Crystal Structure of a Mammalian CTP: Phosphocholine Cytidylyltransferase Catalytic Domain Reveals Novel Active Site Residues within a Highly Conserved Nucleotidyltransferase Fold*†§

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CTP:phosphocholine cytidylyltransferase (CCT) is the key regulatory enzyme in the synthesis of phosphatidylcholine, the most abundant phospholipid in eukaryotic cell membranes. The CCT-catalyzed transfer of a cytidylyl group from CTP to phosphocholine to form CDP-choline is regulated by a membrane lipid-dependent mechanism imparted by its C-terminal membrane binding domain. We present the first analysis of a crystal structure of a eukaryotic CCT. A deletion construct of rat CCT α spanned residues 1–236 (CCT236) lacks the regulatory domain and as a result displays constitutive activity. The 2.2-Å structure reveals a CCT236 homodimer in complex with the reaction product, CDP-choline. Each chain is composed of a complete catalytic domain with an intimately associated N-terminal extension, which together with the catalytic domain contributes to the dimer interface. Although the CCT236 structure reveals elements involved in binding cytidine that are conserved with other members of the cytidylyltransferase superfamily, it also features nonconserved active site residues, His-168 and Tyr-173, that make key interactions with the β-phosphate of CDP-choline. Mutagenesis and kinetic analyses confirmed their role in phosphocholine binding and catalysis. These results demonstrate structural and mechanistic differences in a broadly conserved protein fold across the cytidylyltransferase family. Comparison of the CCT236 structure with those of other nucleotidyltransferases provides evidence for substrate-induced active site loop movements and a disorder-to-order transition of a loop element in the catalytic mechanism.

A key rate-limiting step in the synthesis of phosphatidylcholine in animal cells is the formation of the headgroup donor, CDP-choline, by transfer of a cytidylyl group from CTP to phosphocholine. This reaction is catalyzed by CTP:phosphocholine cytidylyltransferase (CCTα; EC 2.7.7.15), an enzyme subject to many layers of regulation (1–4). The ubiquitous and best studied isoform of mammalian CCT (CCTα, 367 residues) has been described as having four domains (Fig. 1A). An N-terminal domain (−75 residues) housing its nuclear localization signal (NLS) sequence is followed by an −150-residue catalytic domain, an −60-residue membrane binding domain (domain M), and an unstructured phosphorylated tail (−50 residues) (2, 4). CCT functions as a homodimer (5).

CCT activation requires transformation of the enzyme from a soluble form to a membrane lipid-bound form. When the full-length soluble CCT interacts with anionic membrane surfaces, domain M transforms from a mixture of structural elements into an amphipathic α-helix (6–8). Domain M appears to act as an autoinhibitory device, whose interaction with phosphatidylcholine-deficient membranes releases an inhibitory constraint at the active site to enhance $k_{cat}$ by 2 orders of magnitude (9). The primary evidence for this model is the constitutive activity of a construct lacking domain M, CCT236 (9).

To elucidate the mechanism whereby membrane binding activates this important regulatory enzyme, we must first understand how the active form of the enzyme accomplishes catalysis. The cytidylyl transfer reaction catalyzed by CCT involves direct attack at the α-phosphate of CTP by a phosphoryl oxygen of phosphocholine acting as the nucleophile to displace the β,γ-phosphates (pyrophosphate) of CTP and form the product CDP-choline (Fig. 1B). Many aspects of this reaction mechanism resemble those of other adenyl- and cytidylyltransferases, in which the transition state involves conversion of tetrahedral geometry at the α-phosphate to a planar

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‡ The on-line version of this article (available at http://www.jbc.org) contains supplemental Tables 1–3 and Figs. S1–S5.

The atomic coordinates and structure factors (code 3HL4) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

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4 The abbreviations used are: CCT, α isoform of CTP:phosphocholine cytidylyltransferase; GCT, glycerol-phosphate cytidylyltransferase; ECT, ethanolamine-phosphate cytidylyltransferase; aa-tRS, amino acyl tRNA synthetase; PPAT, phosphopantetheine adenylyltransferase; NLS, nuclear localization sequence; r.m.s.d., root mean square deviation; PDB, Protein Data Bank; DTT, dithiothreitol.
bipyramidal penta-coordinate geometry (10–12). One well characterized cytidylyltransferase with a solved structure is glycerol-phosphate cytidylyltransferase (GCT), which catalyzes the synthesis of CDP-glycerol, an intermediate in teichoic acid synthesis for bacterial cell walls. Both GCT from Bacillus subtilis and CCTα from rat show random order (ternary) kinetics and have similar $k_{cat}$ and $K_m$ values (9, 13, 14). The catalytic domain of CCT has 34% sequence identity to GCT overall and very strong homology in several regions (Fig. 2). Crystal structures of GCT with bound CTP (PDB code 1COZ) (15) or with bound CDP-glycerol (PDB code 1N1D) (16) reveal a dimer of an $\alpha/\beta$-nucleotide binding fold composed of a twisted five-stranded parallel $\beta$-sheet flanked by five helices. The ligands are found in a pocket at the base of the $\beta$-sheet, in contact with the conserved HXGH and RTEGISTT motifs that are signatures of the GCT family of nucleotidyltransferases (15). This family includes ethanolamine-phosphate cytidylyltransferases (ECT), phosphopantetheine adenyllyltransferase (PPAT), and the class I aminoacyl tRNA synthetases (aa-tRS) (10, 15–17). Multiple solved structures of PPAT and aa-tRS enzymes have been analyzed, and one as yet unanalyzed structure of human ECT was subjected to gel filtration using a Sephacryl S-200 column (76-ml bead volume) equilibrated with 10 mM Tris-HCl (pH 7.4), 1 mM EDTA, 150 mM NaCl, 2 mM DTT. Fractions containing the purified CCT236 were pooled and concentrated to 12 mg/ml (0.45 mM) using an Amicon Ultra-15 filter. CDP-choline was added to a final concentration of 20 mM. Approximately 2 mg of pure CCT236 was obtained per liter of cell culture suspension. The calculated molecular mass of this construct is 26,755 Da, and its sequence is the first 236 residues of UniProt accession number P19836.

**EXPERIMENTAL PROCEDURES**

**Baculovirus Expression and Purification of CCT236**—An untagged version of residues 1–236 of rat CCTα (CCT236) was expressed in _T. ni_ cells using baculovirus as described (8, 9). After CM-Sepharose and Blue-Sepharose chromatography, the purified protein was concentrated to 10 mg/ml using an Amicon Ultra-15 filter, spun at 2800 × g. The protein was then subjected to gel filtration using a Sephacryl S-200 column (~76-ml bead volume) equilibrated with 10 mM Tris-HCl (pH 7.4), 1 mM EDTA, 150 mM NaCl, 2 mM DTT. Fractions containing the purified CCT236 were pooled and concentrated to 12 mg/ml (0.45 mM) using an Amicon Ultra-15 filter. CDP-choline was added to a final concentration of 20 mM. Approximately 2 mg of pure CCT236 was obtained per liter of cell culture suspension. The calculated molecular mass of this construct is 26,755 Da, and its sequence is the first 236 residues of UniProt accession number P19836.

**E. coli Expression and Purification of Selenomethionine-incorporated CCT236**—The cDNA encoding CCT236 was ferred from the baculovirus shuttle vector into pGEX6p1 via NcoI and BamHI cloning sites, to create a fusion protein with a PreScission protease-cleavable glutathione S-transferase tag. The pGEX6p1-CCT236 plasmid was transformed into the BL21-derived Rosetta cell strain (Novagen) for protein expression (18). The overexpression protocol for selenomethionine-incorporated protein was based on the method of Dublié (19).

Minimal media (1 liter) were inoculated with Rosetta cells from a 10-ml overnight culture and grown at 37 °C for 17 h. A mixture of amino acids was added to reach final concentrations of selenomethionine, isoleucine, leucine, and valine (each 60 mg/liter) and lysine, phenylalanine, and threonine (each 100 mg/liter). Protein expression was induced with 0.5 mM isopropyl-thiogalactoside for ~20 h at 28 °C. The cells were harvested by centrifugation at 2800 × g for 15 min and stored at −80 °C. Thawed cells were resuspended in 40 ml of phosphate-buffered saline (pH 7.4), 1 mM phenylmethylsulfonlfyl fluoride, 0.2 mg/ml lysozyme, 0.1 mg/ml DNase, and 1 mM DTT and were incubated for 1 h on ice. The lysed cells were cycled through a high pressure homogenizer (Avestin). The homogenate was clarified by centrifugation at 27,000 × g for 30 min. The selenomethionine-substituted GST-CCT236 was purified by glutathione-agarose affinity and size-exclusion chromatography. The glutathione beads (0.8 ml) were equilibrated in phosphate-buffered saline with 1% Triton X-100, 1 mM phenylmethylsulfonlfyl fluoride, 1 mM DTT. The clarified lysate was bound to beads with gentle mixing for 3 h at 4 °C. The bound fusion protein was collected by centrifugation at 460 × g for 3
The beads were washed three times with the equilibration buffer. The beads were resuspended in cleavage buffer (50 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 mM EDTA, 1 mM DTT), and CCT236 was released from the matrix by incubation with the PreScission protease (Amersham Biosciences) overnight at 4 °C with gentle mixing. The cleaved CCT236 was applied to a Superdex 200 HR 10/30 column (GE Healthcare), equilibrated, and eluted with 10 mM Tris-HCl (pH 7.4), 1 mM EDTA, 150 mM NaCl, 2 mM DTT. The pure fractions were pooled and concentrated using an Amicon Ultra-Spin Filter at 2800 × g, and a 10-fold molar excess of CDP-choline was added. The final protein concentration used for crystallization was 13 mg/ml (0.5 mM) with 5 mM CDP-choline. This purified CCT contains residues 1–236 of rat CCT with an N-terminal linker (GPLGSA). The specific activities of untagged CCT236 versus CCT236 with the N-terminal linker were the same (2583 ± 377 units/mg for untagged CCT236 (n = 10) and 2783 ± 260 units/mg (n = 4) for CCT236 with the linker). Matrix-assisted laser desorption ionization time-of-flight mass spectrometry gave a mass of 27,396 Da, consistent with selenomethionine incorporation into four of the five possible sites, on average. Purified CCTs were stored at −80 °C.

**CCT236 Crystallization and Diffraction Data Collection**—The native CCT236 (0.45 mM CCT with 20 mM CDP-choline) was crystallized using the sitting drop vapor-diffusion method with a reservoir condition that contained 0.1M sodium acetate (pH 4.6) and 1.4M sodium formate. The protein sample (2 μl) was mixed with 1 μl of the reservoir solution and the drop was incubated over 1 ml of the reservoir. After 7 weeks, the drop was replenished with an additional 1 μl of the same protein sample to extend crystall growth for an additional 5 months. After transfer to a cryo-solution (0.1M sodium acetate (pH 4.6) and 1.4M sodium formate), the crystal was looped and cryo-cooled in liquid nitrogen. A highly redundant diffraction dataset was collected at the Advanced Light Source (ALS) beamline 8.2.2 using an ADSC Q315 detector. 360 images were collected with a 1° oscillation angle and a 5-s exposure time. The initial 300 images were used for data processing and refinement (Table 1).

**FIGURE 2. Alignment of sequences and secondary structure elements of the cytidylyltransferase family.** The aligned sequences are as follows: GCT from *B. subtilis* (P27623); domains N and C of CCT from *Rattus norvegicus* (P19836), *Ciona intestinalis* (XP 002130773) (This sequence is a translation from the *Ciona* genome. The others are from cloned cDNAs), *Caenorhabditis elegans* (P49583), *Arabidopsis thaliana* (Q42555), *Saccharomyces cerevisiae* (P13259), and *Plasmodium falciparum* (P49587); ECT (residues 1–156) from Homo sapiens (Q99447). The alignment was generated by ClustalW2 (62) and then manually adjusted. Residues within a pale gray box are conserved 6/8 times, and residues within a dark gray box are identical. Open boxes surround two sequence elements in segment N of the animal CCTs that are conserved. The secondary structure elements observed in GCT (PDB code 1N1D, gray) and CCT236 (this paper; black) are indicated above the sequence alignment. The layout with secondary structure elements was generated with Espript 2.2 (63). The active site residues involved in CDP coordination are marked with stars, and residues surrounding the choline are marked with circles.
Structure of a Mammalian Cytidylyltransferase

The protein sample (1 μl) was mixed with 1 μl of the reservoir condition and incubated over a 0.5-ml volume of the reservoir. The crystals appeared within 1 week. The cryo-protecotant condition was 20 mM sodium citrate (pH 5.6), 20% PEG 8000, 0.4 M lithium sulfate, and 30% glycerol. The crystal was looped and cryo-cooled in liquid nitrogen prior data collection. A complete SAD dataset was collected at the ALS beamline 8.2.1 using an ADSC Q315R detector as described above for the native crystal (Table 1).

Structure Determination and Refinement—The diffraction data were processed with the program HKL2000 (20). The programs HKL2MAP version 0.2 (21) and AUTOSOL within PHENIX version 1.3 (22) were used to solve the P2 crystal structure of selenomethionine-substituted CCT236 using single anomalous diffraction. The program AUTOBUILD within PHENIX version 1.3 automatically performed density modification and partially constructed the polypeptide chain. The rest of the molecule was built and fitted using the program COOT (23), and restrained refinements were carried out within the program REFMAC5 (24). The P2,2,2 native crystal structure was solved by molecular replacement using the partially refined selenomethionine P2 dimer with the program MOLREP (25). The model was manually adjusted and refined using COOT (23). The data collection, phasing, and refinement statistics are summarized in Table 1. The P2 crystals have four protein chains in the asymmetric unit with a Matthews coefficient (Vm) and solvent content of 2.3 Å3 Da−1 and 47.3% respectively. The P2,2,2 crystals have two protein chains in the asymmetric unit with a Vm and solvent content of 2.3 Å3 Da−1 and 46.4%, respectively.

Structure Analysis—The secondary structure elements were analyzed with programs DSSP (26) and PROMOTIF (27). The program CONTACT within the program suite CCP4 (28) was used to identify interacting residues and measure their distances. The Protein-Protein Interaction Server (29, 30) and PISA web service (31) were used to analyze the dimeric interface, and the SSM Superpose (32) in Coot (23) was used to align the protein structures. The CASTp server (33) was used to identify and analyze pockets on the surface, and SURFACE RACER 1.2 (34) was used to calculate the protein surface area with a probe radius of 1.4 Å. The DALI server (35) was used to find proteins with a similar protein fold. The stereochemistry of the final model was analyzed with PROCHECK (36).

Site-directed Mutagenesis and Purification of H168A and Y173A Mutant CCTs—Site-directed mutagenesis was carried out using QuickChange mutagenesis (Stratagene), following company protocols. The template was the pGEX6p1-CCT236 plasmid described above. The primers used for the H168A mutation were 5′-gattgattcttcggccgctgagatccctctctc-3′ (sense) and 5′-ctactaaaaagcagccggccgactgctatgagttagac-3′ (antisense). The primers used for Y173A mutation were 5′-catgcagatcatcccgcccttcggccgaggg-3′ (sense) and 5′-tcctggcgagagccggaggatgcgtcatgc-3′ (antisense). Annealing temperatures of 78.5 and 80 °C were used during respective PCRs. The H168A, Y173A double mutant was created by introducing the Y173A mutation into the H168A template plasmid, using QuickChange and the same mutagenic primer detailed above. The GST-CCT mutant proteins were expressed in BL21 Rosetta cells (18), purified via glutathione affinity chromatography, and cleaved with PreScission protease, as described above. The protein concentrations in the pooled elution fractions were measured by the Bradford assay (37).

CCT Enzyme Kinetics—CCT activity was measured by monitoring the conversion of [3H]-labeled phosphocholine (Amer- sham Biosciences) into labeled CDP-choline (38). The reaction volume was 40 μl, and each reaction contained 1 μg of CCT236, 20 mM Tris (pH 7.4), 12 mM magnesium chloride, 89 mM NaCl, 10 mM DTT, and variable concentrations of the substrates, phosphocholine and CTP. The reactions were carried out for 10 min at 37 °C, with agitation, in the absence of added lipid vesicles. The phosphocholine dependence was monitored over a range of 0–5 mM with the CTP concentration held constant at 8.8 mM. The CTP dependence was monitored over a range of 0–27 mM with the phosphocholine concentration held constant at 1.5 mM.

The kinetic data were analyzed by nonlinear regression analysis using GraphPad Prism software. The initial velocity (nmol/mg/min) versus substrate concentration data were fit to the sigmoidal Hill equation, ν = (Vmax [S]^n)/(K^n + [S]^n), where ν = reaction rate; Vmax = maximum reaction rate; [S] = substrate concentration; K^n = the Hill constant, i.e. the substrate concentration producing half-maximal velocity; and n = Hill coefficient. We observed substrate inhibition at high phosphocholine concentrations for CCT236 wild type and the Y173A mutant. Thus, the activity data used to calculate the K^n(P-chol) for these constructs was limited to phosphocholine concentrations from 0 to 2 mM (wild type) and 0 to 1.4 mM (Y173A). CTP substrate inhibition was observed for all constructs at very high concentrations; thus the activity data used to calculate the K^n(CTP) was limited to CTP concentrations of 0–20 mM (wild type), 0–5 mM (Y173A) and 0–18 mM, (H168A and H168A, Y173A).

RESULTS

Crystallization and Structure Solution—An untagged version of CCT236 yielded native crystals grown in a condition using sodium formate as a precipitant. These crystals belong to the orthorhombic space group P2,2,2 and diffracted to 2.2 Å (Table 1). No solution to this diffraction set emerged using molecular replacement methods with GCT structures as search models. Crystals of a selenomethionine-incorporated version of CCT236 could not be reproduced in the same condition but were generated in a lithium sulfate/PEG 8000 condition. With this crystal a complete SAD dataset scaled at 3.5 Å resolution yielded the experimental phases when scaled as monoclinic P2.

The partially refined model of CCT236 in the space group P2 was used as a search model to obtain phases by molecular replacement methods for the higher resolution native P2,2,2 dataset. The asymmetric units of both crystal forms consist of identical homodimers. In the orthorhombic unit cell, the asymmetric unit consists of a single dimer, whereas in the monoclinic unit cell, the asymmetric unit consists of two dimers. Both crystals display almost identical crystal packing arrangements. The refined structure at 2.2 Å resolution shows electron density for 353 residues of the possible 472 residues comprising the homodimer, with two bound CDP-choline molecules and 249 solvent molecules (Fig. 3A). Electron density is lacking for the N...
TABLE 1
Crystallographic data collection, phasing, and refinement statistics for the catalytic domain of Rat CCTα
The data collection statistics in parentheses are the values for the highest resolution shell. Se-Met is selenomethionine.

| Crystal parameters | Native | Se-Met incorporated |
|--------------------|--------|---------------------|
| Space group | P2₁,2,2 | P2₁ |
| a, b, c (Å) | 89.0, 129.4, 43.6 | 129.2, 43.8, 88.3 |
| α, β, γ | 90.0, 90.1, 90.0° | 90.0, 90.02, 90.0° |

| Data collection statistics | | |
|-----------------------------|-----------------|
| Wavelength (Å) | 1.00 | 0.9795 |
| Resolution (Å) | 50.0–2.2 (52.8–2.20) | 50.0–3.5 (3.63–3.50) |
| Total reflections | 294,978 | 88,104 |
| Unique reflections | 26,316 | 12,964 |
| Rmerge (%) | 0.105 (0.326) | 0.163 (0.269) |
| Mean (β) (σ) | 32.7 (6.6) | 14.1 (6.8) |
| Completeness (%) | 100.0% (100.0%) | 99.8% (99.8%) |
| Redundancy | 11.2 (10.6%) | 6.4 (6.8%) |

| Phasing statistics | | |
|-------------------|-----------------|
| No. of sites | 17 (out of possible 20) |
| Overall FOM | 0.24 |
| Overall FOM (after density modification) | 0.63 |

| Refinement statistics | | |
|----------------------|-----------------|
| Protein molecules in asymmetric units | 2 |
| Residues | 353 |
| Water molecules | 249 |
| Total no. of atoms | 3176 |
| Rcryst/Rmerge | 0.22/0.27 |
| Average B-factor (all atoms) | 27.7 Å² |
| Average B-factor (CDP-choline) | 19.1 Å² |
| r.m.s.d. on angles | 2.08° |
| r.m.s.d. on bonds | 0.03 Å |

* Rmerge = \( \sum_{h,k,l} I_{hkl} - I_{hkl} \)/\( \sum_{h,k,l} I_{hkl} \), where \( I_{hkl} \) is the observed intensity, and \( I_{hkl} \) is the average intensity obtained from multiple observations of symmetry-related reflections after rejections.

** FOM is figure of merit, \((2P/P_{\alpha}^{2}+2P_{\beta}^{2})/2\), where \( P_{\alpha} \) and \( P_{\beta} \) are the observed and calculated structure factors, respectively.

d Rfree is calculated using 5% of the reflections randomly excluded from refinement.

and C termini of both chains. Chain A accounts for residues 40–215 and chain B for residues 40–216. The crystal packing contacts allow sufficient space between adjacent stacked dimers to accommodate both N- and C-terminal regions as a disordered array.

CCT236 Structure Displays an α/β-Fold with a Novel N-terminal Region—The visible portion of each monomer of the CCT236 dimer (Fig. 3A) contains the catalytic domain (C) plus 35 residues of the N-terminal region (N). Domain C (residues 75–215) conforms to the Rossmann fold, commonly found in nucleotide-binding proteins (39). Domain C is an α/β protein fold composed of five β-strands (β1–β5), and six α-helices (αA, αB, αC, αD, αL, and αE). This nomenclature strives to be compatible with the nomenclature of structurally analogous helices and strands in the previously solved structures of glycerol-phosphate CT (GCT) from B. subtilis (15, 16) and Staphylococcus aureus (40). The αL-helix in CCT236 replaces the 3₁₀ segments in GCT. The five β-strands assemble into a parallel, twisted β-sheet surrounded by the α-helices. Two α-helices (αA and αC) pack against one side of the β-sheet and two other helices (αD and αL) against the opposite side (Fig. 3B and supplemental Fig. S1). The αE-helix extends away from the fold and contacts its partner in the opposite subunit. The region of domain N visible in our structure is not an independent folding unit; rather, it is an intimate part of the CCT catalytic domain fold that crowns the top of domain C (Fig. 3A). In the description that follows, we refer to the residues N-terminal to the catalytic domain as segment N. Residues 40–75 of segment N loop around the entire top of domain C (Fig. 3A). This segment contains a short 3₁₀ helix (residues 48–50) and an α-helix (αN; residues 63–69), positioned near the N terminus of the β₁-strand of domain C. Segment N has no counterpart in GCT (Fig. 2 and Fig. 3C).

A 2-fold rotational axis runs vertically through the center of the dimer (Fig. 3A). The visible N and C termini from each chain meet their counterparts near this rotational axis. Each chain in the dimer is virtually identical in tertiary structure, and their superimposition yields a root mean square deviation (r.m.s.d.) of 0.24 Å over 174 aligned residues. The largest deviation between chain A and B occurs in a region near the N terminus (residues 50–57 in segment N), where lack of direct contact with domain C creates inherent flexibility.

FIGURE 3. Protein fold of CCT236 and comparison to GCT. The structures in this and all other figures were prepared using PyMOL. A, CCT236 homodimer. The asymmetric unit consisting of chain A (residues 40–215) and chain B (residues 40–216) is shown in ribbon representation. Segment N (40–75) is shown in blue in both chains, and domain C of chain A (76–215) and chain B (76–216) are shown in gray and wheat, respectively. B, ribbon diagram of the CCT236 monomer. The coloring of the chain transitions from N terminus (blue) to C terminus (red). The strands are numbered β₁–5 and seven α-helices as well as a 3₁₀ helix are labeled. The active site bound reaction product, CDP-choline, is shown in stick representation (black). C, comparison of CCT236 and GCT structures. A single chain (chain A; PDB code 1N1D) of GCT (magenta; residues 1–126) is shown superimposed on chain A of CCT236 (gray; residues 40–215). The CTD bound CDP-choline (black) and the GCT bound CDP-glycerol (red) are shown in stick representation. The loops in CCT236 that form the active site are labeled L₁, L₂, L₅, and L₆. * indicates the novel extension of the β₂-strand in CCT.
Both Segment N and Domain C Contribute to the CCT Dimerization Interface—Each CCT236 monomer has a surface-accessible area of \( \sim 10,500 \ \text{Å}^2 \). The dimerization interface buries 1,900 Å\(^2\) or 18% of the surface area of each monomer with nonpolar residues constituting 51% of the contact surface. Four structural elements contribute predominantly to the dimer interactions (Fig. 4) as follows: (i) residues 41–44 in segment N; (ii) helix A, the L1 loop preceding it, and the turn following it (residues 87–105); (iii) helix C and the L2 and L3 loops preceding and following it (residues 124–143); and (iv) helix \( \alpha E \) (residues 206–213). Fig. 4B illustrates the charge asymmetry and resulting charge complementarity of the dimer interface. Although there is strong subunit interaction, cross-linking studies suggest that the dimer interface undergoes rearrangement upon transition from the soluble to membrane-bound form (41). It is not known whether this effect contributes to CCT activation. Segment N contributes to dimer stabilization by forging contacts with the \( \alpha A \)-helix and L3 loop of domain C in the opposite chain. The side chains and main chain at Leu-41, Arg-42, and Gln-43 make extensive hydrogen bonding interactions with Lys-100, Asn-101, and Phe-103 at the C terminus of helix \( \alpha A \) in the opposite chain (Fig. 4C). In addition, backbone atoms at Gln-43 and Pro-44 make polar contacts with the side chains of Arg-140 and Tyr-141 in loop L3 in the opposite chain. Further details of the segment N-domain C contacts are found in supplemental Table 1.

The \( ^{140} \text{RYVD}^{143} \) motif, located in loop L3 following helix C is a key contact point in the dimer interface. This motif is a signature sequence of the CT family (15) and likely serves the role of dimer stabilization in all CT folds. The two monomer chains come within 3.6 Å at the \( \text{C}^\alpha \) of Arg-140. There are multiple inter-chain contacts in this loop, with Arg-140 as the key player. Fig. 4D illustrates a key inter-chain contact between the Arg-140 N-\( \eta_1 \) and N-\( \eta_2 \) atoms of one chain with the carbonyl oxygens at Cys-139 and Val-142 of the other chain. The side chains of Tyr-141 and His-138 also make inter-subunit contacts (Fig. 4D; details in supplemental Table 3). In addition to the RYVD motif, there are two other conserved elements in domain C that contribute to dimerization. In the conserved \( ^{85} \text{FDLFXGH}^{92} \) motif, Leu-87 and Phe-88 side chains directly face each other across
the dimer interface, but are too distant for van der Waals contact. Rather, their main chain carbonyls interact via water molecules trapped in the dimer interface (Fig. 4E). The carbonyls at Phe-88 coordinate to w10 and w10′, and the carbonyls at Leu-87 couple to w11 and w11′. Similar water molecules linking the analogous carbonyls are found in the GCT-CDP-glycerol structure (w725 and w777 in PDB code 1N1D) (16) and in ECT (PDB code 3ELB). As well, there are other ordered water molecules trapped in cavities in the dimer interface of CCT that participate in the network of inter-chain hydrogen bonds between elements of the L1 loops and between L1 and helix αC in the partnering subunit.

At the C terminus of domain C, the two αE-helices cross over at the conserved hydrophobic residues, Ile-209 and Val-210. The interacting face of helix E is lined with the hydrophobic residues Ile-206, Ile-209, and Val-210. These three residues create a sterically complementary hydrophobic pocket for the Ile-209 side chain from the opposite helix to pack against (Fig. 4F). Tyr-213 stabilizes this interaction by further packing interactions with these hydrophobic side chains. The Ile-206 side chain of chain A is packed against the aromatic ring of Tyr-213 from chain B, whereas Tyr-213 from chain A packs against Val-210 in the opposite chain. The two Tyr-213 residues occupy different rotamers at the Cα–Cβ bond and may sample these alternate conformations.

Inter-chain contacts at helices A and C constitute important subunit interaction surfaces in both CCT236 and GCT. However, the residues involved are not conserved. In CCT236, a key inter-chain helix C-helix A interaction centers on the side chain of Asp-134 in αC, which couples to the sides chains of Ser-90 and Arg-94 in αA. Another dimer contact unique to CCT is a helix E-helix C interaction involving the side chains of Arg-208, Glu-131, and the main chain carbonyl at Val-126 (Fig. 4F). This contact also serves to anchor helix αE to the main body of the domain C fold.

Domain C Fold of Mammalian CCT Closely Aligns with Bacterial GCT and Human ECT with Several Key Differences—The sequence alignment between CCT236 and other CT enzymes is shown in Fig. 2. Of these C Ts, the best characterized structurally and mechanistically is glycerol-3-phosphate cytidylyltransferase (GCT) from B. subtilis. The full GCT sequence is only 53% of the CCT236 sequence and corresponds to domain C. GCT (B. subtilis) and CCT (rat) share 34% sequence identity and 55% sequence similarity over 129 residues. The alignment of chain A of our CCT236;CDP-choline structure with chain A of the GCT;CDP-glycerol structure (PDB code 1N1D) reveals close superimposition of C-α atoms, with an r.m.s.d. of 1.55 Å over 118 aligned residues (Fig. 3C). The best superimposition is with the main chains of the β-sheets and helices A, C, and E. Segments that contain conserved motifs that forge multiple contacts with the cytidine nucleotide are closely aligned between the two C Ts as follows: the L1 loop/αA-helix that houses the 89HXGH102 motif, and helix E and the preceding RTEGISTS motif. Although single chains overlap well, superimposition of GCT and CCT dimers resulted in less accurate alignment (1.82 Å deviation of C-α atoms over 237 residues corresponding to the domain C region).

Although there is good conservation of the CCT domain C and GCT folds, there are three notable differences (Fig. 3C). The most apparent deviation is in the peripheral region between the β4- and β5-strands (residues 171–191). In GCT, the analogous region (residues 93–106) forms a short β-strand followed by two short 310 helices (15, 16). In CCT, there is a six-residue insertion (173YSSAGS178) that results in a longer extended loop (L5) followed by an α helix (αL) rather than a 310 helix. This L5 loop is the least conserved region in the CT family (Fig. 2). In addition, near the N-terminal end of the β2-strand CCT has a 2-residue insertion (Pro-104 and Asn-105) that is conserved in CCTs (Fig. 2). These two residues contribute to stable intra-chain interactions between segment N and C, as described above. The conformation of loop L2 (referred to as the “40s flap” in GCT; see Ref. 16) also differs substantially between CCT and GCT, both with bound product.

The cytidylyltransferase with the greatest sequence similarity to the CCT catalytic domain is ECT. ECTs identified to date differ from CCT and GCT in that they are monomeric with two nonidentical catalytic domains. The human ECT N-terminal catalytic domain shares 41% identity (60% similarity) with rat CCT236 over 134 residues. The C-terminal catalytic domain lacks key residues in the RTEGISTS motif and may be nonfunctional. It shares 37% identity (55% similarity) with rat CCT236 over 134 residues. After we had analyzed the structure of CCT236 in relation to GCT, the coordinates for the crystal structure of human ECT were released (PDB code 3ELB). As of this writing, no paper describing this ECT structure has been published. Alignment of CCT236 chain A with the two catalytic domains of human ECT revealed excellent superimposition (supplemental Fig. S2, C and D), with an r.m.s.d. of 1.34 Å over 126 residues for the N-terminal catalytic domain and 1.53 Å over 137 residues for the C-terminal catalytic domain. The ECT N-terminal active site is unligated, and the C-terminal active site is complexed with CMP. Like CCT, ECT lacks the two-residue insertion at the beginning of the β2-strand but has a longer L5 loop like CCT. The most interesting differences to emerge from the comparison with our CCT236 structure involve the positions of the L2, L5, and L6 loops. Their position varies depending on ligand occupancy (supplemental Fig. 2B), in support of a role for mobile flaps in the catalytic mechanism (see “Discussion”).

Product CDP-choline Is Buried in a Deep Pocket at the Base of the β-Sheet—The two active sites within the CCT dimer were readily identified by CASTp (33) as the largest pockets on the surface. Each binding pocket has a solvent-accessible surface area of 770–800 Å2 and has two arms (Fig. 5A). One arm extends from the N-terminal end of helix E up to the C-terminal base of the β-sheet, and the other arm extends along the base of the β-sheet toward a pocket bounded by the L4 and L5 loops. Each arm of the pocket is about 16 Å long, and although the first arm is exposed to solvent, the second arm appears buried. Within each active site pocket, there was a strong electron density for a bound ligand (supplemental Fig. S3A). A model for CDP-choline was fit easily into the difference density and refined. The B-factors for the CDP-choline are similar to those of the surrounding protein, suggesting a high occupancy for the CDP-choline. The bound product adopts a zig-zag conforma-
Ordered Water Molecules

Conserved Residues in the L1, L2, L5, and L6 Loops and Several

the 196RTEGISTS motif in the L6 loop. The backbone carbonyls
base is achieved primarily by direct contacts with residues from
the 196 to glutamate or Arg-113 to lysine reduces
\( k_{\text{cat}} / K_m \) by a factor of \( \sim 1000 \) (42). In
CCT, the R196K mutation reduces
\( k_{\text{cat}} / K_m(\text{CTP}) \) by a factor of \( \sim 100 \) (43).

In addition to direct interactions, the cytidine moiety is anchored by
well coordinated water molecules. Water3 links the cytosine carbonyl
oxygen to elements in \( \alpha \Lambda \) and L6. The \( \alpha \Lambda \) contacts involve the back-
bone carbonyl of Gly-91 in the
\[^{10}\]HSGH motif and the side chain carbonyl of Gln-98, whereas the L6 loop contacts involve Thr-197 side chain O-\( \gamma \) and Gln-195 backbone carbonyl. Water4 bridges the cyto-
sine carbonyl to the ribose 2'-hydroxyl and also coordinates to the
L6 loop by Thr-194 side chain O-\( \gamma \) and Gln-195 backbone carbonyl. Finally, water1 links the ribose
3'-hydroxyl to backbone carbonyls in loop L1/\( \beta \) (at Asp-82) and \( \beta 4 \) (at Ala-167). Analogous waters are
found in ECT for w1, w3, and w4, and in CCT there are water analogs
for w1 and w3.

Compared with the cytosine and ribose groups, the \( \alpha \)- and \( \beta \)-phos-
phates\(^5\) engage many more residues
via direct hydrogen bonds with multiple structural elements in
helix \( \alpha \Lambda \) and the L1, L2, and L5 loops. The O-1 exo-oxygen of the
\( \alpha \)-phosphate directly H-bonds to the N-\( e \)2 atom of His-92 in
\( \alpha \Lambda \) and the backbone NH group at Phe-85 in L1, whereas the
NH of Ile-84 coordinates both exo-oxygen atoms. The Lys-122
side chain N-\( \zeta \) in L2 bridges the \( \alpha \)- and \( \beta \)-phosphates (Fig. 5D).
Analogous direct links to the \( \alpha \)-phosphate occurring in
GCT-CDP-glycerol (PDB code 1N1D) at Lys-46 and at Thr-9
and Phe-10 (16), and in the ECT-CMP structure there are also
direct links to the \( \alpha \)-phosphate involving residues analogous to
Phe-85, His-92, and Lys-122 in L1, \( \alpha \Lambda \), and L2. Finally, there are
contacts with the \( \beta \)-phosphate unique to CCT236 involving

\(^5\) \( \beta \)-Phosphate refers to the phosphoryl group of CDP-choline that originates
from phosphocholine. We also sometimes refer to it as the nucleophile
phosphate. This nomenclature distinguishes this phosphate from the
\( \beta \)-phosphate of CTP, the other substrate.
The listed distances are averages of the values measured from each active site within the homodimer. The water molecules are labeled by the last digit of their ID numbers in the PDB file.

| Residue | Distance | GCT analog |
|---------|----------|------------|
| Cytosine-4-NH₂ | L6 2.9 | Thr-114 |
| Cytosine N-3 | L6 2.8 | Thr-117 |
| Cytosine O-2 | Water3 2.9 | |
| Water4 | 2.8 | |
| Ribose 2⁺ OH | Water4 3.0 | |
| Ribose 3⁺ OH | Water1 2.8 | |
| α-Phosphate O-1 | αA 3.4 | His-17 |
| Ile-84 N | L1 3.0 | Thr-9 |
| Phe-85 N | L1 2.8 | Phe-10 |
| Water2 | 3.2 | |
| Water5 | 2.6 | |
| β-Phosphate O-1 | Lys-122 N-ζ 3.2 | Lys-46, Lys-44 |
| Ile-84 N | 3.3 | Thr-9 |
| β-Phosphate O-2 | Lys-122 N-ζ L2 2.7 | Lys-46 |
| Tyr-173 O-η | L5 2.7 | |
| β-Phosphate O-2 | His-168 N-ε 2.6 | Trp-95 |
| Water6 | 2.6 | |

The role of His-168 and Tyr-173 in catalysis and phosphocholine binding—The ligand contacts to His-168 and Tyr-173 observed in the CCT236 structure and described above are novel to CCT. These residues are not conserved in GCTs (Fig. 2), and their role in CCT has not been tested previously. His-168 and Tyr-173 make hydrogen bonding contacts with the β*-phosphate of CDP-choline, which is the nucleophilic phosphate of the phosphocholine substrate (Fig. 1B). They are also part of the pocket complementary to the choline group. To test the hypothesis that these two residues participate in substrate binding and/or catalysis, we mutated each to alanine singly and also as a double mutation. These mutations had negligible consequence on global tertiary structure, as assessed by lack of change in the proteolytic accessibility. A time course of limited proteolysis using chymotrypsin shows essentially identical cleavage pattern for wild-type CCT236 and the three mutant proteins (supplemental Fig. S4).

The role of His-168 and Tyr-173 in catalysis is clearly seen in the plots of substrate dependence versus reaction velocity (Fig. 6). Both the phosphocholine and CTP dependences were ana-
lyzed by a sigmoidal curve fit, as the Hill coefficients were greater than 1.0 (Table 4). Positive cooperativity toward CTP has been reported previously for both C-terminal truncated mutants (47) and full-length CCTs (43). For the Tyr-173 mutant, the $V_{\text{max}}$ value (determined from the CTP dependence curves) was reduced by a factor of 20, but the $K^*$ value for phosphocholine was unaffected. For the His-168 mutant, the $V_{\text{max}}$ value was reduced by a factor of 100 and the $K^*$ value for phosphocholine increased by $\sim$8-fold. The double mutant revealed a 162-fold reduction in $V_{\text{max}}$ and a 10-fold increase in $K^*$ (phosphocholine), translating into a reduced catalytic efficiency ($V_{\text{max}}/K^*$) of $\sim$300-fold. None of the mutations significantly affect the $K^*(\\text{CTP})$ value. These results indicate that both His-168 and Tyr-173 side chains are required for optimal catalysis, whereas His-168 appears crucial for efficient phosphocholine binding.

**DISCUSSION**

This study presents the first structural analysis of an eukaryotic cytidylyltransferase, providing a view into the workings of the active site of the enzyme that controls the rate of phosphatidylcholine synthesis and helps maintain a balanced membrane phospholipid composition (38, 48, 49). The CCT236 structure, which provides a complete catalytic domain as well as an intimately associated N-terminal extension, expands the repertoire of folds within the nucleotidyltransferase family and reveals unique features that have evolved to bind and position the phosphocholine group.

**Residues 40–75 of Segment N Are an Integral Portion of the Catalytic Domain**—The N-terminal region (residues 1–75) was previously regarded as a distinct domain comparable with other domains with specialized functions. However, our crystal structure of CCT236 reveals that the C-terminal half of the N “domain” (residues 40–75) wraps around the “top” of the catalytic domain, making specific and intimate contacts with it within and between individual subunits. Although it appears to be part of the catalytic domain fold of CCT, this N-terminal extension is absent from other cytidylyltransferases, including GCT and ECT, but is present in CCTs that have regulatory tail domains, from yeast to humans (see Fig. 2). GCT and CCT have similar $k_{\text{cat}}/K_m$ values (9, 14). This argues against a participation of segment N in catalysis, and it hints at cooperation with regulatory domains. Although the N-terminal 39 residues of animal CCTs are not well conserved, sequences are conserved in the C-terminal portion corresponding to the $3_{\alpha}$ (44PAPFSDE$^{39}$) and $\alpha$-helices ($^{52}$VTMEEA$^{57}$; mammalian CCTa numbering) that contribute to segment N-domain C contacts (Fig. 2 and supplemental Table 2).

A role for segment N in stabilization of folding and dimerization is also supported by several observations in addition to the structural data provided here. First, N-terminal truncation mutants of CCT$\alpha$ missing more than 39 amino acids do not express well, and the expressed protein is insoluble. Second, a disulfide bridge readily forms across the dimer interface at Cys-37 (41), showing that this portion of segment N contacts the corresponding region in its partner subunit. Third, a yeast two-hybrid probe of dimerization domains indicated that segment N was required for stable dimer formation (41).

Residues 1–39 are not visible in the structure. Several secondary structure predictive algorithms predict no ordered structure for this region, while correctly predicting the $\alpha$-helix at residues 63–70. Other evidence in support of a flexible N terminus is that the arginine residues within the NLS are highly accessible to the protease ArgC (50). CCT$\beta$, an enzymatically active isoform of CCT, initiates at a residue corresponding to Asp-28 of CCT$\alpha$ (51), and truncation of the first 28 residues of CCT$\beta$ produces a functional CCT (52). These data emphasize the lack of a structural requirement for the beginning section of CCT.

**TABLE 4**

Kinetic parameters for wild-type CCT236 and active site mutants of CCT236

Parameters were obtained from the plots of initial velocity versus substrate concentration shown in Fig. 6, using GraphPad Prism 4 software. The $K_m$ and $V_{\text{max}}$ values are derived from the best fit to the Hill equation (see Fig. 6, legend) $\pm$ S.E. of the fit. The wild-type $V_{\text{max}}$ value from the CTP curve is more accurate, because the CTP concentration was subsaturating in the phosphocholine curves. Units for $V_{\text{max}}$ are nanomoles of CDP-choline formed (min$^{-1}$) (mg CCT$^{-1}$). Units for $K^*$ are in millimolar.

| CCT construct          | CTP dependence $V_{\text{max}}$ (nmol/min/mg) | $K^*$ (mM) | Hill coefficient | Phosphocholine dependence $V_{\text{max}}$ (nmol/min/mg) | $K^*$ (mM) | Hill coefficient |
|------------------------|-----------------------------------------------|------------|------------------|----------------------------------------------------------|------------|------------------|
| CCT236 wild type       | 3399 ± 365                                    | 7.3 ± 1.1  | 1.6 ± 0.3        | 1724 ± 108                                               | 0.33 ± 0.09| 1.4 ± 0.2        |
| Y173A                  | 167 ± 21                                      | 7.7 ± 1.2  | 1.9 ± 0.3        | 116 ± 11                                                 | 0.48 ± 0.15| 1.4 ± 0.2        |
| H168A                  | 34 ± 3                                        | 7.7 ± 1.0  | 2.0 ± 0.3        | 69 ± 12                                                  | 2.5 ± 0.6  | 1.3 ± 0.3        |
| H168A, Y173A           | 21 ± 6                                        | 8.8 ± 2.9  | 2.0 ± 0.6        | 55 ± 16                                                  | 3.2 ± 1.2  | 1.4 ± 0.4        |

6. Z. Ding and J. Lee, unpublished observations.
In all four enzymes the α-phosphate also coordinates with backbone atoms in the loop preceding the HXGH motif (L1 in the CT nomenclature). The second element resides in the loop following the β5-strand of the fold. It is a poorly conserved loop that houses the RATEGIST motif in CT's, a \( \Phi XXXXMS(K/A)S \) motif in the class I aa-tRS enzymes, and WSIFFFSS in PPAT. The boldface positions are sites of main chain direct hydrogen bonds to the cytosine or adenine base. The arginine in the CT's or the first lysine in the Trp-tRS in this motif contacts the α-phosphate of the product, and it may assist in transition state stabilization. In PPAT, a functionally analogous arginine (Arg-91) is contributed from a separate site at the end of the β4-strand. In the third feature, three amine-containing side chains coordinate to phosphate or carboxylate oxygens of the nucleophile (illustrated with different colors in Fig. 7) as follows: a backbone NH from the L1 loop (cyan), a lysine ε-amino at the C-terminal end of the second helix of the fold (magenta), and a side chain NH from a residue in the β4-strand of the fold (yellow). Finally, all four enzymes share another common feature, the participation of moving loops during a catalytic cycle, as discussed below.

**Novel Features of CCT Operate within a Conserved Architecture to Specify Recognition for Phosphocholine**—The major differences between the CCT and GCT active site architecture relate to loops L2 and L5, which provide contacts with the nucleophilic substrate. In CCT, the L2 loop backbone positions the single Lys-122 ε-amino group for contact with the nucleophilic phosphate, whereas in GCT two lysines (44 and 46) engage in this interaction. The importance of Lys-122 in CCT (and Lys-46 and -44 in GCT) is well established by mutagenesis (16, 43).

The L5 loop has an extension in CCT, found also in ECT but not in GCT (Fig. 3C and supplemental Fig. S2). The side chains in the middle of this loop \( ^{177}SSAGSDD^{180} \) do not contact CDP-choline. Rather, hydrophobic side chains on either side of it (Tyr-173, Val-181, and Tyr-182) make van der Waals interactions with the choline methyl groups. The L5 loop also positions the Tyr-173 hydroxyl for coordination with the nucleophilic phosphate in concert with His-168 at the C terminus of β4. The tyrosine functions as an analog to Lys-44 in GCT, although part of a separate loop; and His-168 is the functional counterpart for Trp-95 in GCT. Our kinetic analysis confirmed roles for His-168 and Tyr-173 in phosphocholine binding and/or catalysis. The histidine residue is buried deeper in the active site cleft and appears crucial for binding (Table 4), whereas the tyrosine likely assists in positioning the nucleophilic phosphate for attack. Thus, although the specific residues differ, both CTs feature three contact points from the L2 and L5 loops with the nucleophilic phosphate (see Fig. 7).

**Proposal for the CCT Catalytic Mechanism**—Standard kinetic analysis shows that the CCT catalytic mechanism is via a random ordered ternary complex (13), just like that of the GCT from *B. subtilis* (14), but different from mammalian ECT, which operates via a sequential mechanism (54). Mutational analysis in CCT and/or GCT reveals important catalytic roles for the two histidines in the HXGH site, Arg-196 (Arg-113) in L6, Lys-122 (Lys-44/46) in L2, and Asp-94 in GCT, corresponding to Asp-169 in β4 (13, 16, 42, 43) and His-168 and Tyr-173 in...
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β4/L5 (this work). Together with the insights from the GCT and CCT236 solved structures, we propose the following catalytic mechanism.

Substrates bind in random order. CTP binds to Arm 1 of the active site cleft (Fig. 5A) via multiple contacts in the L1 and L6 loop and Asp-169 at the start of the L5 loop. Precise contacts with the β- and γ-phosphates of CTP cannot be discerned from our structure. We speculate from its position in our CCT236 structure that ordered w7 occupies a portion of the active site designated for an exo-oxygen of the CTP β-phosphate, and from analogous interactions in GCT (15) we speculate that His-89 and Thr-202 function to bind the β- and γ-phosphates, respectively. In a further analogy to GCT, Arg-196 in L6, as well as the αE-helix dipole, may contribute electrostatic stabilization of the β- and γ-phosphates (15). Phosphocholine binds within Arm 2 of the pocket. We propose, based on analyses of human Trp-tRS (55), that the same contacts with the phosphocholine in the CCT236-CDP-choline complex are used to bind the phosphocholine substrate. Those contacts are with Asp-82, Lys-122, Trp-151, and His-168 and a complementary hydrophobic pocket for the three methyl groups contributed by Cys-113 at the base of β2 and hydrophobic side chains from L4 and L5.

The trigonal bipyrimidyl transition state at the α-phosphate is generated by surrounding the electronegative oxygens with NH groups (Fig. 5). The N-e2 of His-92 of the HXGH motif and the backbone NH at Ile-84 and Phe-85 pull on one of the exo-oxygens of the α-phosphate. Orientation of the nucleophilic phosphate is accomplished by three simultaneous contacts with His-168 N-e2, Lys-122 N-ζ, and Tyr-173 O-η. By analogy to the aa-tRS enzymes (55–58) and to GCT, this catalytic cycle may feature open and closed active site conformations created by loop movements of L2, L5, and L6 so that these binding elements can converge on their target atoms in the ligand.

Moving Loops during a Catalytic Cycle—A comparison of the crystal structures of unliganded GCT, GCT with CTP, and GCT with CTP glycerol/sulfate (15, 16, 40) point to two major mobile elements in the GCT fold that make adjustments during a catalytic cycle. First, the L6/αE segment containing the RTEGISTT motif is ordered upon binding CTP as it engages the cytosine, ribose, and phosphates at multiple points (40, 59). Second, the αB/L2 segment housing Lys-44/46 that helps orient the nucleophilic phosphate moves like a flap to close up the active site. The residues that anchor the L2 flap to the main body in GCT (Arg-55 and Tyr-52) are critical for efficient catalysis (15). The αB/L2 moving flap likely operates in CCT as well, because the opening to the choline binding pocket in CCT236 is too small for efficient entry or exit of the substrate and product. Also, the hinge-forming residues and their anchors are closely preserved in CCT and GCT. In CCT (GCT) Cys-113 (Ser-36) and Val-126 (Tyr-49) act as the hinges and are anchored to Arg-132 (Arg-55) and Asp-86 (Asp-11). An alignment of the ECT active sites with and without CMP suggests a similar movement of L2 about a hinge (supplemental Fig. 5). This alignment also provides evidence for L5 as a moving flap.

In the CMP-occupied active site, the flap is drawn closer toward the ligand to enable three residues at the N-terminal side of L5 (His-307, Gly-308, and Thr-310) to forge contacts with the ribose hydroxyls. As with the L2 loop, the movement of L5 resembles a rotation about a hinge anchored on either side of the loop. We envision that in the catalytic cycles of CCT and ECT, the L2 and L5 flap movements function to allow access to the active site, forge specific contacts with the phosphocholine, exclude water, and allow product release.

The various elements of the active site share a web of hydrogen-bonding interactions, including ordered water molecules that likely coordinate movements during a reaction cycle to generate cooperative reaction kinetics. This web of interactions is charted in supplemental Fig. S3B. Coupling of the movements of the two active sites of the CCT dimer could be attributed to the location of key active site residues in L1 and in L6 that are near or at the dimer interface, as described under “Results.”

Proposed Model for Regulation of CCT Catalysis by the M Domain—A prominent model for autoinhibition by domain M proposes direct inhibitory contacts with some part of the active site (2, 4, 9). Such a mechanism was recently elucidated for the E. coli amphitropic membrane protein, pyruvate oxidase (60). Its C terminus forms an ~23-residue amphipathic helix for membrane binding and activation, and the same segment forms an ordered structure blocking the active site and causing misalignment of a key catalytic residue in the inactive form. However, there are at least three observations that do not easily fit with this contact-inhibition model for CCT regulation. First, M domains from diverse species that exhibit lipid regulation show very little sequence conservation, an expected feature for a domain that forges specific interactions with a conserved catalytic domain. Second, the accessibility of over 30 protease sites distributed throughout domain C was not altered when comparing soluble and membrane-bound CCT (50), incompatible with a reversible docking of domain M onto C. Finally, circular dichroism analysis of the secondary structure of whole CCT and CCT fragments suggested that in the absence of lipid vesicles domain M takes on a mix of conformations, with none dominating, i.e. it appears flexible (8).

We suggest an alternative to the direct contact model, which posits that the flexibility of the domain M in the soluble form of CCT may contribute to inactivation of the enzyme by creating too much disorder in its active site. The crystallographic B-factors for the CCT236 structure (supplemental Fig. S5) suggest that although the β-sheet and helices A, C, D, and L are rigid, there is some disorder in αN, αB/L2, L5, the C terminus of L6, and especially the αE-helix. Several observations indicate that in C6 L6 and αE become more ordered commensurate with CTP binding. NMR analysis of GCT indicates that CTP binding is accompanied by a loss of conformational freedom, including at the C-terminal segment (59). In the CCT-CDP-glycerol/sulfate complex, one of the monomers of the four chain unit lacked the sulfate (mimic of pyrophosphate), and this resulted in disordering of helix E (16). In the structure of unliganded GCT from S. aureus, part of the L6 loop and all of the αE-helix are not visible (40). These findings suggest that CTP binding orders this region
and that L6/αE may be a suitable structural element for regulation. As suggested by Fong et al. (40), high flexibility of helix E could prevent binding or transition state positioning of CTP. It is worth noting that CCT236, although having lipid-independent activity, has a very high CTP Kᵣ, compared with full-length lipid-bound CCT or GCT (9). The apparent weaker affinity for CTP could be a reflection of disorder at the C terminus of the catalytic domain due to lack of tethering (e.g. to a membrane surface via domain M). Interestingly, the N-terminal catalytic domain of ECT has a very highly ordered helix E (supplemental Fig. S2), due to extensive interactions with the linker segment between the two catalytic domains. This enzyme is also constitutively active (61), which supports the idea that helix αE ordering could regulate catalytic function in CCT.

Domain M of CCT, which is thought to begin at residue ρ–234 (7, 50), is linked to helix αE via a linker region of ~20 residues. This linker region may be quite flexible as there are highly accessible primary protease sites at residues 223 and 225 (50), and because it was unresolved in the CCT236 crystal. The turnover time for fully active CCT is on the order of 50–100 ms (5, 9); thus the active site residues require 50–100 ms to carry out their ordered dance, uninterrupted by the excursions of a chaotic tail domain that likely has rotational frequencies orders of magnitude higher. Binding of the M domains to membranes reduces their conformational entropy. The ordering of the M domain would facilitate the ordering of the linker segment, helix αE and loop L6. The entropic model for CCT regulation presented here is novel, yet it can be incorporated into a model invoking inhibitory contacts, if those contacts are transient and perhaps confined to a small sub-region. Our model can be tested by spectroscopic examination of the mobility of the αE-helix and the succeeding linker segment in soluble, inactive versus membrane-bound, active forms of CCT.

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