding sequence from exon 2 and 122 bp of the downstream intron. The 1.2-kb DTA selection cassette (a gift from James Ihle, Department of Biochemistry, St. Jude Children’s Research Hospital) was inserted by using XhoI and Clal. Finally, the 1.2-kb XbaI-EcoRV Neo resistance gene cassette was inserted. The DNA sequence of the plasmid was verified, and the insert is depicted in Fig. 1A.

Generation and identification of gene-targeted ES cell clones. Kml was used to linearize the replacement vector prior to transfection into the 129/SvJ-l+ /-Tyr-cMgfSl-J/+ substrain of embryonic stem cells, W9.5 (gift from Peter McKinnon, Department of Genetics, St. Jude Children’s Research Hospital), grown on mitotically inactivated mouse embryonic fibroblasts which carry resistance to neomycin. More than 200 clones resistant to G418 were selected and screened by Southern blot analysis by using a 207-bp PstI-HindIII probe corresponding to the 5′ end of exon 2 outside the targeted portion (Fig. 1A). Genomic DNA (10 μg) from individual embryonic stem (ES) cell clones was digested with BamHI and separated on a 0.8% agarose gel. After transfer to Nylon membrane (Hybond-N+; Amersham Pharma Biotech), hybridization was performed by using Quickhyb solution (Stratagene) containing 1 × 106 to 2 × 106 cpm of 32P-labeled probe/ml at 42°C overnight. After being washed the blot was exposed to a phosphor screen for 1 to 4 days and was scanned. ES cells that had undergone homologous recombination on the X chromosome with the replacement vector were identified by a 2.8-kb band instead of the 9.0-kb wild-type band (Fig. 1B).

Cell culture. Rat pheochromocytoma (PC12) cells were obtained from the American Type Culture Collection (ATCC) and were maintained in Ham’s F12 medium (ATCC) with 15% horse serum (Atlanta Biologicals, Inc.), 2.5% fatty bovine serum (Atlanta Biologicals, Inc.), 10 mM HEPES (Invitrogen), 2 mM glutamine (Invitrogen), and 5% CO2 at 37°C. For experiments with nerve growth factor (NGF; Invitrogen), cells were harvested without trypsinization and were plated on 100-mm-diameter collagen-coated petri dishes (Becton Dickinson) at a density of 20%, followed by incubation for 24 h. NGF (100 ng/ml) was then added to the indicated cultures and also when the medium was changed 3 days later. After 5 days of incubation with NGF, both the untreated control dishes and the NGF-treated cultures were confluent and cells were washed with phosphate-buffered saline (PBS), lysed with TRIZOL reagent (Invitrogen) for RNA isolation, scraped into PBS without calcium or magnesium (Invitrogen) for protein determination (10) by using rabbit immunoglobulin as a standard, or trypsinized for cell counting by using a hemocytometer.

Generation and identification of CCTβ2-disrupted mice. Cells from a 129/SvJ-l+ /-Tyr-cMgfSl-J/+ (W9.5) ES cell colony containing the recombined Pestyb DNA at the correct locus were injected into C57BL/6J blastocysts, which were then implanted into pseudopregnant female mice by the St. Jude Transgenic Core Facility. Male offspring with 75 to 90% agouti color, the coat color contributed by the ES cells, were bred with C57BL/6J females. Pups that were 100% agouti, indicating germ line transmission, were screened. Tail clips from weanling male offspring were digested overnight at 55°C in buffer containing 0.25 mg of proteinase K/ml, 0.2 M NaCl, 100 mM Tris-HCl (pH 8.5), 5 mM EDTA, and 0.2% SDS. The DNA was extracted by using an equal volume of phenol-chloroform/isoamyl alcohol, and the upper phase was precipitated with 0.8 volumes of cold isopropanol. The DNA precipitate was washed with 80% ethanol and was dissolved in 50 to 100 μl of water at 50°C for 30 min. The DNA was then subjected to

**MATERIALS AND METHODS**

Isolation of murine CCTβ genomic clones. A mouse genomic 129/SvJ library in λEML3 (gift from Gerard Grosfeld, Department of Genetics, St. Jude Children’s Research Hospital) was screened with the d[32P]CTP-labeled human 0.9-kb (BamHI-EcoRI) fragment of the CCTβ2 cDNA from plasmid pAL2 (35). Plaque hybridization was performed for 14 h at 42°C in a solution of 25% formamide, 0.75 M NaCl, 75 mM sodium citrate, 2 × Denhardt’s solution (0.02% Ficoll, 0.02% polyvinylpyrrolidone, and 0.02% sodium phosphate buffer (pH 6.5), 0.1% sodium dodecyl sulfate (SDS), 100 μg of salmon sperm DNA/ml, 10% dextran sulfate. The nitrocellulose filters (Schleicher & Schuell, Dassel, Germany) were washed at 55°C for 1 h in a solution of 0.1% SSC (1 × SSC is 0.15 M NaCl plus 0.015 M sodium citrate) and 0.1% SDS and were exposed to film. After secondary and tertiary screening, a positive plaque was isolated by using the OLEGAN Lambda kit, the DNA was digested with BamHI, and a 9.0-kb fragment was subcloned into the vector pBluescript KS (+). The sequence of the mouse CCTβ2 fragment was determined and found to contain the entire exon 2 in the murine Pestyb gene structure (32) plus 857 bp of upstream and 5,948 bp of downstream intronic sequences.

Construction of the CCTβ2 replacement vector. In general, the Neo resistance gene cassette was inserted into exon 2 of the CCTβ gene and a portion of the 3′ coding sequence from exon 2 was deleted. A negative selection cassette encoding the diphtheria toxin antigen (DTA) was added at the 3′ end of the mouse genomic DNA insert. To accomplish this addition, synthetic oligonucleotides were used to construct a new multiple cloning site that was inserted in pBlue-script KS (+) to yield pMAK6. Next the 1,459-bp HindIII-NcoI fragment of CCTβ2 DNA was subcloned into pMAK6, followed by the 6-kb PSI-BamHI gene fragment. This removed 35 bp of coding sequence from exon 2 and 122 bp of the downstream intron. The 1.2-kb DTA selection cassette (a gift from James Ihle, Department of Biochemistry, St. Jude Children’s Research Hospital) was inserted by using XhoI and Clal. Finally, the 1.2-kb XbaI-EcoRV Neo resistance gene cassette was inserted. The DNA sequence of the plasmid was verified, and the insert is depicted in Fig. 1A.
Breeding and superovulation of mice. The significant drop in fertility found upon breeding the CCTβ2 homozygous females with the CCTβ2-deficient males made it necessary to mate the heterozygous females with the knockout males to maintain a breeding colony. Colony littermates were used as female heterozygotes and male wild-type controls, and commercial wild-type females and age-matched control mice (129/Sv × C57BL/6J) were purchased from Jackson Laboratory (Bar Harbor, Maine). For ovary maturation and ovulation of immature females, CCTβ2 homozygous and wild-type controls were given intraperitoneal injections of 2.5 U of pregnant mare’s serum (Sigma) as a source of follicle-stimulating hormones (FSH) followed by 2.5 U of human chorionic gonadotropin (Sigma) 48 h later (late afternoon) as a source of luteinizing hormone (LH). Females were immediately placed with males and checked for vaginal plugs, indicative of mating, the next morning. Mice were maintained on regular Purina rodina chow #5013 at a room temperature of 72°F ± 2°F, room humidity of 50% ± 10%, and a 12-h light, 12-h dark cycle, with the dark cycle starting at 1800 h. All procedures concerning the care and use of animals were done according to St. Jude Children’s Research Hospital Institutional Animal Care and Use Committee approved protocols.

Detection of CCT isoform mRNAs by RT-PCR. Total RNA was isolated from mouse tissues by using TRIZOL reagent according to the manufacturer’s instructions. Pelleted RNA was resuspended in nuclease-free water, digested with DNase I to remove any contaminating genomic DNA, and aliquoted and reprecipitated with ethanol, and stored at −20°C. Reverse transcription (RT) was done by using SuperScript II RNase H− Reverse Transcriptionase (Invitrogen) and the RNA template and random primers to make the corresponding cDNAs. For conventional RT-PCR, multiplex PCR amplification and detection by visual inspection of the CCT cDNAs was done to measure the relative levels of expression by using the ABI Prism 7700 Sequence Detection System using primers and probes listed in Table 1. The Taqman Rodent GAPDH Control Reagent (Applied Biosystems) was the source of the primers and probe for quantitating the control GAPDH mRNA. RNA was isolated from liver, brain, lung, ovary, and testis of at least three mice of each gender individually, and each RNA sample was quantitated in quintuplicate and, by using the comparative Ct method, the amount of target CCT (Ct) and was related to the amount of target CCTα in liver (ΔCtC), which was set as the calibrator at 1.0. Standard deviations from the mean ΔCt values (not shown) were <10%. Tissues from both female (stippled bars) and male (cross-hatched bars) mice were analyzed. (E) Distribution of CCT and PEMT transcripts was calculated for individual organs by using data obtained from real-time PCR quantification (shown in panels A to D).

Table 1. Real-time PCR reagents

| Primer  | Sequence                              | Probe                   |
|---------|---------------------------------------|-------------------------|
| mCCTα-F | 5'-TGATGCAAGAAGGTTTACGCTA          | 6FM-AATTGTGCTTGTGCACGAT-TAMRA |
| mCCTα-R | 5'-TGTCCTCATTAGGGCCAGGT            | 6FM-TGCTCTTGGACATGCTTACACGAC-TAMRA |
| mCCTβ2-F | 5’-TTCTTGTGCTGGAGGAGAGCT          | 6FM-AATTGTGCTTGTGCACGAT-TAMRA |
| mCCTβ2-R | 5’-AAGTACTGCTGGCAGGCTGTA            | 6FM-TGCTCTTGGACATGCTTACACGAC-TAMRA |
| mCCTβ3-F | 5’-GCGGCACACCGCGCGGACT            | 6FM-AATTGTGCTTGTGCACGAT-TAMRA |
| mCCTβ3-R | 5’-GAGATGCAACTGCTGGACACGC          | 6FM-AATTGTGCTTGTGCACGAT-TAMRA |
| mPEMT-F | 5’-GGCATCTTGACTGGCAGA            | 6FM-GGCTGCACTGGACATGCTTACACGAC-TAMRA |
| mPEMT-R | 5’-TGAGGCTTGACATGCTTACACGAC        | 6FM-GGCTGCACTGGACATGCTTACACGAC-TAMRA |

Southern analysis by using the exon 2 outside probe as described above. Heterozygous (+/−) females and homozygous (−/−) males were identified and separated into homozygous (+/−) females as identified by PCR with FPr1 (5' - GACATTTGGCTGGTCTGATCC) (Fig. 1A) plus RPr2 (5’ - GAAACAGTGTCGGTATGAAG) (Fig. 1A) or RNeo1 (5’ - GAGGATCTCTGCGTGACCA) (Fig. 1A). This PCR yielded either a 1.8-kb product indicative of the wild-type allele or a 2.8-kb product indicative of the disrupted or null allele using FPr1 and RPr2 (Fig. 1A) and a 1.8-kb product for the mutant allele using only FPr1 and RNeo1. For faster genotyping on a routine basis, a multiplex PCR assay was designed with FPr3 (5’ - AATAGACAGACAGCTGGCCAGGCC) (Fig. 1A), RPr4 (5’ - AAGCACAACCATATCCCCAGACCC) (Fig. 1A), and FNeo1 (5' - ATAGCCGGAATGCCTTCCTACACCAAG) to yield products of 274 bp for the wild-type allele and 500 bp for the disrupted or null allele (Fig. 1C).
alkaline phosphatase reaction, the immunoblot was scanned by using a Typhoon 9200 Variable Mode Imager (Molecular Dynamics, Amersham Pharmacia Bio-tech). CCTβ2 (approximately 42 kDa) was distinguished from CCTβ3 (approximately 38 kDa) by its migration distance and comparison with authentic CCTβ2 and CCTβ3 cDNA standards (32).

Tissue histology and fluorescent immunocytochemistry. Mouse tissues were fixed in 10% formalin and processed by dehydration in 70% ethanol, absolute ethanol, and then xylene. Tissues were infiltrated and embedded in paraffin, cut at 4 μm, mounted on microscope slides, and stained with hematoxylin and eosin. For immunocytochemistry, paraffin sections were dewaxed with xylene three times for 3 min each and were rehydrated with 100% ethanol and then 95% ethanol and were then permealized with 0.3% Triton X-100 in PBS for 20 min at room temperature. Tissues were blocked with 10% fetal bovine serum (FBS) in PBS for 1 h and rinsed in PBS. Affinity-purified primary antibody (rabbit anti-mouse CCTα [38] or rabbit anti-human CCTβ [35]) or preimmune serum at 10 to 20 ng/ml in 2% FBS in PBS was loaded onto the slices and incubated overnight at room temperature. The next morning, slides were washed three times with PBS, secondary goat anti-rabbit Alexa Fluor 488-labeled IgG (Molecular Probes, Inc.) was applied at a 1:250 dilution in PBS for 1 h at room temperature, and the sections were washed three times with PBS. Sections were mounted in 60% glycerol–1.5% propyl gallate in PBS and sealed with nail polish, and a Nikon E800 microscope was used to examine the tissues for fluorescence.

Serum hormone determinations. Whole blood was collected from mice by orbital bleed or cardiac puncture, allowed to clot at room temperature for 1 h, and centrifuged again and serum was collected off the top and stored at −80°C until processed for hormone analyses. Serum FSH and LH were measured by the Ligand Assay and Analysis Core Lab, University of Virginia, Charlottesville.

Phospholipid determinations. Flash-frozen brain tissue was thawed and weighed, and approximately 50 mg was extracted by the method of Bligh and Dyer (8). The organic phase containing lipid was concentrated under nitrogen and resuspended in 400 μl of chloroform:methanol (2:1). A 1-μl aliquot was loaded onto a thin-layer silica gel rod and was developed first in ether, dried, and developed in chloroform:methanol:acetic acid:water (50:25:8:3). Lipid mass was calculated by using a standard curve prepared with egg PtdCho (Matreya, Inc.) and a Nikon E800 microscope was used to examine the tissues for fluorescence.

RESULTS

Quantitative determination of CCT transcripts. We developed quantitative real-time PCR assays to critically evaluate the relative contributions of the different CCTs to PtdCho production in mouse tissues. The PEMT transcripts were also quantified because this enzyme, in addition to the CCTs, mediates a reaction significant for PtdCho biosynthesis in liver tissue (54). This technique has a linear range of more than 4 orders of magnitude which allows for wide-ranging detection of both high and very low copy number transcripts in a sample, and the probe designs are based on a common hybridization condition optimized for all the transcripts. The levels of the transcripts encoding CCTα, CCTβ2, CCTβ3, and PEMT were determined in selected tissues in wild-type 129/Sv × C57BL/6J male and female mice. Both of the CCTα transcripts encode the same protein sequence, and thus the primers were designed to signal both mRNAs. The amounts of target RNA were normalized to GAPDH mRNA and were compared to the level of CCTα in liver (normalized to 1.0) (Fig. 1A). CCTα was most highly expressed in the testis and lung and at lower levels in liver (female) and ovary. The data correlate with the fact that testes and lung secrete large amounts of PtdCho in the seminal fluid and as surfactant, respectively, and the PEMT is expressed at much lower but still significant levels in these organs. On the other hand, liver also has a large capacity for PtdCho production and secretion as a component of the lipoprotein biosynthetic pathway, but the bulk of the PtdCho synthesis that contributes to lipoproteins is governed in large part by the PEMT, which was expressed at a level about 15 times higher (Fig. 1A and D). The high-density lipoprotein deficiency in choline-deficient PEMT knockout mice (44) is consistent with this view. In comparison, brain did not contain large amounts of either of the CCTs per milligram of tissue weight, and PEMT was almost undetectable despite the fact that the brain is a lipid-rich organ. CCTβ2 and CCTβ3 transcripts, which encode proteins that differ at the amino terminus (32), were consistently expressed together in the same tissues and were most highly expressed in brain, lung, and gonad, although in most cases their levels were at least 10 times lower than the expression level of liver CCTα (Fig. 1B and C). Only in brain did the levels of CCTβ2 and β3 approach that of CCTα. This was evident when relative levels were compared to one another within individual organs as shown in Fig. 1F. CCTβ2 was at its highest level in brain, where it constituted 30% of the four transcripts (Fig. 1F), whereas CCTβ3 was expressed at a relative value of only about 0.1 compared to that of liver CCTα (Fig. 1A and B). CCTβ3 in the brain constituted about 50% of the four transcripts (Fig. 1F) and was expressed at a value of about 0.17 compared to that of liver CCTα (Fig. 1A). CCTβ2 expression constituted about 30% of the transcripts expressed in female brain, compared to 50% CCTα and 12% PEMT (Fig. 1F). CCTβ2 contributed 5% to the transcript abundance in lung and 1.6% in ovary. These data suggested that the absence of CCTβ2 expression might affect brain or lung most severely.

Deriving CCTβ2-deficient mice. The role of the Pcyt1b gene encoding the CCTβ2 protein was investigated by deriving a mouse model that lacked the expression of the full-length CCTβ2 isoform due to disruption of the first coding exon. Genomic clones were isolated and a replacement vector was constructed in which the 3′ end of exon 2 (32) was replaced with the neomycin (Neo) resistance gene, and the DTA gene was ligated to the 3′ end of the genomic DNA as a negative selection agent to enrich for homologous recombinants (Fig. 2A). The construct was electroporated into W9.5 ES cells, and several hundred Neo-resistant clones were isolated and screened by Southern blotting and PCR as described in Materials and Methods and as illustrated in Fig. 1B. The ES cells were XY; thus, there was only a single X-linked CCTβ (Pcyt1b) allele present in the genome, and only a single band on the Southern blots or PCR screens was detected for either the wild-type or recombinant gene (Fig. 2B and C). A clone (#25) where the targeting construct had integrated by homologous recombination into the CCTβ gene was identified, and a normal karyotype (n = 40) was confirmed. Clone 25 was injected into C57BL/6J blastocysts and was implanted into pseudopregnant F1 B6/CBA founder mothers at the St. Jude Transgenic Core Facility. Male chimeras (129Sv × C57BL/6J) were bred with C57BL/6J females, and the F1 agouti pups that arose from germline transmission of the disrupted CCTβ2 allele as determined by Southern blot analysis were interbred to generate F2 animals for genotyping (as shown in Fig. 2C) and subsequent study. Female mice with the CCTβ2−/− genotype (25%) and male mice with the CCTβ2−/− genotype (25%) were routinely obtained in the litters derived from breeding the +/- × +/- animals, indicating that there was no fetal death of the homozygous knockout animals. However, a reproductive de-
we surveyed the expression of both of the CCT Neo isoforms by the insertion of the DTA gene into exon 2 at the NcoI site, eliminating part of the coding portion of the exon. The targeting construct inserts the DTA gene by homologous recombination into exon 2 of the CCT gene. (B) An example of Southern blot analysis of mouse genomic DNA. Genomic DNA was isolated and digested with BamHI, and the fragments were separated by agarose gel electrophoresis. The blot was probed with a 9-kb band, whereas the mutant allele gives a 2.8-kb band due to the presence of a new BamHI site within the inserted Neo gene. Sample #25 is the ES cell clone used for blastocyst injections. Probe B was used in Southern blot experiments to detect the presence of the 7.2-kb Neo gene fragment in the BamHI digest to confirm the genotype of the mice (data not shown). (C) An example of the PCR analysis used to genotype mice. A multiplex PCR consisting of a mixture of primers FPr3, FNeo1, and RPr4 were used to signal either the 274-bp wild-type allele or the 500-bp null allele. PCRs used to screen the ES cell clones consisted of FPr1 coupled with RPr2 or RNeo1 to give a 1.8-kb band in the wild-type cells and a 2.8-kb band in the mutant cells (data not shown).

CCT isoform expression in knockout mice. The expression, or lack thereof, of CCTβ2 was confirmed in the mouse pups by performing conventional RT-PCR analysis of the RNA from selected organs from the wild-type and knockout animals. These experiments also addressed whether expression from the CCTβ3 exon 1 lurking 18.5 kb upstream would be affected by the insertion of the Neo gene into exon 2. At the same time, we surveyed the expression of both of the CCTα transcripts, α1 and α2, arising from the alternate gene Pcyt1a (Fig. 3). The CCTβ transcripts were detected visually after 40 cycles of PCR, whereas detection of CCTα transcripts required only 30 cycles, confirming our results from the real-time PCR measurements that CCTβ was expressed at a much lower level in most mouse tissues. Male CCTβ2−/− mice showed the expected distribution of CCTα expression (Fig. 3A) (32), which included the unique expression of CCTα2 in the testis. All male CCTβ2−/− tissues lacked the expression of the CCTβ2 transcript but retained CCTβ3 expression. The same general pattern was true in the female CCTβ2−/− mice (Fig. 3B), which expressed CCTα1, CCTα2, and CCTβ3 transcripts as detected by this methodology. These data established that the expression of the full-length CCTβ2 transcript was abolished in both the male and female knockout mice.

Because CCTβ2 was most highly expressed in brain tissue, we confirmed that the full-length CCTβ2 protein was present in wild-type brains and that it was absent in the CCTβ2−/−
FIG. 4. Immunoblot analysis of CCTβ2 protein expression. Western blotting was performed by using mouse brain tissue. Wild-type (+/+) and knockout (−/−) female brains were homogenized and separated on 8% NuPAGE gels prior to blotting with a CCTβ isoform-specific antibody (32). CCTβ2 migrates at a molecular size of 42 kDa.

brains by immunoblotting tissue homogenates with a CCTβ isoform-specific antibody (Fig. 4). However, measurements of brain PtdCho (in micrograms per milligrams of weight) did not reveal a significant difference between wild-type 129/Sv × C57BL/6J mice (18.97 ± 3.69, n = 24) and knockout mice (18.73 ± 2.36, n = 24). Our results suggested that there may be excess capacity for PtdCho synthesis, represented by the copy number of the rate-controlling enzymes in brain (and other tissues), and the loss of about one-third of the CCT transcripts does not have a gross effect.

We were somewhat surprised that disruption of CCTβ2 expression in mice did not have any overt effect brain development, content, or morphology in light of recent reports that CCTβ2 expression was selectively upregulated in the hippocampus by the neuropeptide arginine-vasopressin (4-8) (63) and during neurite outgrowth of NGF-treated PC12 cells (12). We addressed this inconsistency by using our real-time PCR tools to quantify the changes in the CCT transcript levels in PC12 cells stimulated with NGF to determine whether the information obtained by using an immortalized cell line was distinctly different from that obtained from the whole animal. PC12 cells respond to NGF by ceasing cell division and extending neurite-like outgrowths associated with a fourfold increase in the total protein per cell (23). In our experiments, NGF-treated PC12 cells exhibited neurite outgrowth and the total protein increased from 796.2 μg per 10^7 cells to 2,168.7 μg per 10^7 cells following addition of 100 ng of NGF/ml to the cultures for 5 days. The amount of protein and the amount of total RNA per dish was equivalent for comparisons of NGF-treated and untreated cells cultured for 5 days. However, the cell number of the NGF-treated cultures was significantly lower (2.6 × 10^7 cells in untreated 100-mm-diameter dishes compared to 8.83 × 10^7 cells in those exposed to NGF), consistent with the known functions of NGF to halt proliferation, trigger the differentiation, and support survival of PC12 cells (22, 23, 50). Whereas the relative amount of CCTβ2 transcript per cell increased by a factor of 3.8, from 0.268 to 1.02 in our measurements, in agreement with the previous report (12), the amount of CCTα transcript per cell also increased, from 1.0 to 2.48 (Fig. 5A). We also quantified the CCT transcript levels normalized to total RNA, and the relative amounts of CCTα and CCTβ2 transcripts were not significantly different in control and NGF-treated cells. The ratio of CCTα to CCTβ2 transcript was about 3.7 to 1 in undifferentiated PC12 cells and was 2.5 to 1 in PC12 cells treated with NGF (Fig. 4), in general agreement with our data for primary brain tissue, where there was a 2:1 ratio of CCTα and CCTβ2 and -β3 expression (Fig. 1F). CCTβ3 was found not to be expressed in the PC12 cells in both labs. Therefore, these data point out an increase in CCTα expression as well as CCTβ2 expression upon neuronal differentiation and are consistent with our CCT expression results for mouse brain. These data also support the view that sufficient CCTα is present in the brain to support cellular and tissue function and structure in the absence of CCTβ2.

We investigated whether there were compensatory changes in CCT expression to explain normal organ development and function in CCTβ2-deficient mice. Previously in mice where CCTα expression was eliminated in macrophages, the cells survived due to an upregulation of CCTβ2 expression (66). There was no change in the CCTα, CCTβ3, or PEMT transcript levels in liver, brain, or gonads of the knockout CCTβ2 mice (data not shown). However, in lung there was a 28% increase in CCTα expression in the male and about a 50% increase in the female CCTβ2-deficient mice. The result was selective for CCTα, as the PEMT transcripts which constituted a small but significant portion of those surveyed in lung did not change. The amount of CCTβ3 transcript also did not change (data not shown). The reason for this apparent upregulation of CCTα expression in the lungs of CCTβ2 knockout animals is not clear.

Reproductive phenotype of the CCTβ2-deficient mice. Heterozygous CCTβ2+/− females were mated with CCTβ2−/− males, and the CCTβ2−− female pups appeared outwardly
FIG. 6. Fertility of CCTβ2/-/- females. Female mice (2 to 4 months old) with the indicated genotypes were paired 1:1 with either wild-type or CCTβ2-/- males as indicated for 2 months and were scored as fertile if they were pregnant or had borne pups at least once by the end of that time period. Comparison of the mating groups was done by the Fisher exact test. For CCTβ2 +/- x CCTβ2-/- (n = 27) compared to CCTβ2-/- x CCTβ2-/- (n = 19), P < 0.001; for CCTβ2-/- x CCTβ2 +/- (n = 19) compared to CCTβ2-/- x CCTβ2-/- (n = 16), P = 0.022; for CCTβ2 +/- x CCTβ2-/- (n = 6) compared to CCTβ2-/- x CCTβ2-/- (n = 6), P < 0.001; for CCTβ2 +/- x CCTβ2-/- or CCTβ2 +/- x CCTβ2-/- (n = 30) compared to CCTβ2-/- x CCTβ2-/- or CCTβ2-/- (n = 36), P < 0.001.

normal. However, the adult homozygous CCTβ2-/- females had reduced numbers of pregnancies. CCTβ2-/- females delivered normal numbers of pups per litter (8.3 ± 2.4, n = 22) compared to the wild-type 129/Sv x C57BL/6J females (7.7 ± 2.3, n = 14), and the female and male pups in both groups were average weight (CCTβ2 +/- mice, 17.3 ± 0.9 g, n = 10; CCTβ2-/- mice, 17.8 ± 0.85 g, n = 7; CCTβ2-/- mice, 19.9 ± 1.4 g, n = 8; CCTβ2-/- mice, 19.7 ± 0.7, n = 10 at 5 weeks). A study was done where 1:1 mating pairs of homozygous, heterozygous, and wild-type virgin mice between the ages of 2 and 4 months were maintained together for 2 months and the females were scored as fertile or barren. The data in Fig. 6 report the percentages of mature females who had at least one litter. Those mice which were scored as barren did not show evidence of pregnancy. Less than 50% of the homozygous CCTβ2-/- females were fertile when mated with wild-type 129/Sv x C57BL/6J males (P < 0.001 compared to wild-type females), and fertility decreased when the CCTβ2-deficient females were mated with knockout CCTβ2-/- males (P < 0.001) (Fig. 6). The percentage of heterozygous females mated to either wild-type or knockout males and that birthed pups was also reduced significantly (P < 0.001) (Fig. 6). Keeping in mind that the Pexylb gene is located on the X chromosome, this result may have been due to random X inactivation of the single wild-type allele which would occur with about a 50% frequency, conferring a knockout phenotype on the heterozygous females with two effectively dysfunctional alleles. Alternatively, the lower pregnancy percentage of the heterozygous females could have been due to a gene dosage effect. There was a significant drop (P < 0.022) in female pregnancy when the CCTβ2 homozygous knockout females were bred with CCTβ2-deficient males compared to results for breeding with wild-type males, suggesting a reproductive defect in the male knockout mice as well (Fig. 6). Tissue samples were obtained from the 4- to 6-month-old females and males in this study for histological analysis (see below).

Ovary pathology in CCTβ2-/- knockout females. An explanation for the reduced fertility in the groups of CCTβ2-/- and CCTβ2 +/- females was evident from microscopic examination of the ovaries. Ovaries from a 4-month-old wild-type mouse and a knockout mouse are shown in Fig. 7. The wild-type ovaries exhibited normal ovary morphology with developing follicles and multiple corpora lutea. Higher magnification revealed highly organized germinal epithelial cells and interstitial stromal cells and follicles at several stages of development with ova. However, the infertile CCTβ2-/- mice had no follicles or corpora lutea. Higher magnification revealed epithelial and interstitial stromal cell hyperplasia, with invagination of the epithelial cells into the interstitial stromal cell layer forming epithelial tubules and cords of interstitial stromal cells. Fertile CCTβ2-/- mice, on the other hand, had ovaries with normal morphologies. The ovaries of a subset of the heterozygous females (n = 10) were also examined, and 40% exhibited aberrant ovarian morphology, with the absence of follicles and corpora lutea. Those mice with the abnormal ovarian pathology were also barren. The data support the idea that the reproductive defect in CCTβ2-deficient females was due to a failure in ovary maturation.

We compared ovarian pathologies as a function of age. At 4 weeks, 89% of knockout virgin females had normal ovaries (n = 9); 45% of the animals between 2 to 5 months of age were normal (n = 17), at 6 months 38% were normal (n = 8), and at 16 months 17% of the ovaries were normal (n = 6). Examination of wild-type 129/Sv x C57BL/6J ovaries on multiple occasions between ages 4 weeks to 6 months showed 100% normal ovaries. In a separate study, when immature 4-week-old CCTβ2-/- female mice were treated with FSH and LH to induce follicle maturation and ovulation and then were mated for 1 month, 3 out of 10 were fertile compared to age-matched wild-type 129/Sv x C57BL/6J females, all of which successfully became pregnant (100%, n = 10). These data suggested that that ovary dysfunction and infertility preceded the overt pathology of the defective ovaries in the younger CCTβ2-/- females and that the ovaries of an increasing percentage of females were not able to develop follicles as the animals aged. The abnormal ovaries from the 16-month-old CCTβ2 knockout females had infiltrating tubulostromal tumors (Fig. 8). The tumor cells were similar to the epithelial and interstitial stromal cells of the ovaries of the CCTβ2-/- females, but the tumor cells invaded the adipose tissue of the ovary's hilus. Although the tumor cells were invasive, their cytological morphology was benign and mitotic figures were not evident. We concluded from these results that long-term absence of CCTβ2 expression in ovarian tissue resulted in the development of benign ovarian tumors. The older CCTβ2-/- tumors varied in their proportion of epithelial and interstitial stromal cell components, and occasionally there were foci of granulosa cells.

The serum FSH and LH levels of 4-month-old mice were investigated and correlated with ovarian morphologies. We found a 100% correlation between aberrant ovarian pathology and elevated FSH and LH levels. Wild-type and CCTβ2-deficient animals with normal ovaries had average serum FSH values (± standard error) of 7.1 ± 0.5 ng/ml and LH of 0.4 ± 0.05 ng/ml (n = 40), whereas the CCTβ2-/- animals with ovaries that lacked follicles or corpora lutea had significantly higher FSH values of 63.7 ± 3.4 ng/ml (P < 0.0001) and LH values of 5.5 ± 1.2 ng/ml (P < 0.0001) (n = 10). Occasionally
an ovary from either a CCTβ2−/− or CCTβ2−/+ female would exhibit evidence of follicles along with hyperplasia of the interstitial stromal cells, and the FSH levels in these cases were midrange.

Testis pathology and serum hormone levels in CCTβ2−/− males. An explanation for the reduced fertility of the CCTβ2−/− males (Fig. 6) was evident from microscopic examination of the testes from the 4- to 6-month-old mice. A testis from a 6-month-old CCTβ2−/− male is shown in Fig. 8. Wild-type testes exhibited normal organization of seminiferous tubules with spermatogenesis, mature sperm, and tubular interstitial Leydig cells (Fig. 9A and B). Testes from CCTβ2-deficient mice showed seminiferous tubular degeneration at multiple foci and reduced spermatogenesis (Fig. 9C and D). The number of animals with abnormal testicular pathology and the severity of testicular degeneration increased with age. Two- to 4-month-old CCTβ2-deficient males exhibited mild seminiferous tubular degeneration (58%, n = 12), while at 6 months 71% showed severe multifocal seminiferous tubular degeneration (n = 7 animals). These data are consistent with the reduced fertility seen with the CCTβ2−/− males (Fig. 6). The fact that successful insemination by males can still occur in animals with lower than normal sperm production explains why the male reproductive defect was not as obvious until the males aged. In contrast with the CCTβ2−/− females, the aberrant testis morphology of the CCTβ2-deficient males was not ac-

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**FIG. 7.** Aberrant ovary morphology in CCTβ2−/− mice. The ovaries from wild-type and CCTβ2−/− knockout mice were removed, fixed, and stained for pathology analysis. (A) View (magnification, ×4) of a normal ovary showing developing follicles and multiple corpora lutea. Structures indicated are the following: 3°F, tertiary follicle with ovum; O, ovum; G, granulosa cells; CL, corpus luteum; and I, interstitial stromal tissue. (B) View (magnification, ×40) of a normal ovary showing the following: Pr, primordial follicle with ovum; 1°F, a primary follicle with ovum surrounded by a layer of granulosa cells; and GE, organized germinal epithelial cell layer. (C) View (magnification, ×4) of a CCTβ2−/− knockout ovary illustrating the absence of ova, follicles, and corpora lutea. (D) View (magnification, ×40) of a CCTβ2−/− ovary showing hyperplastic epithelial cells, invaginating into and dissecting the interstitial stromal cell tissue forming epithelial tubules (ET) and cords of interstitial stromal cells with foamy cytoplasm (I).
accompanied by successively higher levels of sperm production.

Deletion of CCTβ2 expression in mice revealed an essential role for this isoform in spermatogenic function and the maintenance of sperm production. Considering the low level of CCTβ2 expression in gonads that was revealed by quantitative real-time PCR, it is remarkable that the animals have a gonadal phenotype. It would be virtually impossible to link a biochemical defect in PtdCho metabolism to the loss of CCTβ2 expression due to the very low expression level of CCTβ2 compared to that of CCTα, which is expressed at a 20-fold higher level in ovaries and a 350-fold higher level in testes (Fig. 1). CCTβ3 is also expressed in the CCTβ2-deficient animals and probably performs the same cellular function as CCTβ2 due to its nearly identical protein sequence. The challenge of repeated maturation of the gametes during adulthood accompanied by successive rounds of cellular differentiation may have allowed the phenotypic deficiency to be revealed.

Animals had reduced fertility due to defective ovarian follicular development and testicular spermatogenesis. Postnatal ovarian follicles were evident, as were testicular spermatogeno-
nia, Sertoli cells, and interstitial Leydig cells in CCTβ2 knockout weaning animals, and both tissues arise from the same embryonic origin (25), suggesting that CCTβ2 is not necessary for the embryonic development of the gonads. However, a large percentage of the older female mice that lacked CCTβ2 expression were unable to develop mature ovarian follicles, and CCTβ2-deficient testes became progressively atrophic with reduced spermatogenesis. Follicle and oocyte maturation, as well as testicular stem cell renewal and differentiation into Sertoli cells and spermatids, occur in response to circulating pituitary hormonal stimuli in mature animals. However, the gonadal problems did not arise from deficient reproductive hormone production from the pituitary. The high levels of FSH and LH in the CCTβ2−/− females with abnormal ovaries coupled with the lack of elevated FSH and LH levels in the CCTβ2−/− males with degenerating testes argue that the pituitary gland is capable of synthesizing and secreting ample amounts of the reproductive hormones. The lack of a growth defect in the CCTβ2-deficient mice is consistent with sufficient secretion of growth hormone, another pituitary product, and microscopic examination of brain slices through the hypothalamic-pituitary axis did not reveal alterations in aged knockout animals (data not shown). Thus, the reproductive defect probably arises from a unique requirement for CCTβ2 activity in

FIG. 9. Aberrant testis morphology in CCTβ2−/− mice. The testes from 6-month-old CCTβ2−/− knockout mice and wild-type littermate controls were removed, fixed, and stained for pathology analysis. (A) View (magnification, ×2) of a wild-type testis showing seminiferous tubules with small lumens (S). (B) View (magnification, ×20) of testis showing seminiferous tubules with 4 to 5 germinal cell layers composed of spermatogonia and Sertoli cells and spermatogenesis progressing from the basal germinal layer (G) to mature spermatids (S) with their tails extending into the lumen. Interstitial Leydig cells (I) are located in the space between the seminiferous tubules. (C) View (magnification, ×2) of a CCTβ2−/− knockout testis illustrating multiple foci of degenerate seminiferous tubules with a large lumen (D) and also normal seminiferous tubules with a small lumen (S). (D) View (magnification, ×20) of a CCTβ2−/− knockout testis showing multiple small degenerative seminiferous tubules with reduced number of germinal cell layers and absence of spermatozoa. Sertoli cells (Sr) are the predominant cells associated with the basal germinal layer, and interstitial Leydig cells (I) are increased between the tubules.
the gonads, where the CCTβ2 protein functions in a pathway critical to maintaining the repeated germ cell differentiation that takes place in the gonads after puberty and through adulthood. CCTβ3 is still expressed in the knockout mice and is governed by a different promoter (32); therefore, the loss of response of the ovary and testis to stimulatory ligands or growth factors may be due to total CCTβ expression falling below a threshold level.

There are several signaling pathways that are essential for normal ovary maturation and maintenance. Development of the gonads after birth is dependent on FSH and LH production by the pituitary gland and on the production of steroid hormones in the testis and ovary (33, 61). Gonad maturation is also dependent on growth factors as shown in mice with mutations at the white-spotting (W), steel (Sl), or deficiency in growth differentiation factor 9 (Gdf9) loci or sheep with the FecX (Inverdale) defect (9, 11), which all have reduced fertility due to the absence of normal ovarian follicle development followed by ovarian degeneration. Mice with the W/W defect also develop ovarian tumors (1). The W gene encodes the c-kit growth factor receptor (13, 43), which is expressed in growing oocytes as well as other cell types, and the Sl (steel) locus encodes the ligand for the c-kit receptor and is expressed in follicle cells (26, 67). Mice with homozygous mutations in either locus lack germ cells, and their ovaries develop to the stage when germ cells would enter meiosis but have arrested differentiation (41, 42). GDF-9 (also known as FecX) is an oocyte-derived growth factor required for ovarian somatic cell function (21, 39, 62). Similar to the CCTβ2−/− mice, ovarian degeneration or regression in animals with these mutant alleles is associated with significantly elevated serum FSH and LH (39, 49).

CCTα is the dominant isoform (Fig. 1) and is expressed in virtually all gonadal cells (Fig. 10), although its highest level occurs in the nuclei of the interstitial stromal cells (Fig. 10). These cells produce progesterone, providing an essential step in female cycling. The immunocytochemistry also reveals that CCTβ2 is a major determinant of PtdCho synthesis in wild-type maturing ova (Fig. 10). CCTβ2−/− ovaries have staining, indicating that CCTβ3 is still expressed in the ova and suggesting that disruption of both β2 and β3 expression might yield a completely infertile female phenotype. There is an intimate relationship between ova and granulosa cells that is essential to the maturing follicle illustrated by the research on GDF-9 (40), and epidermal growth factor (EGF)-like receptors are critical mediators of LH action in ovaries (46). One interpretation of our data is that CCTβ2-mediated PtdCho synthesis may be required to support the paracrine stimulation between these

FIG. 10. Fluorescent immunocytochemistry of CCT expression in ovary. Wild-type C57BL/6J × 129J ovary (age, 4 months) was fixed and stained with hematoxylin and eosin (A), rabbit anti-mouse CCTα antibody (B), or rabbit anti-human CCTβ antibody (C), followed by goat anti-rabbit Alexa Fluor 488-coupled secondary IgG (H + L). Substitution of preimmune rabbit serum for the primary antibody did not yield a fluorescent image at the same exposure. Structures indicated are the following: I, interstitial stromal cells; O, ova; and G, granulosa cells. CCTα is a dominant isoform (Fig. 1) and is expressed in virtually all gonadal cells (Fig. 10), although its highest level occurs in the nuclei of the interstitial stromal cells (Fig. 10). These cells produce progesterone, providing an essential step in female cycling. The immunocytochemistry also reveals that CCTβ2 is a major determinant of PtdCho synthesis in wild-type maturing ova (Fig. 10). CCTβ2−/− ovaries have staining, indicating that CCTβ3 is still expressed in the ova and suggesting that disruption of both β2 and β3 expression might yield a completely infertile female phenotype. There is an intimate relationship between ova and granulosa cells that is essential to the maturing follicle illustrated by the research on GDF-9 (40), and epidermal growth factor (EGF)-like receptors are critical mediators of LH action in ovaries (46). One interpretation of our data is that CCTβ2-mediated PtdCho synthesis may be required to support the paracrine stimulation between these

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cells. Flux through a CCTβ2-dependent pathway for PtdCho production associated with membrane trafficking may support a turnover pathway stimulated by a growth factor(s) (29), thus leading to the failure to maintain differentiated ovaries in CCTβ2−/− mice. This latter idea is supported by recent work with the Drosophila system that implicates CCT in EGF receptor and Notch signaling (60), where disruption of one of two CCT genes leads to defective ovarian morphogenesis (24). Flies deficient in CCTI have reduced signal transduction through these two pathways that correlates with an increase in the sequestration of EGF receptors in an enlarged endocytic compartment. The genetic connection between CCTI, Egfr, and Notch (60) supports an emerging role for PtdCho metabolism in metazoan vesicular trafficking (6, 27). Future work with the CCTβ2−/− mice will investigate this hypothesis in more detail.

The real-time PCR data indicate that both CCTα and CCTβ2 transcripts increase during neurite outgrowth of PC12 cells. The inconsistency of this result with those recently reported (12) may be explained by the limited linear range, particularly at the high end of the scale, of the visual detection used in conventional RT-PCR methodology compared to that of the multilog linear range for amplicons in the real-time PCR assays used here. Our data (Fig. 5) and those of Araki and Wurtman (1) show that CCT expression and enzyme activity are not selectively regulated by NGF treatment of PC12 cells. Rather, both CCTα and CCTβ2 increase in a ratio of about 2:1, and these data agree with the expression measurements in the mouse brain. Also, the lack of a neural phenotype argues that CCTβ2 expression is not an essential component of NGF signaling as compared to the severe defect in neuronal cell survival in mice lacking the NGF receptor (16).

In general, CCTβ2 transcripts are found at levels 10-fold lower than those encoding CCTα, and therefore CCTβ2 is likely to contribute only a small percentage of the CDP-Cho required for a tissue to sustain bulk PtdCho synthesis. If the steady-state mRNA abundance reflects the relative amount of CCT isoforms in cells, then CCTβ2 usually contributes 10% or less to the total CCT activity in most tissues. The exception is brain, where CCTβ transcripts are approximately in equal abundance with CCTα, and CCTβ3 transcripts constitute a small portion of the CCT pool (Fig. 1). This selective expression pattern still may suggest a specific role for CCTβ2 in brain physiology, and future work will investigate the development of primary neurons in a more detailed manner. Our data with CCTβ2-deficient mice, together with the results in mice with a macrophage-specific CCTα deletion (66), suggest that CCT
expression is in excess in most wild-type primary cells and that selected cell types can increase CCT expression when the PtdCho synthetic capacity for normal growth and function is challenged.

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