Crystal Structure of Urea Carboxylase Provides Insights into the Carboxyltransfer Reaction*§

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Background: Urea carboxylase (UC) plays an essential role in urea utilization and belongs to the biotin-dependent carboxylase superfamily.

Results: We report the crystal structure of the Kluyveromyces lactis UC.

Conclusion: The structure provides insights into the UC carboxyltransfer reaction.

Significance: Our study sheds light on the carboxyltransfer catalysis of biotin-dependent carboxylases in general and the function of the KipA-KipI complex involved in sporulation regulation.

Urea carboxylase (UC) is conserved in many bacteria, algae, and fungi and catalyzes the conversion of urea to allophanate, an essential step in the utilization of urea as a nitrogen source in these organisms. UC belongs to the biotin-dependent carboxylase superfamily and shares the biotin carboxylase (BC) and biotin carboxyl carrier protein (BCCP) domains with these other enzymes, but its carboxyltransferase (CT) domain is distinct. Currently, there is no information on the molecular basis of catalysis by UC. We report here the crystal structure of the Kluyveromyces lactis UC and biochemical studies to assess the structural information. Structural and sequence analyses indicate the CT domain of UC belongs to a large family of proteins with diverse functions, including the Bacillus subtilis KipA-KipI complex, which has important functions in sporulation regulation. A structure of the KipA-KipI complex is not currently available, and our structure provides a framework to understand the function of this complex. Most interestingly, in the structure the CT domain interacts with the BCCP domain, with biotin and a urea molecule bound at its active site. This structural information and our follow-up biochemical experiments provided molecular insights into the UC carboxyltransfer reaction. Several structural elements important for the UC carboxyltransfer reaction are found in other biotin-dependent carboxylases and might be conserved within this family, and our data could shed light on the mechanism of catalysis of these enzymes.

In most living organisms, urea is generated in the degradition of nitrogen-containing molecules. In mammals, urea is excreted and is the main nitrogen-containing compound in the urine. Plants, fungi, algae, and bacteria possess an enzymatic activity that converts urea to ammonium, enabling them to utilize urea as a nitrogen source (1–3). In agriculture, urea is widely used as a fertilizer. In the digestive tracts of many animals, the residing microbes utilize urea from the host to produce biomass, which contributes to the host’s nutrient intake (1, 4). Two distinct enzymes catalyze the conversion of urea to ammonium: the nickel-containing enzyme urease and the biotin-dependent enzyme urea amidolyase.

Urea amidolyase is composed of urea carboxylase (UC)2 and allophanate hydrolase (AH) activities (Fig. 1A). UC converts urea to allophanate, and AH subsequently converts it to ammonium (Fig. 1B). In some organisms, UC and AH are encoded as separate polypeptides (5–8). Urea amidolyase has been identified in fungi (9, 10), algae (11), and bacteria (5, 6); recent studies revealed that it is widely distributed in fungi and bacteria (7, 8). In many of these organisms urease is not found. By adopting urea amidolyase for their urea utilization, they avoid the transition metal nickel, whose cellular level has to be tightly regulated (7). In addition to urea utilization, urea amidolyase is involved in a number of other processes. It is an essential component of a pyrimidine nucleic acid precursor degradation pathway (12). In the human pathogen Candida albicans it is required for the yeast–hyphal transition, an important mechanism to escape the host defense (13, 14).

UC catalyzes a two-step, ATP- and biotin-dependent carboxylation reaction of urea. It is composed of biotin carboxylase (BC), carboxyltransferase (CT), and biotin carboxyl carrier protein (BCCP) domains (Fig. 1A). Biotin is covalent linked to a conserved lysine residue in the BCCP domain. The BC domain carboxylates biotin with concomitant ATP cleavage, and the

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‡ The atomic coordinates and structure factors (code 3VA7) 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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2 The abbreviations used are: UC, urea carboxylase; AH, allophanate hydrolase; ACC, acetyl-CoA carboxylase; BC, biotin carboxylase; BCCP, biotin carboxyl carrier protein; BisTris, bis(2-hydroxyethyl)iminotris(hydroxymethyl)methane; CT, carboxyltransferase; KIAH, Kluyveromyces lactis AH; KlUC, Kluyveromyces lactis UC; PC, pyruvate carboxylase; PCC, propionyl-CoA carboxylase; PDB, Protein Data Base; r.m.s., root mean square.
Crystal Structure of Urea Carboxylase

CT domain subsequently transfers the carboxyl group from biotin to urea, generating allophanate (Fig. 1B). The reaction mechanism and domain organization share common features with other biotin-dependent carboxylases, including pyruvate carboxylase (PC), propionyl-CoA carboxylase (PCC), acetyl-CoA carboxylase (ACC) and 3-methylcrotonyl-CoA carboxylase (PC), propionyl-CoA carboxylase (PCC), acetyl-CoA carboxylase (ACC) and 3-methylcrotonyl-CoA carboxylase (ACC). These enzymes catalyze a variety of carboxylation reactions, playing important roles in many pathways (15, 16). Like UC, they contain BC, CT, and BCCP domains and catalyze two-step carboxylation reactions (16, 17). The BC and BCCP domains of biotin-dependent carboxylases are highly homologous, but their CT domains are distinct. Recently, high resolution crystal structures of the PC (18, 19) and PCC (20) holoenzymes have been reported, which greatly advanced the understanding of biotin-dependent carboxylases. However, the molecular mechanism for how UC recognizes the urea substrate and catalyzes its carboxylation is currently not known.

We report here the crystal structure at 2.6 Å resolution of the Kluyveromyces lactis urea amidolase UC domain (KlUC). Structural and sequence analysis indicate that its CT domain belongs to a large family of proteins with diverse functions, including the KipA–KipI complex, which has important regulatory functions in the Bacillus subtilis sporulation (21). A structure of the KipA–KipI complex is not currently available, and the UC structure provides a framework to understand the function of this complex. Most interestingly, the current UC structure captured a carboxytransfer reaction intermediate, in which the BCCP domain interacts with the CT domain and delivers biotin into its active site. This structural information and our follow-up biochemical experiments provide molecular insights into the UC carboxytransfer reaction. Analyses suggest that other biotin-dependent carboxylases might adopt similar modes of biotin coordination and CT-BCCP interactions in their carboxytransfer reaction, and our data could shed light on the mechanism of the catalysis of these enzymes.

EXPERIMENTAL PROCEDURES

Protein Expression and Purification—The UC domain of the K. lactis urea amidolase (KlUC, residues 617–1829) was cloned into vector pET28a (Novagen). The resulting recombinant protein contains a His6 tag and a thrombin cleavage sequence at its N terminus. The plasmid was co-transformed with a pTYB12 (New England Biolabs)-based plasmid, containing the Escherichia coli biotin ligase gene birA, into the E. coli strain BL21 Star (DE3). Cells were cultured in theuria broth supplemented with 50 mg/liter kanamycin, 100 mg/liter ampicillin and were induced with 0.3 mM isopropyl-β-d-thiogalactopyranoside (Bio Basic Inc.), at 20 °C for 12–16 h. 30 min before the induction, 15 mg/liter d-biotin was added to the medium. Cells were harvested by centrifugation and lysed by sonification. KlUC was purified with nickel-nitriolitracetic acid-agarose (Qiagen) and size exclusion (Sephacryl S300 HR; GE Healthcare) columns. Purified KlUC, in a buffer containing 20 mM Tris-HCl, pH 7.5, 200 mM NaCl, 2 mM dithiothreitol (DTT), and 5% (v/v) glycerol, was concentrated to 10 mg/ml, flash-cooled in liquid nitrogen, and stored at −80 °C.

The selenomethionine (SeMet)-substituted protein was produced by growing cells in the M9 medium with specific amino acids to block methionine synthesis and supplementing with SeMet (22). Its purification was the same as the native protein, except that the DTT concentration was increased to 10 mM.

Protein Crystallization—Rod-shaped crystals of KlUC were obtained with the vapor diffusion sitting-drop method at 20 °C. The reservoir liquid contains 2.4 mM ammonium sulfate and 0.1 mM BisTris, pH 7.0. Before crystallization setups, urea and ADP (2 mM each) were added to the protein solution. Crystallization of the SeMet-substituted protein was the same as the native protein, except that 10 mM EDTA (Hampton Research) was added to the protein solution before crystallization setups. Crystals were cryoprotected by equilibrating in the reservoir liquid supplemented with 20% v/v glycerol, flash cooled, and stored in liquid nitrogen.

Data Collection, Structure Determination, and Refinement—Data collection was performed at 100 K. Diffraction data were collected on an ADSC Q315 CCD detector at the Shanghai Synchrotron Radiation Facility beamline BL17U. A native data set was collected at 0.979 Å. A single wavelength anomalous diffraction data set was collected at the selenium absorption edge on a SeMet-substituted crystal. Diffraction data were processed with MOSFLM (23), integrated with SCALA (24), and the intensities were converted to structure factors with CTRUNCATE (25).

Molecular replacement calculations by MOLREP (26), with the structure of the Staphylococcus aureus ACC BC subunit (Protein Data Bank (PDB) code 2VPQ, 43% sequence identity to the KlUC BC domain) (27) as the search model, identified a solu-
TABLE 1
Data collection and refinement statistics

Numbers enclosed in parentheses are for the highest resolution shells.

| Crystal parameters          | P4322,2 | Cell dimensions (Å) |
|-----------------------------|---------|---------------------|
| a, b, c (Å)                 | 126.7, 126.7, 217.9 |
| α, β, γ (°)                 | 90.0, 90.0, 90.0 |
| Data collection             |         |                     |
| Wavelength                  | 0.979   | Resolution (Å)       |
|                             | 50.0–2.6 (2.74–2.60) | Rmerge (%) |
|                             | 9.2 (38.8) | Completeness (%)     |
|                             | 6.1 (1.9) | Redundancy (%)       |
|                             | 99.4 (99.4) | Bond lengths (Å)     |
|                             | 4.7 (4.6) | Bond angles (°)      |

Refinement

Resolution (Å) 50.0–2.6 (2.67–2.60)
No. reflections 51,949 (3,967)
Rmerge 18.9 (26.8)/26.0 (33.6)
No. atoms Protein 8,837
Ligand/ion 19
Water 444

RESULTS AND DISCUSSION

Structure Determination—UC from different organisms are highly conserved (supplemental Fig. S1). After screening through a number of organisms, we purified and crystallized KlUC (residues 617–1829 of the K. lactis urea amidolyase). The crystals belong to space group P4322,2, and contain one KlUC molecule in the asymmetric unit. The structure was determined with a combination of molecular replacement and single-wavelength anomalous scattering methods and was refined to a resolution of 2.6 Å. The refined structure agrees well with crystallographic data and expected geometric values (Table 1), with most of the residues (87.3%) in the most favored region of the Ramachandran plot and 11.4% in the additionally allowed region.

Overall Structure—The KlUC monomer adopts an elongated structure. The CT domain (residues 1069–1741) is located at the bottom, with the BC (residues 621–1068) and BCCP (residues 1760–1829) domains on top of it. In the observed conformational interactions are found between the BC and the CT domains, and the BCCP and the CT domains, but not between the BC and the BCCP domains (Fig. 2 and supplemental Fig. S2). The BC B subdomain (36) (residues 755–824) and part of the CT-BCCP linker (residues 1742–1759) are poorly defined in the electron density map and were not included in the model. Urea and ADP (2 mM each) were included in the crystallization setups. A urea molecule was found at the CT domain active site, but no electron densities for ADP were observed.

In contrast to PC (18, 19) and PCC (20), which form oligomeric holoenzymes, the functional unit of KlUC appears to be the monomer. No substantial intermolecular interactions are observed in the crystal. In our gel filtration experiments the elution volume of KlUC corresponded to a monomeric form (supplemental Fig. S8, B and C), consistent with a molecular mass of 141 kDa measured by dynamic light scattering experiments (the molecular mass of the KlUC monomer deduced from its amino acid composition is 137 kDa). Our biochemical studies indicated that this monomeric enzyme is active. Its Km for urea is comparable with that reported for the Candida utilis urea amidolyase (9), and the kinetic parameters indicate that it is more active than the Oleomonas sagaranaensis UC (5) (Table 2).

In the structure, the BC and CT domain active sites face each other and are separated by a distance of 60 Å (Fig. 2 and supplemental Fig. S2). This distance is comparable with that reported for PC (18, 19) and PCC (20) and is consistent with a model that the entire BCCP domain translocates during the catalysis. The CT-BCCP linker is partially disordered, suggesting it is flexible in nature, which could facilitate the translocation of the BCCP domain.
Crystal Structure of Urea Carboxylase

**FIGURE 2.** Overall structure of KIUC. A and B are roughly related by a 110° rotation along the vertical axis. The biotinylated lysine (Lys-1795), biotin, the urea, and water molecules found at the CT domain active site are shown in stick and ball representations. The red stars indicate active sites of the BC and the CT domains. Structural figures were produced with PyMOL.

**TABLE 2**
Summary of the KIUC kinetic parameters

| Mutation       | $k_{\text{cat}}$ s$^{-1}$ | $K_m$ for urea | $k_{\text{cat}}/K_m$ s$^{-1}$ mm$^{-1}$ |
|----------------|---------------------------|----------------|----------------------------------------|
| Wild type      | 210.8 ± 4.0 (1.0)         | 0.32 ± 0.03 (1.0) | 659 ± 74 (1.0)                          |
| D1321A         | 3.16 ± 0.20 (0.015)       | 78 ± 20 (2.4×10$^2$) | 0.041 ± 0.013 (6.2×10$^{-5}$)          |
| Y1324F         | 45.8 ± 1.2 (0.22)         | 7.7 ± 1.0 (24)   | 5.9 ± 0.9 (9.0×10$^{-2}$)              |
| D1580N         | 61.8 ± 1.8 (0.29)         | 11.3 ± 1.4 (35)  | 5.5 ± 0.8 (8.3×10$^{-2}$)              |
| K1605A         | 1.44 ± 0.06 (0.0068)      | 493 ± 91 (1.5×10$^2$) | 0.0029 ± 0.0007 (4.4×10$^{-6}$)        |
| Y1628F         | 64.9 ± 1.6 (0.31)         | 15.0 ± 1.7 (47)  | 4.3 ± 0.6 (6.5×10$^{-2}$)              |
| E1792A         | 68.7 ± 1.2 (0.33)         | 0.39 ± 0.03 (1.2) | 176 ± 17 (0.27)                        |

The CT Domain Belongs to a Large Family of Proteins with Diverse Functions—The UC CT domain does not share any sequence homology with other biotin-dependent carboxylases. Consistently, its structure is distinct from the CT domains of other biotin-dependent carboxylases (Fig. 3A). It is composed of four subdomains: A (residues 1069–1253), B (residues 1254–1415), C (residues 1416–1512), and D (residues 1513–1741). Subdomain A is a β sandwich with an α helix on one face (supplemental Fig. S3A). Subdomain C contains a five-strand antiparallel β sheet, and two α helices at one side (supplemental Fig. S3C). Both subdomains B and D adopt the cyclophilin fold (37) (supplemental Fig. S3, B and D) and can be superimposed with a r.m.s. deviation for related Ca atoms of 1.6 Å (supplemental Fig. S3E), but the sequence conservation between them is not obvious. Subdomains B, C, and D form a tight bundle. Subdomain A is located on top of the bundle, interacting mainly with subdomain B. In the structure biotin and a urea molecule are bound at the cleft between subdomains B and D (Fig. 3A).

Structural and sequence analyses suggest that the CT domain of UC belongs to a large family of proteins with diverse functions. Most prominent among these homologues is the B. subtilis KipA-KipI complex, which has important regulatory functions in the sporulation process of this organism. KipI binds to the sporulation master kinase KinA and inhibits its autophosphorylation, whereas KipA interacts with KipI and prevents it from inhibiting KinA (21). KipA is equivalent to CT subdomains A and B, with sequence identity of 35%. KipI is equivalent to CT subdomains C and D, with sequence identity of 21%. Structures of the KipI N- (38) and C- (39) terminal domains have been determined. They can be superimposed onto the KIUC CT subdomains C and D, with r.m.s. deviations for related Ca atoms of 1.6 Å (Fig. 3, B and C). Interestingly, the operon encoding KipI and KipA is regulated by the availability of nitrogen, in a manner similar to that of urea amidolyase (21). However, residues that are important for the carboxyltransfer reaction in the UC CT domain are not conserved in the KipA-KipI complex (see below), and these proteins have entirely different functions.

Although structural information is currently available for the two domains of KipI, structures of KipA and the KipA-KipI complex are not known. The structure of the UC CT domain may provide a framework to understand the function of this complex. It has been proposed that KipI dissociates from KipA to bind and inhibit KinA (38), but the molecular mechanism of this process is currently unclear. In UC the cleft between CT subdomains B and D accommodates biotin and urea. These subdomains correspond to the C-terminal domains of KipA and KipI; therefore binding of small molecule (s) at the similar location in the KipA-KipI complex could modulate the KipA-KipI interaction and might induce the formation or dissociation of the complex. It remains to be seen whether the KipA-KipI complex is regulated this way. Such a process could enable the KipA-KipI complex to sense nutrient or signaling molecule...
levels and integrate this information into sporulation regulation.

Besides the KipA-KipI complex, the CT domain is also homologous to the *Thermus thermophilus* protein TTHA0988 (PDB codes 3OPF and 3ORE) (38, 40) and a heterodimeric protein complex Msmeg0435-Msmeg0436 from *Mycobacterium smegmatis* (PDB code 3MML, Msmeg0435/0436 hereafter). The r.m.s. distances for equivalent Cα atoms between the CT domain and these structures are 1.9 and 1.6 Å, respectively, and the sequence identities are ~30%. Msmeg0435 is equivalent to CT subdomains C and D, whereas Msmeg0436 is equivalent to subdomains A and B. In contrast, the domain order of TTHA0988, C-D-A-B, is different compared with that of CT, suggesting distinct fusion events in the evolution of these proteins. The structural overlays also show that subdomain D can assume different positions in the three proteins (Fig. 3, D and E, and supplemental Figs. S4, A–D, and S5).

The functions of neither TTHA0988 nor Msmeg0435/0436 have been characterized. Both proteins are annotated as allop-hanate hydrolase but have no detectable sequence homology to the AH component of urea amidolyase. An attempt to demonstrate such an activity for TTHA0988 was unsuccessful (38, 40). UC CT domain residues important for the carboxyltransfer catalysis are not conserved in these proteins, and it is unlikely that they possess CT activities (see below). It remains to be seen whether they have functions similar to the KipA-KipI complex. It is also possible that they have yet unidentified, completely different activities.

**Structure of the CT Domain Active Site Provides Insights into Carboxyltransfer Reaction**—In the current KIUC structure, the biotinylated lysine (Lys-1795) side chain and the biotin valeric acid arm are fully extended, and the biotin head group (the tetrahydrothiophene and ureido rings) is bound at the cleft between subdomains B and D (Fig. 4A). The biotin group is well defined in the electron density map, a patch of electron density in the proximity of its N1 nitrogen was modeled as urea and water molecules (supplemental Fig. S6). Conserved residues concentrate at the cleft, suggesting that it is the CT domain active site. The structural information thus provides molecular insights into the carboxyltransfer reaction in this unique CT domain.

Highly conserved residues interact with the biotin head group and the urea molecule. The biotin head group forms three hydrogen bonds with the CT domain. Its carbonyl oxygen hydrogen bonds with the Tyr-1324 side chain hydroxyl and the Gly-1348 main chain amide, its N3 nitrogen hydrogen bonds with the Asn-1330 side chain carbonyl. The tetrahydrothiophene ring is located in a pocket formed by residues Pro-1344,
Val-1585, Pro-1634, and Gly-1635, and the ureido ring forms van der Waals interactions with side chains of Thr-1332, Phe-1349, and Ser-1609. The urea molecule is sandwiched between side chains of Val-1585 and Tyr-1637. Its carbonyl oxygen hydrogen bonds with the Asp-1584 side chain carboxyl. One of the amine groups is within hydrogen bond distance with the Tyr-1628 side chain hydroxyl and the Gly-1636 main chain amide. The amine group to be carboxylated is located 3 Å away from the biotin N1 nitrogen (Fig. 4A). An Asp-1584–Val-1585 cis-peptide bond is found at the active site.

It has been proposed that in the carboxyltransfer reaction, biotin-dependent carboxylases utilize a general base to extract a proton from the substrate atom to be carboxylated and transfer it to the biotin N1 nitrogen (17) (Fig. S7). In PC, a conserved threonine (Thr-908 in the human PC, Thr-876 in the S. aureus PC, Thr-882 in the Rhizobium etli PC) plays such a role (41, 42). In the current UC structure, the urea amine group to be carboxylated is located 3 Å away from the biotin N1 nitrogen (Fig. 4A). An Asp-1584–Val-1585 cis-peptide bond is found at the active site.

To gain further insights into the carboxyltransfer reaction, we introduced point mutations in the relevant regions and compared the activities of the wild-type protein and the mutants. All of the mutants were fully biotinylated (supplemental Fig. S8A) and behaved identically to the wild-type protein on a gel filtration column (supplemental Fig. S8B), suggesting that any change in their activity is not due to insufficient biotinylation or protein misfolding. Mutating Lys-1605 to alanine greatly reduced the activity, so did substituting its ion pair partner Asp-1321 with alanine. Other mutations in the active site, Y1324F, D1584N, and Y1628F, decreased $k_{cat}$ 3–5-fold and increased $K_m$ for urea 20–50-fold (Table 2 and supplemental Fig. S9). These data are consistent with our mechanism of the UC carboxyltransfer catalysis. That the K1605A mutant retains minimal activity suggests that direct “unassisted” proton trans-
transfer between urea and biotin might also occur (43). Our model could also explain the importance of Asp-1321 observed in the activity assays. The observed conformation might mimic a carboxyltransfer reaction intermediate, in which Lys-1605 has extracted a proton from urea, but has not protonated biotin, and the extracted proton is stored in the Lys-1605–Asp-1321 ion pair (supplemental Fig. S7). However, having not directly observed the conformation in which the general base interacts with biotin and the urea molecule, our hypotheses for the functions of Lys-1605 and Asp-1321 remain speculative. Further experiments are required to fully elucidate the mechanism of the UC carboxyltransfer reaction.

Some of the essential residues in the CT domain active site are not conserved in its homologues TTHA0988, Msmeg0435/0436, and the KipA-Kipl complex. In these proteins there is a one-residue deletion at the position corresponding to Asp-1584, Asp-1321 is not conserved (serine), neither is Tyr-1324 (threonine in TTHA0988, alanine in Msmeg0435/0436, valine in KipA) (supplemental Fig. S4E). Mutations at these locations in KIUC reduced the activity a few hundred to more than 10,000-fold (Table 2); therefore, despite the homology, these proteins are not likely to possess the CT activity.

**Function of the BCCP Domain in the CT Catalysis**—In the carboxyltransfer reaction the BCCP domain interacts with the CT domain to deliver biotin. Our structure provided a detailed picture of the CT-BCCP interactions. Excluding the biotin group, the CT-BCCP interface buries 680 Å² of surface area. The β4–β5 hairpin in the BCCP domain mediates a significant portion of the interactions. The amphipathic side chain of the biotinylated Lys-1795 is completely buried and is surrounded by the side chains of Asn-1291, Phe-1293, and Thr-1611. Hydrogen bonds are formed between its side chain amine and the Asn-1291 side chain carbonyl and the Gly-1294 main chain carbonyl, and between its main chain amide and the Asn-1289 side chain carbonyl. The side chain of the adjacent and strictly conserved Glu-1792 is also completely buried. Its carboxyl oxygen hydrogen bond with side chains of the conserved Arg-1538 and of Asn-1289 and Thr-1611 (Fig. 4B). Side chains of Tyr-1767, Thr-1768, Arg-1770, Trp-1772, and Met-1794 in the BCCP domain, and Leu-1127, Tyr-1288, Pro-1368, Arg-1538, Glu-1541, and Thr-1542 in the CT domain contribute to additional van der Waals interactions. Additional hydrogen bonds are formed between the Arg-1770 side chain guanidinium group and the Glu-1541 main chain carbonyl, and side chains of Trp-1772 and Glu-1541 (Fig. 4B).

Consistent with the structural observations, substituting Glu-1792 with alanine decreased $k_{cat}$ 3-fold, without significantly affecting the $K_m$ for urea (Table 2 and supplemental Fig. S9A). This mutation did not cause any decrease in protein biotinylation or apparent misfolding (supplemental Fig. S8, A and B). The loss of activity is presumably due to decreased CT-BCCP affinity, which would hinder biotin delivery to the CT domain active site, slowing down the carboxyltransfer reaction and in turn the overall catalysis. The function of the interactions between the BCCP and the catalytic domains has also been demonstrated by a recent study on the *R. etli* PC. Introducing the T882A mutation in this enzyme, located at its CT domain active site, increased its BC activity. Residue Thr-882 contributes to the CT-BCCP interactions in the *R. etli* PC by forming a hydrogen bond with biotin. The authors of this study propose that the T882A mutation decreases the CT-BCCP affinity and consequently biotin occupancy at the CT domain active site, which shifts the equilibrium toward increased biotin occupancy at the BC domain active site, leading to an increase in the BC activity. Consistent with this, they found that enhancing CT-BCCP affinity with small molecules in this mutant decreased its BC activity (42).

The BCCP domain might play additional roles in the catalysis. In the observed conformation, the position of the CT subdomain D is dramatically different from its counterparts in the CT domain homologues TTHA0988 and Msmeg0435/0436. Consequently, the KIUC CT domain active site cleft between subdomains B and D is much narrower compared with the equivalent regions in its homologues (Fig. 3, D and E, and supplemental Fig. S5). Such a “closed” conformation is stabilized by the BCCP domain, which interacts simultaneously with subdomains B and D (Fig. 4B). Therefore, when the BCCP domain translocates away from the CT domain during the catalysis, the CT domain might adopt an “open” conformation resembling that of TTHA0988 and Msmeg0435/0436, with a wider active site cleft. Such open/close cycles could facilitate substrate binding and release of the reaction products.

**Conserved Structural Elements in the Carboxyltransfer Catalysis of Biotin-dependent Carboxylases**—Detailed structural information of biotin and substrate coordination at the CT domain active site provides insights into the carboxyltransfer reaction. To date, such information is available for PC (19), PCC (20), and for UC from the present study. In PCC the biotin group is in a partially folded, unproductive conformation (20), whereas in PC (19) and UC it is positioned for the carboxyltransfer reaction. There are no homologies between the CT domains of PC and UC, but biotin coordination at their active site shows unexpected similarities. In both enzymes, the biotin carbonyl oxygen forms two hydrogen bonds with the CT domain. In PC, it hydrogen bonds with the side chain hydroxyl of a conserved serine (Ser-911 in the human PC, Ser-879 in the *S. aureus* PC) and the main chain amide of a neighboring lysine (Lys-912 in the human PC, Lys-880 in the *S. aureus* PC). If the biotin head groups are aligned, hydrogen bond partners of the biotin carbonyl in KIUC and PC occupy similar spatial locations (Fig. 5A). Mutations affecting these hydrogen bonds in KIUC (Y1324F) and in the *S. aureus* PC (S879A) (41) reduced the enzyme activity. These hydrogen bonds could stabilize the enolate biotin intermediate, which explains their importance in the catalysis (supplemental Fig. S7). It remains to be seen whether similar hydrogen bonds form during the carboxyltransfer reaction of other biotin-dependent carboxylases and whether they have similar functions.

During the carboxyltransfer catalysis, the BCCP domain interacts with the CT domain, delivering biotin for the reaction. In KIUC, hydrogen bonds between the conserved Glu-1792 in the BCCP domain and residues in the CT domain contribute significantly to the CT-BCCP interactions (Figs. 4B and 5B), substituting it with alanine reduced the enzyme activity. CT-BCCP interactions are also observed in the crystal structures of the human and *S. aureus* PC (19) and PCC (20). In
the human PC the glutamate is not conserved (serine) and does not form hydrogen bonds with the CT domain. In both the *S. aureus* PC and PCC, the glutamate is conserved, and potential hydrogen bond partners in the CT domain are found. In the *S. aureus* PC, the Lys-983 side chain amine is located 4.6 Å away from its carboxyl oxygen (Glu-1109, Fig. 5C) (19). In PCC, the side chain amine of Asn-311 in the β subunit is located 3.5 Å away from its carboxyl oxygen (Glu-691 in the α subunit, Fig. 5D) (20). Different side chain rotamers of these neighboring residues could reduce the distance, enabling hydrogen bond formation. The glutamate is conserved in many biotin-dependent enzymes (16), and the above analysis suggests that in these enzymes it might have a conserved function: mediating interactions between the BCCP and CT domains.

**FIGURE 5.** Common structural elements shared by UC and other biotin-dependent carboxylases. A, biotin coordination at the CT domain active site. CT domain active sites of KlUC and the *S. aureus* PC (gray) are aligned on the biotin head group and are shown in stereo. Reaction substrates, potential general bases, and hydrogen bond partners of the biotin carbonyl are highlighted. B–D, CT-BCCP interactions. Crystal structures of KlUC (B), the *S. aureus* PC (C), and PCC (D) are aligned on their BCCP domains (orange) and are shown side by side. Biotin, the biotinylated lysine, the conserved glutamate in the BCCP domain, and its potential hydrogen bond partners in the CT domain (green for the *S. aureus* PC and PCC) are highlighted.

**FIGURE 6.** The BC domain. A, the BC domain active site. Crystal structure of the *E. coli* ACC BC subunit (gray) (45) is superimposed for reference. ADP, Mg$^{2+}$ ion (gray sphere), bicarbonate (black for the carbon atom), and biotin were modeled based on the *E. coli* BC structure. Important active site residues are highlighted; residue numbers for the *E. coli* BC are enclosed in parentheses. B, conformation differences at the BC dimer interface. Crystal structure of the *E. coli* BC (36, 44) dimer is aligned for reference and is shown in different shades of gray for the two monomers. Secondary structure elements and the *E. coli* BC residue Phe-363 are indicated; those labeled with the prime sign belong to the second monomer in the dimer. The red star indicates conformational differences at the β18–β19 hairpin. C, the BC-CT interface. Residues contributing to the BC-CT interactions are highlighted. Labels for those contributing to the conserved interactions are underlined.
erved BC and BCCP domains, our analysis suggests that biotin-dependent carboxylases share homologous structural elements in biotin coordination at their CT domain active site and in CT–BCCP interactions, which might play important roles in their carboxyltransfer reaction. Further studies are required to test these hypotheses, which will advance the understanding of this important enzyme family.

The BC Domain—The structure of the BC domain is homologous to the BC components of other biotin-dependent enzymes. After structure alignment, the r.m.s. deviation for related Ca atoms in the KIUC BC domain and the BC subunit of E. coli ACC (36) is 0.97 Å. Essential residues at the BC domain active site (44, 45) are conserved in KIUC (Fig. 6A). In contrast to the E. coli BC, which form homodimers, the KIUC BC domain is monomeric. The protein surface mediating dimer interactions in the E. coli BC is not involved in protein–protein interactions in KIUC and faces a solvent channel in the crystal. Structural differences at this surface provide molecular basis for the monomeric form. KIUC contains two fewer residues in the β18–β19 hairpin, Phe-363, a major contributor to the E. coli BC dimer interactions (46), does not exist in KIUC (Fig. 6B). The Glu-23–Arg-401 intermolecular ion pair, another major contributor to the E. coli BC dimer interactions (46), is not conserved in KIUC, in which Glu-23 is replaced by an arginine (Arg-644). A similar case has recently been reported for PCC (20).

The BC domain interacts with CT subdomains A and D (Fig. 6C). The interface buries 780 Å² of surface area. It is composed of 25% nonpolar residues, 42% polar ones, and 33% charged ones (47). At the interface with the CT subdomain D, two hydrogen bonds are formed between side chains of Tyr-660 and Asn-997 in the BC domain, and Asp-1685 and Asn-1688 in the CT domain, respectively. Side chains of Tyr-660, Gln-976, and Lys-994 in the BC domain, and Leu-1650, Ile-1652, Glu-1684, and Lys-1690 in the CT domain, contribute to the van der Waals interactions. No hydrogen bonds are formed between the BC domain and the CT subdomain A. Buried surface areas are contributed by side chains of Val-967, Thr-978, Asn-1028, Tyr-1033, Ile-1036, and Asp-1040 in the BC domain, and Glu-1072, Pro-1192, Asn-1210, Val-1219, Phe-1221, and Val-1229 in the CT subdomain A (Fig. 6C). An analysis with the Consurf server (48) on 159 selected UC sequences indicated that among the BC-CT interactions, those mediated by Thr-978, Pro-1192, and Asn-1210 are conserved. These residues are located in the center of the BC-CT interface (Fig. 6C). Therefore, although in different UC the BC-CT interactions appear to be somewhat variable, the general domain architecture is likely to be preserved.

In summary, our structural and biochemical studies provided molecular insights into the UC carboxyltransfer reaction. Because some structural elements important for the catalysis might be conserved in the biotin-dependent carboxylase family, our work also shed light on the mechanism of the carboxyltransfer reaction catalyzed by biotin-dependent carboxylases in general. In addition, structure and function of the CT domain and its homologues gave yet another interesting example of diverse functions for proteins with a common fold, and our studies on UC provided a framework to understand the function of these homologues, including the KipA–KipI complex involved in sporulation regulation in B. subtilis. Furthermore, analysis of the BC domain revealed a monomeric state similar to that reported for PCC; and even though the BC-CT interactions are somewhat variable, the UC quaternary structure is likely to be conserved.

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