Structure-Function Studies with the Unique Hexameric Form II Ribulose-1,5-bisphosphate Carboxylase/Oxygenase (Rubisco) from *Rhodopseudomonas palustris*\(^*\)\(^{*}\)

Received for publication, May 7, 2014, and in revised form, June 16, 2014. Published, JBC Papers in Press, June 18, 2014, DOI 10.1074/jbc.M114.578625

Sriram Satagopan\(^1\), Sum Chan\(^1\), L. Jeanne Perry\(^2\), and F. Robert Tabita\(^1\)^\(^2\)

From the \(^4\)Department of Microbiology, The Ohio State University, Columbus, Ohio 43210-1292 and \(^8\)UCLA-Department of Energy (DOE) Institute for Genomics and Proteomics, UCLA, Los Angeles, California 90095-1570

**Background:** Rubisco fixes atmospheric CO\(_2\) to organic carbon and sustains life on earth.

**Results:** A form II Rubisco structure has been solved, and functional analysis was conducted with divergent residues.

**Conclusion:** A form II Rubisco structure has been solved, and functional analysis was conducted with divergent residues.

**Significance:** This is the first high resolution structure of an activated transition-state analog (CABP)-bound form II Rubisco.

The first x-ray crystal structure has been solved for an activated transition-state analog-bound form II ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco). This enzyme, from *Rhodopseudomonas palustris*, assembles as a unique hexamer with three pairs of catalytic large subunit homodimers around a central 3-fold symmetry axis. This oligomer arrangement is unique among all known Rubisco structures, including the form III or a form I enzyme, which varied both in length and carboxyl-terminal sequence with termini from either a form II Ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco)\(^3\) accounts for most of the biologically fixed CO\(_2\) on earth (1–3). It catalyzes the rate-limiting reaction of the Calvin-Benson-Bassham (CBB) reductive pentose phosphate pathway whereby CO\(_2\) attacks the enediol of the keto sugar ribulose 1,5-bisphosphate (RuBP) to form two molecules of 3-phosphoglycerate. There has been much interest in manipulating Rubisco because it is an inefficient biological catalyst, and the carboxylation efficiency of the enzyme is further diminished by its competition with O\(_2\). With increased potential to use Rubisco and other catalysts for enhancing carbon capture strategies and to engineer autotrophic organisms for biofuel production, it is clear that even small but incremental additions to knowledge about the catalytic mechanism of this enzyme may have a significant impact (4, 5).

There is natural variation in both structural and functional properties of Rubiscos from diverse sources (6–8). Although traditionally associated with its role in the CBB pathway for CO\(_2\) assimilation in plants and autotrophic bacteria, the involvement of archaenal Rubiscos in nucleic acid metabolism and that of the Rubisco-like proteins (RLPs) in sulfur metabolism exemplify the diverse physiological contexts in which this enzyme family may function (6). With an explosion in data obtained from “omics” approaches, it is becoming clear that several organisms utilize more than one form of Rubisco to cater to the diverse physiological needs arising from changing environmental conditions (6, 8). Because of its abundance and importance, Rubisco is one of the most highly sequenced genes (9). The rates of consumption and release of atmospheric CO\(_2\) largely determine the severity of ecological problems such as global warming, pollution, and food and energy shortages. Autotrophic organisms play a key role in maintaining this balance because of their ability to fix inorganic carbon dioxide from the environment into reduced organic carbon that is then utilized by all organisms. The enzyme ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco)\(^3\) accounts for most of the biologically fixed CO\(_2\) on earth (1–3). It catalyzes the rate-limiting reaction of the Calvin-Benson-Bassham (CBB) reductive pentose phosphate pathway whereby CO\(_2\) attacks the enediol of the keto sugar ribulose 1,5-bisphosphate (RuBP) to form two molecules of 3-phosphoglycerate. There has been much interest in manipulating Rubisco because it is an inefficient biological catalyst, and the carboxylation efficiency of the enzyme is further diminished by its competition with O\(_2\). With increased potential to use Rubisco and other catalysts for enhancing carbon capture strategies and to engineer autotrophic organisms for biofuel production, it is clear that even small but incremental additions to knowledge about the catalytic mechanism of this enzyme may have a significant impact (4, 5).

There is natural variation in both structural and functional properties of Rubiscos from diverse sources (6–8). Although traditionally associated with its role in the CBB pathway for CO\(_2\) assimilation in plants and autotrophic bacteria, the involvement of archaenal Rubiscos in nucleic acid metabolism and that of the Rubisco-like proteins (RLPs) in sulfur metabolism exemplify the diverse physiological contexts in which this enzyme family may function (6). With an explosion in data obtained from “omics” approaches, it is becoming clear that several organisms utilize more than one form of Rubisco to cater to the diverse physiological needs arising from changing environmental conditions (6, 8). Because of its abundance and importance, Rubisco is one of the most highly sequenced genes

\(^*\) This work was supported, in whole or in part, by National Institutes of Health Grant GM095742. This work was also supported by Department of Energy Grant DE-FC02-02ER63421 (to the Protein Expression Technology Center Core Facility at UCLA).

\(^3\) The abbreviations used are: Rubisco, ribulose-1,5-bisphosphate carboxylase/oxygenase; RLP, Rubisco-like protein; RuBP, ribulose 1,5-bisphosphate; CBB, Calvin-Benson-Bassham; \(K_{c,}^c\) substrate specificity; \(K_{c,}^m\) Michaelis constant for CO\(_2\); \(K_{c,}^e\) Michaelis constant for O\(_2\); \(K_{rub}\) Michaelis constant for RuBP; CABB, 2-carboxyarabinitol 1,5-bisphosphate.
with 14 unique structures and several variations also available in the protein database. Eleven of these structures are of homologous form I enzymes, all of which assemble as hexadecamers with eight large, catalytic subunits (~475 amino acids) and eight small subunits (~125 amino acids) to form an L8S8 structure. There is one form II structure (Rhodospirillum rubrum L2 dimer) and two archaeal form III structures (Pyrococcus horikoshii L8 octamer and Thermococcus kodakaraensis L10 decamer) (9). The average sequence identity between the catalytic subunits of different forms of Rubiscos is about 30%, and the in-group identity ranges from about 50 to 85% (8). The existing structures of R. rubrum Rubisco, the only form II enzyme studied in detail so far, have been useful for deducing the roles of several active site residues in the reaction mechanism (10–13). However, the identification of structural determinants of functional differences between the different forms of Rubisco has been somewhat limited by the absence of a form II structure bound to the transition-state analog 2-carboxyarabinitol 1,5-bisphosphate (CABP). CABP locks the enzyme into a “closed” conformation and positions active site residues in those conformations that likely resemble the arrangement in the enzyme bound to the carboxylation intermediate (14). By contrast, excellent transition-state analog-Rubisco structural complexes are available for form I (several) and form III (T. kodakaraensis) Rubiscos. In part, the current study was undertaken to provide the structure of a form II transition-state analog complex that might provide a further understanding as to how unique enzymatic properties attributed to form II enzymes might be resolved at the structural level.

Previous studies have shown that the genomes of Rhodobacter sphaeroides, Rhodobacter capsulatus (15), and more recently Rhodopseudomonas palustris (16) all encode for form II Rubiscos, which are homologs of the R. rubrum Rubisco, sharing about 75% sequence identity among each other. R. palustris is a metabolically versatile Gram-negative purple non-sulfur bacterium best known for its ability to switch between the different modes of metabolism that support life, thereby making this organism suitable for use in biotechnological applications like biodegradation and hydrogen production (16). The amino acid sequence of R. palustris form II Rubisco is 74% identical and 83% similar to its R. rubrum homolog. In addition to the form II Rubisco, the genome of R. palustris encodes for a form I enzyme as well as two RLPs (RPL1 and RLP2). Whereas the regulation and functions of Rubiscos (forms I and II) as part of the CBB pathway have been well studied (17, 18), the precise physiological role(s) of the RLPs is yet to be determined. The evolution of both structure and function of Rubiscos and RLPs appears to be linked (8). The form II enzyme from R. palustris was chosen for analysis as part of a broader investigation to understand the role of its Rubisco function in the physiological context. From a structural perspective, it will be useful to find out how these divergent active sites may account for functional differences between the different forms of Rubiscos and RLPs and whether Rubiscos may complement for RLP function (8).

Studies aimed at better understanding the reaction mechanism for engineering a more efficient Rubisco have relied on biological selection/directed enzyme evolution carried out with different Rubiscos in the metabolically diverse native/non-native host systems provided by Chlamydomonas, Escherichia coli, and R. capsulatus (19–21). With versatile growth capabilities, simple medium requirements, and the potential to use foreign Rubiscos for complementing autotrophic growth requirements, R. capsulatus has proven to be a valuable heterologous host for the selection and eventual identification of several residues and their functional significance (20, 22, 23). Thus, in the current study, structural analysis was combined with functional assessment using the Rubisco deletion strain of R. capsulatus complemented with wild-type and mutant enzymes of R. palustris form II Rubisco.

The x-ray crystal structure of the activated CABP-bound R. palustris form II Rubisco was solved in this study, making this the first instance of a form II-CABP enzyme-activated structural complex. Two residues near the active site were chosen for further analysis via site-directed mutagenesis based on previous studies with the R. rubrum (form II) (13, 24), Chlamydomonas (25), Synechococcus (26), and tobacco (27) form I enzymes. The carboxyl terminus, which varies both in length and composition among different Rubiscos, was also targeted for structure-function analysis by constructing chimeric enzymes. Using random mutagenesis, a second-site suppressor was isolated for a negative mutant using a previously described biological selection system (23). The structure-function analyses presented here help in better understanding the structural basis for catalysis in form II Rubisco, and these studies might prove useful for engineering a Rubisco with optimized properties for a variety of applications (4, 5).

**EXPERIMENTAL PROCEDURES**

**Plasmids, Strains, DNA Manipulations, and Culture Conditions—**Genomic DNA from R. palustris strain CA6009 was used as a template to amplify the cbbM gene, which encodes for the form II Rubisco. The amplified gene with its native ribosome binding site (15 bp upstream of the start codon) was cloned as a blunt end PCR product into the plasmid pCR Blunt II-TOPO using the Zero Blunt TOPO PCR cloning kit (Invitrogen) to generate pCR-RpccB. Site-directed mutagenesis was performed with the QuikChange kit (Agilent) using appropriate primers and pCR-RpccB as the template. Chimeric carboxyl-terminal sequences were introduced using inverse PCR with pCR-RpccB as the template. The PCR primers, which annealed to the coding sequence for the carboxyl terminus, were designed to introduce the substituted sequences in one or two successive rounds of PCRs depending on the length of the chimera. After verifying the constructs via DNA sequencing at the Plant-Microbe Genomics Facility at The Ohio State University, the exact coding regions were amplified from wild-type and mutant plasmids with primers that included an NdeI site overlapping the start codon and a BamHI site immediately 3’ of the stop codon. The 1.5-kb PCR products were digested with the restriction enzymes and cloned between NdeI and BamHI sites in plasmid pET11a or pET28a (Novagen) for recombinant protein expression in E. coli strain BL21(DE3). Use of pET28a resulted in the introduction of an amino-terminal hexahistidine tag to the various proteins, thereby facilitating protein purification via a single step nickel affinity column chromatography. All the aforementioned PCRs were carried out using

---

**Structure-Function Analysis of a Hexameric Form II Rubisco**

**Plasmids, Strains, DNA Manipulations, and Culture Conditions—**Genomic DNA from R. palustris strain CA6009 was used as a template to amplify the cbbM gene, which encodes for the form II Rubisco. The amplified gene with its native ribosome binding site (15 bp upstream of the start codon) was cloned as a blunt end PCR product into the plasmid pCR Blunt II-TOPO using the Zero Blunt TOPO PCR cloning kit (Invitrogen) to generate pCR-RpccB. Site-directed mutagenesis was performed with the QuikChange kit (Agilent) using appropriate primers and pCR-RpccB as the template. Chimeric carboxyl-terminal sequences were introduced using inverse PCR with pCR-RpccB as the template. The PCR primers, which annealed to the coding sequence for the carboxyl terminus, were designed to introduce the substituted sequences in one or two successive rounds of PCRs depending on the length of the chimera. After verifying the constructs via DNA sequencing at the Plant-Microbe Genomics Facility at The Ohio State University, the exact coding regions were amplified from wild-type and mutant plasmids with primers that included an NdeI site overlapping the start codon and a BamHI site immediately 3’ of the stop codon. The 1.5-kb PCR products were digested with the restriction enzymes and cloned between NdeI and BamHI sites in plasmid pET11a or pET28a (Novagen) for recombinant protein expression in E. coli strain BL21(DE3). Use of pET28a resulted in the introduction of an amino-terminal hexahistidine tag to the various proteins, thereby facilitating protein purification via a single step nickel affinity column chromatography. All the aforementioned PCRs were carried out using

---

**Plasmids, Strains, DNA Manipulations, and Culture Conditions—**Genomic DNA from R. palustris strain CA6009 was used as a template to amplify the cbbM gene, which encodes for the form II Rubisco. The amplified gene with its native ribosome binding site (15 bp upstream of the start codon) was cloned as a blunt end PCR product into the plasmid pCR Blunt II-TOPO using the Zero Blunt TOPO PCR cloning kit (Invitrogen) to generate pCR-RpccB. Site-directed mutagenesis was performed with the QuikChange kit (Agilent) using appropriate primers and pCR-RpccB as the template. Chimeric carboxyl-terminal sequences were introduced using inverse PCR with pCR-RpccB as the template. The PCR primers, which annealed to the coding sequence for the carboxyl terminus, were designed to introduce the substituted sequences in one or two successive rounds of PCRs depending on the length of the chimera. After verifying the constructs via DNA sequencing at the Plant-Microbe Genomics Facility at The Ohio State University, the exact coding regions were amplified from wild-type and mutant plasmids with primers that included an NdeI site overlapping the start codon and a BamHI site immediately 3’ of the stop codon. The 1.5-kb PCR products were digested with the restriction enzymes and cloned between NdeI and BamHI sites in plasmid pET11a or pET28a (Novagen) for recombinant protein expression in E. coli strain BL21(DE3). Use of pET28a resulted in the introduction of an amino-terminal hexahistidine tag to the various proteins, thereby facilitating protein purification via a single step nickel affinity column chromatography. All the aforementioned PCRs were carried out using
Structure-Function Analysis of a Hexameric Form II Rubisco

either Phusion (New England Biolabs) or PrimeSTAR GXL (Clontech) high fidelity DNA polymerase.

Random mutagenesis was carried out with negative mutant pCR-RpbbM plasmids as templates. This was accomplished using error-prone PCR amplification in the presence of 0.1 mM MnCl₂ and the enzyme Taq DNA polymerase (Invitrogen). The forward and reverse primers contained the recognition sites for the restriction enzymes KpnI and SacI, respectively. The KpnI site was designed such that the PCR product would include the native ribosome binding site for the cbbM gene. Wild-type and mutant genes were cloned into the broad host range vector pRPS-MCS3 (23) using T4 DNA Ligase (New England Biolabs). These constructs were transformed into E. coli strain S17-1 (ATCC47055) and mobilized into the Rubisco deletion strain of R. capsulatus (SB I/II”) via diparental matings to aid heterologous protein expression and functional analysis (23). The plasmid-complemented R. capsulatus colonies were selected either on rich peptone yeast extract medium (chemoheterotrophy) or on Ormerod's minimal medium plates with no organic carbon (photoautotrophic growth) as described previously (20). Liquid photoautotrophic growth curves were generated by inoculating 20 ml of Ormerod’s minimal medium with a 1:100 inoculum and bubbling illuminated cultures with a mixture of either 1.5 or 5% CO₂ balanced with H₂. The cells for the inoculum were obtained by growing single colony isolates in liquid chemoheterotrophic medium for 2 days; subsequently, the preculture cells were harvested and washed thoroughly with minimal medium. Growth was monitored by measuring the optical density of the cultures at 660 nm using a Genesys 20 visible spectrophotometer (Thermo Scientific). The cultures for protein purification were scaled up to 3 liters in transparent culture bottles with home-made fittings that enabled constant gas bubbling.

All E. coli strains were grown in Luria-Bertani medium at 37 °C using 12.5 μg/ml tetracycline or 36 μg/ml kanamycin to maintain the plasmids. For growth of R. capsulatus strains, the plates and cultures were maintained in the dark (chemoautotrophic growth) or in light (photoautotrophic growth) at 30 °C. When appropriate, the antibiotics kanamycin and spectinomycin were used at 25 μg/ml and tetracycline was used at 2 μg/ml, respectively. The plates for photoautotrophic growth selection were packed in a tightly sealed jar that contained a palladium catalyst (for removing oxygen) and CO₂/H₂-generating kits, resulting in CO₂ concentrations in the 5–10% range (Oxoid). Mixotrophic growth was assessed on rich peptone yeast extract medium with the plates incubated in tightly sealed jars containing air and a CO₂/H₂-generating kit (Oxoid).

Expression and Purification of Recombinant Proteins—For proteins expressed in E. coli strain BL21(DE3) with either pET11a or pET28a constructs of wild-type and mutant R. palustris cbbM genes, cells were routinely cultured in 500–1000 ml of LB medium. Once the cells reached an optical density of 0.4–0.6 (at 600 nm), protein expression was induced by adding isopropyl-1-thio-β-D-galactopyranoside to 1 mM final concentration. After 5–7 h, the cells were harvested, and the pellets were stored at −20 °C prior to protein purification. Prior to purification, cells were rinsed and resuspended in either a 50 mM phosphate buffer, pH 8.0 with 300 mM NaCl and 15 mM imidazole (for nickel-nitrilotriacetic acid affinity purification) or a 50 mM Tris-HCl buffer, pH 8.0 with 1 mM DTT, 10 mM MgCl₂, and 10 mM NaHCO₃ and lysed using a French pressure cell at 1000 p.s.i. (two passages). The lysates were clarified using high speed centrifugation at 4 °C.

Hexahistidine-tagged proteins were purified using one-step nickel affinity gravity flow chromatography (nickel-nitrilotriacetic acid-agarose, Qiagen). The recommended procedure (Qiagen) was followed but with a few modifications. The lysis and wash buffers were prepared with 15 and 30 mM imidazole, respectively, to reduce nonspecific binding of other cellular proteins. All the buffers also contained 20% glycerol. The imidazole in the eluted proteins was removed by dialysis against a buffer containing 20 mM Tris-HCl, pH 8.0, 10 mM MgCl₂, 300 mM NaCl, 1 mM DTT, 10 mM NaHCO₃, and 20% glycerol.

For purification of non-tagged proteins expressed in either E. coli or R. capsulatus, a two-step procedure was optimized. The first purification step utilized a Green Separopore resin (biowORLD, Dublin, OH) in which Rubisco bound to the reactive green dye was eluted using a 0–1 mM NaCl gradient in a buffer that contained 50 mM Tris-HCl, 10 mM MgCl₂, 1 mM DTT, and 10 mM NaHCO₃, pH 8.0. Fractions with Rubisco were identified via RuBP carboxylase activities as determined via radiometric assays in which the 14C from NaH14CO₃ was incorporated into acid-stable 3-phosphoglycerate (28). The Rubisco fractions were pooled, dialyzed against a no-salt buffer, and subjected to ion exchange chromatography using a 12-ml UnoQ column (Bio-Rad). Again the Rubisco fractions were identified via radiometric assays, pooled, dialyzed against a no-salt buffer, concentrated, and mixed with 20% glycerol. Dialysis and protein concentration were carried out over Amicon filters (50- or 100-kDa molecular mass cutoff; Millipore), and the protein concentrations were determined using the Bradford assay (29). Aliquots of each protein were snap frozen in liquid nitrogen and stored at −80 °C for further biochemical analysis.

Enzyme Assays and Kinetic Measurements—The kinetic constants kcat (for carboxylation), Km (for CO₂), and Ks (for O₂) were determined from simultaneous radiometric assays performed in vials flushed with either 100% N₂ or 100% O₂ as outlined previously (20). The values for K RuBP (Km for RuBP) were obtained from similar assays performed in sealed vials by varying the RuBP concentration. The substrate specificity (Ω) values were measured with purified enzymes (25–250 μg) in 400-μl assays performed in 1.5-mI vials under saturating (1.23 mM) O₂ concentrations (23). [1-14C]RuBP used in these assays was synthesized and purified as described elsewhere (30). Rates of Rubisco activation were measured with unactivated Rubisco enzymes using a modified procedure adapted from elsewhere (31). Purified enzymes (500–2500 μg) were unactivated by four to five buffer exchanges against a 100 mM HEPES-NaOH buffer, pH 8.0 using 0.5-ml Amicon Ultracel filters (50-kDa molecular mass cutoff; Millipore). The unactivated enzymes (in 500 μl) were flushed with N₂ in sealed vials and incubated with 0.5 mM RuBP. After incubation at room temperature for 1–1.5 h, a 100-μl aliquot of the enzyme-RuBP complex was treated with 400 μl of “activation” buffer comprising 100 mM HEPES-NaOH, pH 7.7, 12.5 mM MgCl₂, 25 mM NaHCO₃, 1.25 mM DTT, and 0.5 mM RuBP. The time course of activation was followed
by periodically removing 20-μl aliquots of this mixture and measuring RuBP carboxylase activities at 30 °C in 0.5-min reactions, following the stoichiometric incorporation of 14C from NaHCO3 into acid-stable 3-phosphoglycerate. The assay reaction mixtures contained 20 mM NaHCO3 (containing 8 μCi of NaH14CO3), 10 mM MgCl2, and 0.8 mM RuBP in 100 mM HEPES-NaOH, pH 8.0.

Crystallography, Data Collection, Structure Determination, and Refinement—Purified Rubisco was buffer-exchanged into 20 mM Tris-Cl, pH 8.0, 0.3 mM NaCl, 10% glycerol, 10 mM MgCl2, and 20 mM NaHCO3 using a 5-kDa Ultra-15 concentrator (Amicon), and the protein concentration was determined to be 14 mg/ml (46 nM). CABP, which had been synthesized using previously published procedures (32), was added to a final concentration of 4 mM. Seven 96-well crystal screens were set up for crystallization using the sitting drop vapor diffusion method. Drops were prepared by mixing the Rubisco-CABP solution with an equal amount of crystallization reservoir solution (varying composition). The trays were incubated at 20 °C, and crystals were observed after 2 days. The best diffracting apo crystals were grown using a crystallization reservoir solution comprising 170 mM lithium sulfate, 85 mM Tris-HCl, pH 8.5, 24% (w/w) PEG 4000, and 14% (w/v) glycerol. Similarly, good quality CABP-bound crystals could be obtained in a reservoir solution with 60 mM Tris-Cl, pH 8.5, 20% (w/w) PEG 4000, and 10% (v/v) glycerol. The crystals were harvested and swiftly passed through a cryoprotectant solution before freezing at −170 °C for data collection. The cryoprotectant solution was prepared by mixing 15 parts of 100% glycerol with 85 parts of the mother liquor. X-ray diffraction data sets were collected at beam line 8.2.1 of the Advanced Light Source at Lawrence Berkeley National Laboratory that was equipped with an ADSC Quantum 315 charge-coupled device detector. Data were collected using a wavelength of 1.000 Å at −170 °C and processed using DENZO and SCALEPACK (33). The quality of anisotropic data for the sulfate-bound crystal was improved by imposing an icosahedral resolution boundary using the UCLA MBI Diffraction Anisotropy Server.

The sulfate-bound structure (Protein Data Bank code 4LF2), which had resulted from using high sulfate concentrations in the initial crystallization conditions, was solved by molecular replacement using Phaser (34) with the structure of R. rubrum Rubisco form II (Protein Data Bank code 5RUB) as a search model. The CABP-bound structure (Protein Data Bank code 4LF1) obtained subsequently with the use of sulfate-free crystallization buffer was also solved by molecular replacement using the sulfate-bound structure as the search model. Electron densities were observed for the CABP molecule and a coordinating Mg2+ ion in all six oligomers in the hexamer. Model building was performed manually with Coot (35), and the models were refined with REFMAC (36) in which anisotropic displacement parameters were modeled with translation/libration/screw group definitions and non-crystallographic symmetry restraints. Structure validation was done using the UCLA MBI SAVES and the Protein Data Bank ADIT Servers. Subunit-subunit interface areas were calculated from accessible surfaces on each subunit that were buried in either the interface with the paired subunit of the dimer or the interface with another subunit from the neighboring dimer. The calculations were done using the program ArealMol (37, 38). Surface electrostatic potentials were calculated using the program Ezprot (39). All structure images were prepared using The PyMOL Molecular Graphics System (Schrödinger, LLC).

RESULTS

Structure Determination and Features of R. palustris Form II Rubisco—X-ray crystal structures of both the activated apo- and CABP-bound proteins were determined. The apo crystal structure was solved first and revealed two sulfate ions bound at each active site coordinated by the same residues that would interact with two phosphate groups of the CABP molecule. Attempts to obtain CABP-bound protein crystals using these crystallization conditions were unsuccessful, and in retrospect, it is obvious that the high sulfate concentration allowed sulfate ions to outcompete CABP molecules for active site occupancy. The CABP-bound crystal structure was subsequently solved from crystals obtained under phosphate- and sulfate-free conditions. Both structures show a novel hexameric arrangement in the same P1 space group with similar unit cell dimensions and an overall root mean square deviation of only 0.554 Å between the α-carbon positions of the two enzymes (Table 1). The hexameric assembly is consistent with native molecular weight determinations from size exclusion chromatography and static light scattering experiments yielding an approximate molecular mass of 300 kDa. It is thus likely that this quaternary structure arrangement is biologically relevant. The average root mean square deviation between α-carbon chains of all the dimers from both structures is 0.318 Å with a range from 0.11 to 0.45 Å. The mean value for individual subunits is 0.249 Å with a range from 0.05 to 0.45 Å. The overall arrangement of all secondary structural elements is identical in both sulfate-bound and CABP-bound structures of the activated R. palustris enzyme. Structural features around the active site region of our sulfate-bound structure are quite different from those in the previously solved sulfate-bound structure of the unactivated form I Rubisco from Galludia partita (44). As the CABP-bound structure of the activated enzyme is comparable with the reaction intermediate-bound Rubisco during catalysis, further

4 S. Satagopan and M. Arbing, unpublished results.
analysis and discussions presented below will focus on this structure. The hexamer is a trimer of homodimers with the three dimers arranged along the 3-fold symmetry axis forming a toroid tertiary structure (Fig. 1A). The two subunits within each dimer are positioned in a “head-to-tail” fashion forming two active sites per dimer with both subunits contributing to each active site, similar to what is found in other Rubisco structures (45). However, unlike the arrangement in form I (for example, \( \text{Synechococcus} \) sp. strain PCC6301 Rubisco) or form III (for example, \( \text{T. kodakaraensis} \) Rubisco), the inclination of the dimer interface relative to the cylindrical axis passing through the solvent channel is 39° to the right in the \( \text{R. palustris} \) form II

| Active-Site Ligand | Sulfate | 2-CABP |
|-------------------|---------|--------|
| PDB accession ID  | 4LF2 4LF1 |
| Metal             | Mg\(^{2+}\) Mg\(^{2+}\) |
| Off-site bound small molecules | CO\(_3\)\(^{2-}\) |

**Data collection statistics**

| Description               | Form II | Form I |
|---------------------------|---------|--------|
| Resolution range, overall (Å) | 90.00 – 1.85 | 80.00 – 2.38 |
| Resolution range, last shell (Å) | 1.92 – 1.85 | 2.47 – 2.38 |
| Space group               | P1      | P1     |

**Unit cell dimensions**

| Parameter | Form II | Form I |
|-----------|---------|--------|
| \(a\) (Å) | 74.2    | 75.0   |
| \(b\) (Å) | 99.8    | 100.7  |
| \(c\) (Å) | 100.0   | 100.7  |
| \(α\) (°) | 67.3    | 66.5   |
| \(β\) (°) | 71.3    | 108.3  |
| \(γ\) (°) | 86.6    | 95.4   |

**Radiation source**

| Source | Form II | Form I |
|--------|---------|--------|
| ALS 8.2.1 | 1.000 | 1.000 |

Statistics in parentheses were obtained after ellipsoidal truncation by the Diffraction Anisotropy Server.

**Table 1**

| Active-Site Ligand | Sulfate | 2-CABP |
|-------------------|---------|--------|
| PDB accession ID  | 4LF2    | 4LF1   |
| Metal             | Mg\(^{2+}\) | Mg\(^{2+}\) |
| Off-site bound small molecules | CO\(_3\)\(^{2-}\) |

**Diffraction Data Processing Statistics**

| Description               | Form II | Form I |
|---------------------------|---------|--------|
| Measured reflections      | 649,061 | (647,067) | 333,523 |
| Unique reflections        | 185,533 | (184,523) | 101,179 |
| Overall completeness (%)  | 85.9    | (85.4) | 98.1 |
| Last shell completeness (%)| 42.6 | (39.1) | 97.0 |
| Overall \(R_{sym}\) \(^a\) | 0.142   | (0.143) | 0.147 |
| Last shell \(R_{sym}\) \(^b\) | 0.363   | (0.483) | 0.460 |
| Overall \(I/\sigma(I)\)    | 7.9     | (7.9)  | 6.3 |
| Last shell \(I/\sigma(I)\) | 1.7     | (1.8)  | 2.1 |

**Refinement Statistics**

| Description               | Form II | Form I |
|---------------------------|---------|--------|
| \(R_{work}\) \(^c\)       | 0.227   | 0.203  |
| \(R_{free}\) \(^c\)       | 0.275   | 0.258  |
| \(R_{free}\) test set size (%) | 5 (random) | 5 (random) |
| rmsd bond length (Å)      | 0.009   | 0.010  |
| rmsd bond angle (°)       | 1.226   | 1.462  |

**Structure Validation Statistics**

| Description               | Form II | Form I |
|---------------------------|---------|--------|
| Most favored (%)          | 93.1    | 92.6   |
| Additional allowed regions (%) | 6.4 | 6.9 |
| Generously allowed regions (%) | 0.3 | 0.3 |
| Disallowed regions (%)    | 0.3     | 0.3    |

\(^a\) \(R_{sym}(I) = \frac{\sum |I_{obs} - \langle I_{sym}\rangle|}{\sum I_{sym}}\).

\(^b\) \(R_{work} = \frac{\sum |I_{obs} - F_{calc}|}{\sum |I_{obs}|}\).

\(^c\) \(R_{free} = \frac{\sum |I_{obs} - F_{calc}|}{\sum |I_{obs}|}\), where all reflections belong to a test set of 5% of randomly selected data.
enzyme (Fig. 1B). Furthermore, tertiary structure alignment of a subunit from this structure with an equivalent large subunit from other CABP-Rubisco structures shows the unique relative arrangement of amino and carboxyl termini with the termini positioned farther away from each other and pointing in different directions (Fig. 2). The closed conformation of catalytic loop 6 in the *R. palustris* structure is comparable with its position in all other analogous CABP structures. This is quite different, however, from the “open” arrangement of loop 6 in the activated, RuBP-bound structure of the homologous *R. rubrum*
enzyme (Fig. 2A). Structure alignments with other representative Rubisco sequences highlight the presence of form II-specific α helices at both the amino (αA) and carboxyl (αl and αj) termini of the R. palustris enzyme, similar to what was observed with R. rubrum form II Rubisco (Figs. 2 and 3) (45). As with the T. kodakaraensis and Galdieria enzymes, there is an α helix (α-1) preceding βC (44, 46) (Fig. 2B). However, unlike the equivalent residues from the form I and form III enzymes, residues in this region of the R. palustris enzyme are not involved in interdimer (L2-L2) interactions. Yet another distinguishing feature is the presence of a 3_{10} helix in the amino-terminal domain (Helix 0) between αC and βE. The active site residue Asn^{112} is found in this structure (Fig. 3).

It could be concluded from subunit interface analyses that the region involved in interactions with a neighboring subunit within the same dimer of the R. palustris form II Rubisco (Fig. 4A) is identical to what has been described before for other Rubiscos (9) (supplemental Movie S1). However, different forms appear to utilize different sets of regions for interactions with other neighboring subunits, including small subunits in the case of form I enzymes. The R. palustris form II enzyme uses a central core region (supplemental Movie S1), whereas the form III enzymes seem to use two flanking regions for these interactions (Fig. 4A). The form I Rubiscos utilize all these regions and a few more residues from elsewhere for the interdimer and large subunit-small subunit interactions. The interdimer interfaces of R. palustris and R. rubrum form II enzymes both have a higher concentration of positively charged residues, although the latter exists only as a free dimer. The equivalent surfaces in the form I and form III enzymes analyzed here are predominantly negatively charged (Fig. 4B).

Mutagenesis, Screening for Function, and Suppressor Selection—Comparison of all residues within 4 Å of CABP in the active sites of previously solved Rubisco structures highlights the conservation and predominantly polar nature of the active site (Figs. 5 and 6). Despite the involvement of a common set of active site residues in the reaction mechanism, there are subtle differences in both the spatial arrangement of the active site and identities of two vicinal residues between the structures of the R. palustris form II Rubisco and other homologs (Fig. 5). The active sites of sulfate-bound and CABP-bound R. palustris form II Rubisco structures are virtually superimposable with the exception of the position of residue Glu^{19}, which appears to be closer to all other active site residues in the sulfate-bound structure (Fig. 5A). A representative comparison of the CABP structure of R. palustris enzyme with its form I homolog from Synchococcus (Fig. 5A) and phylogenetic comparisons of residue identities in this region (Fig. 5B) elicit other differences. These differences could potentially explain the functional distinctness of the different enzyme forms. Although not ascribed a direct role in catalysis, Thr^{173} (Synchococcus Rubisco numbering) is two residues away from the well characterized active site Lys^{173}, and its identity is conserved as a Thr residue in all form I enzymes. However, its equivalent is an Ile in form II (Ile^{165} in R. palustris enzyme) and a Val in form III sequences (Fig. 5B). Situated on the opposite side of the active site is loop 6, which undergoes substantial conformational changes during catalysis (9). Lys^{334} (Synchococcus Rubisco numbering) from loop 6 has long been known to play a role in coordinating with the incoming CO₂ molecule during catalysis (27). The side chain of the neighboring residue (Leu^{335} in tobacco) is hydrophobic in nature and is either a Met (Met^{331} in R. palustris form II...
Rubisco) or a Leu in other Rubisco sequences (Fig. 5). Based on phylogenetic comparisons, residue Ile\(^{165}\) was changed to either Thr (as found in form I) or Val (as found in form III), and Met\(^{331}\) was changed to a Leu (as in forms I and III). A neutral substitution was also introduced in the form of an Ala at these positions. A double mutant (I165T/M331L) was also created to mimic a form I-like sequence. When identical constructs in the broad host range vector were mobilized into \textit{R. capsulatus} Rubisco deletion strain SB1/II\(^{−}\), only the wild type and mutant I165V supported autotrophic CO\(_2\)-dependent growth (Fig. 7), indicating that the rest of the mutations led to structural and/or functional defects. However, it was evident from multiple growth experiments that the mutant I165V enzyme supported significantly poorer growth relative to the wild-type enzyme.

Several studies have established the carboxyl terminus as a hot spot for affecting Rubisco catalysis (47–49). Structure comparisons and unique placement of the carboxyl terminus in the activated C4BP-bound \textit{R. palustris} Rubisco (Fig. 6) prompted further analysis of phylogenetic differences in this region. Chimeric \textit{cbbM} constructs were prepared by replacing the native \textit{R. palustris} carboxyl-terminal sequence with equivalent divergent sequences from \textit{R. rubrum} (form II), \textit{Synechococcus} (form I), or \textit{T. kodakaraensis} (form III) (Fig. 8). Functional complementation analysis in \textit{R. capsulatus} showed that only the chimeric RrCt with a form II-like carboxyl terminus could support CO\(_2\)-dependent growth and at a rate comparable with the \textit{R. palustris} wild-type enzyme (Fig. 7). After expression in either \textit{R. capsulatus} or \textit{E. coli}, equal amounts of all mutant and chimeric proteins could be purified, indicating that the substitutions did not destabilize the holoenzyme structure. Furthermore, size exclusion chromatography confirmed that the sizes of all mutant proteins were identical to that of wild-type protein (data not shown).

Random mutagenesis was carried out \textit{in vitro}, and suppressor mutations were selected with the \textit{R. capsulatus} strain SB I/II\(^{−}\)system as described previously (20). Whereas only true revertants (\textit{i.e.} T165I) and pseudo revertants (\textit{i.e.} A165V) could be obtained from Ile\(^{165}\) mutants, a second-site suppressor was isolated for the negative mutant M331A. This resulted from a mutation in the codon for Ala\(^{331}\) that substitutes it with a Val. To identify the mechanism of suppression, the mutation resulting in an A47V substitution was introduced into an otherwise wild-type gene. Unlike the double mutant A47V/M331A, which could barely support CO\(_2\)-dependent growth of the Rubisco deletion strain of \textit{R. capsulatus}, the suppressor-alone mutation could complement for near wild-type levels of photoautotrophic growth (Fig. 7). After several rounds of mutagenesis experiments, no suppressors could be isolated for mutants M331L, I165T/M331L, SyCt, or TkCt. It is thus likely that mutations introduced via the random mutagenesis technique described here were insufficient to sample the sequence space for those mutations that may be required to suppress the defects of some of the negative mutants analyzed in this study. However, these single base mutations were sufficient to result in true reversion, pseudo reversion, or second-site suppression of mutants I165T, I165A, and M331A, respectively.

**Purification and Biochemical Characterization of Wild-type, Mutant, and Chimeric Rubiscos**—Following previously published procedures for the expression and purification of common bacterial Rubisco enzymes, the wild-type and mutant proteins were initially expressed as recombinant proteins in \textit{E. coli} (20, 22, 50). The use of an amino-terminal hexahistidine fusion tag facilitated purification of all proteins using single step nickel affinity chromatography. However, upon determination of enzymatic properties from these histidine-tagged proteins, the \(K_m\) value for CO\(_2\) (\(K\)) determined for the recombinant A47V/M331A double mutant enzyme was substantially high (~1500 mM) and failed to explain why it was recovered as a suppressor (Fig. 7). A reciprocal experiment was conducted in which the \textit{R. capsulatus} Rubisco deletion strain was complemented with the histidine-tagged A47V/M331A double mutant gene. However, this construct failed to support CO\(_2\)-dependent autotrophic growth, further confirming that the histidine tag was possibly interfering with the function of the enzyme. This prompted reanalysis of all enzymatic properties with untagged proteins that had been found to complement \textit{R. capsulatus} strain SB I/II\(^{−}\)for autotrophic growth. A two-step protein purification starting with large batches of autotrophically grown \textit{R. capsulatus} yielded proteins that were sufficiently pure (about 75%) to facilitate further analysis. The autotrophic growth requirement for Rubisco expression precluded the purification of negative mutant enzymes from \textit{R. capsulatus}. However, they were purified as recombinant proteins from \textit{E. coli} after overexpression from respective pET11 (non His-tagged) constructs following isopropyl \(\beta\)-\(D\)–galactopyranoside induction. The properties of the wild-type enzyme, which was used as a control for \textit{E. coli} expression, as well as the other enzymes that had been purified from \textit{R. capsulatus} were identical to those of the respective recombinant proteins purified from \textit{E. coli}. Whereas the specific activities of \textit{R. rubrum} and \textit{R. palustris} wild-type and the chimeric RrCt enzymes were all in the range of 2–3 \(\mu\)mol of CO\(_2\) fixed/min/mg, mutant I165T/M331L and chimeric SyCt and TkCt enzymes had insignificant levels of activity, precluding them from further analysis. The specific activities of I165V (about 30% of wild type) and all other mutant enzymes (10–15%) were sufficient for the determination of enzymatic properties.

The \(\Omega\) values determined from HPLC fractionation of tritium-labeled carboxylation and oxygenation products were identical for the wild-type form II enzymes (\textit{R. palustris} and \textit{R. cap}

**FIGURE 3.** Structure-based sequence alignment of catalytic large subunits from \textit{R. palustris} (\textit{Rp}) form II, \textit{R. rubrum} (\textit{Rr}) form II ([Protein Data Bank code 9RUB]), \textit{Synechococcus} (\textit{S. elongatus} (\textit{Se}) form I ([Protein Data Bank code 1RBL]), spinach (\textit{S. oleracea} (\textit{So}) form I ([Protein Data Bank code 8RUC]), \textit{P. horikoshii} (\textit{Ph}) form III ([Protein Data Bank code 2D69]), and \textit{T. kodakaraensis} (\textit{Tk}) form III ([Protein Data Bank code 3A12]) enzymes. The secondary structural assignments for the \textit{R. palustris} enzyme are displayed above the sequences using a schematic that is consistent with that of Fig. 28. The asterisks below the sequences designate strictly conserved residues among known Rubiscos, and the inverted triangles above the sequences indicate the positions of the active site residues. Residues involved in interactions with the dimer-forming neighboring subunit are colored green; the highlighted residues (in salmon color) are involved in interactions with neighboring subunit(s) other than the dimer partner. The dotted boxes indicate contiguous regions in different Rubiscos that appear to be involved in interactions with a neighboring subunit, excluding its dimer partner, as also highlighted in the space-filling surface representations (Fig. 44).
rubrum), chimeric RrCt enzyme, and mutant A47V enzyme (Table 2). The $k_{cat}$ value (for carboxylation) measured for the chimeric RrCt enzyme was 37% higher relative to the value for R. palustris wild type and 55% higher than what was measured for the R. rubrum wild-type enzyme. Mutant enzymes I165V, A47V, and A47V/M331A all have lower $k_{cat}$ values, ranging from 27 to 5% of the wild-type value (Table 2). With all other properties comparable with the wild type, the improved $k_{cat}$ of
the chimeric RrCt enzyme does not appear to confer a discernible growth advantage to the *R. capsulatus* host strain under the conditions reported here (Fig. 7 and Table 2). Comparable values for \( \Omega \) and \( K_m \) (for CO\(_2\) and RuBP) and an 82% lower \( k_{\text{cat}} \) possibly explain the moderate effect in growth complementation observed for the mutant A47V. Substantially lower values of \( \Omega \) and higher values of \( K_m \) (for CO\(_2\) and RuBP) likely explain the slower growth conferred by mutants I165V and A47V/M331A (Fig. 7). The higher \( K_m \) values for O\(_2\) (\( K_o \)) suggest that these mutant enzymes are less inhibited by O\(_2\). With only 4% of the wild-type value of \( k_{\text{cat}} \), it is not surprising that the A47V/M331A suppressor mutant confers the longest lag and slowest growth among all the mutants that can complement for Rubisco function. With such poor enzymatic properties, it seems likely that this suppressor may not have been identified if selection had been carried out under ambient CO\(_2\) concentrations. In fact, when autotrophic growth of the mutant A47V/M331A was monitored in liquid medium bubbled with either 5 or 15% CO\(_2\), the measured lag time increased from \( \sim 100 \) to \( \sim 150 \) h. There was very little difference in the lag times measured for the wild type and other mutants (data not shown).

The mutant enzymes, which failed to complement *R. capsulatus* for CO\(_2\)-dependent autotrophic growth, all had varying levels of detrimental changes to the catalytic properties. Barring the mutant I165A enzyme, which still retained 16% of the wild-type level of \( k_{\text{cat}} \), the other mutant enzymes retained only 4–6% of the wild-type \( k_{\text{cat}} \). Although the \( \Omega \) values of mutant enzymes I165T and M331L were comparable with the wild-type value, the \( K_m \) (I165T, 218 \( \mu \)M; M331L, 322 \( \mu \)M) and \( K_{\text{RuBP}} \) (I165T, 58 \( \mu \)M; M331L, 320 \( \mu \)M) values for these enzymes were both substantially higher. Mutant enzymes I165A and M331A both had \( \Omega \) values of 8. The lower \( \Omega \) value combined with a \( k_{\text{cat}} \) value of only 0.2/s and high substrate \( K_m \) values (\( K_c \), 345 \( \mu \)M; \( K_{\text{RuBP}} \), 115 \( \mu \)M) possibly explains the negative phenotype attributable to the mutant M331A. However, with superior values for \( k_{\text{cat}} \) (0.8/s), \( K_c \) (151 \( \mu \)M), and \( K_{\text{RuBP}} \) (13 \( \mu \)M) relative to those measured...
for the suppressor mutant A47V/M331A enzyme (0.2/s, 433 μM/ and 20 μM, respectively), it was surprising that the mutant I165A confers a “negative” phenotype in *R. capsulatus* (Fig. 7 and Table 2).

Time courses for activation were measured with the intent to explain the possible differences in the phenotypes conferred by mutants I165A and A47V/M331A. Sugar phosphates, including RuBP, are known to bind the unactivated form of Rubisco (i.e. in the absence of CO₂ and Mg²⁺), thereby forming an inactive complex *in vivo* (51). Reactivation of the enzyme involves release of the inhibitor from the active site followed by the carbamylation of an active site Lys residue and coordination with Mg²⁺. The tightness of inhibitor binding influences the rate of enzyme activation *in vivo* and is a well known mechanism to
regulate Rubisco activity in various organisms (51). Upon inhibition of the unactivated form of the enzyme by RuBP, the rates of reactivation were measured for wild-type and mutant I165A and A47V/M331A enzymes. The mutant I165A enzyme was indeed the slowest to be reactivated when compared with the wild-type and mutant A47V/M331A enzymes (Fig. 9). Initial slopes from these activation profiles were used to calculate levels of activation per unit time for each enzyme (Fig. 9). The calculated values were 63%/min for the wild type, 14%/min for the double mutant A47V/M331A, and 5%/min for the mutant I165A. The rapid reactivation of the wild-type enzyme indicates weak inactivation by RuBP binding. Unlike eukaryotic and cyanobacterial Rubisco enzymes, which are known to interact with accessory proteins to relieve sugar phosphate inhibition in vivo (31, 51), no such regulatory proteins have been identified in R. palustris or R. capsulatus. It might be tentatively concluded that inactivation of the wild-type enzyme by RuBP (and potentially other sugar phosphates) is probably too weak to have a potential regulatory role in vivo. However, detrimental changes to the catalytic properties combined with tight binding inhibition caused by various sugar phosphates could account for the negative phenotype conferred by the mutant I165A enzyme and the poor but positive complementation conferred by suppressor mutant A47V/M331A (Fig. 7). To summarize, although the residues Ile165 and Met331 have not been ascribed direct roles in catalysis, even subtle alterations to the side chains of these residues appear to cause significant changes in the functional properties of the enzyme.

**DISCUSSION**

Distinguishing Features of the CABP-bound R. palustris Form II Rubisco Structure—The form II Rubisco structure solved as part of this study allows for a direct comparison with previously described CABP structural complexes of form I and form III Rubiscos. Besides the unique placement of the carboxyl terminus in the R. palustris form II enzyme, which could not be deduced from the previously solved structures of R. rubrum Rubisco, the surface areas involved in the interdimer interactions and the nature of the constituent surface residues are quite different among the three forms of Rubisco (Figs. 3 and 4 and Table 3). Structural alignments show that the residues involved in the interdimer interactions in the R. palustris form II Rubisco are indeed similar to those in the R. rubrum form II Rubisco (Figs. 2 and 3). However, subtle differences in residue identities and the consequential interactions involving at least two surface regions potentially explain the higher order hexameric association that is observed only with the R. palustris enzyme. In the R. palustris structure, the residue Arg149 in Lys457 is stabilized by van der Waals interactions with Asp146 from the same chain and carbonyl oxygen of Leu359 from the adjacent dimer (Fig. 10). The R. rubrum enzyme has a more flexible Lys148 in place of Arg149 of the R. palustris enzyme; likewise Ala145 is found in place of the Asp146 residue of the R. palustris...
enzyme. Although the carbonyl oxygen of the \( R. \) \( \text{rubrum} \) Gly\(^{358} \) could serve the same function as the equivalent oxygen of Leu\(^{359} \) in \( R. \) \( \text{palustris} \), any potential interaction to form a higher order structure would be sterically precluded by the side chain of Glu\(^{154} \) in the \( R. \) \( \text{rubrum} \) enzyme (Fig. 10C). Furthermore, the \( R. \) \( \text{palustris} \) structure has an intricate network of hydrogen bonds between adjacent chains involving three residues from the loop between \( \alpha 4 \) and \( \beta 5 \). The side chain of Gln\(^{283} \) is stabilized by two carbonyl oxygen atoms from Ala\(^{256} \) and Ile\(^{259} \) of the same chain and by hydrogen bond interactions with the guanidyl group of Arg\(^{99} \) from the adjacent dimer. The carbonyl oxygen of the adjacent Lys\(^{282} \) is also within hydrogen bond distance from the guanidyl group of Arg\(^{99} \). This network is further stabilized by a water-mediated interaction of the side-chain amino group of Lys\(^{282} \) with the same Arg\(^{99} \) (Fig 10C). Such an intricate network seems unlikely with the \( R. \) \( \text{rubrum} \) structure where the equivalent of \( R. \) \( \text{palustris} \) Gln\(^{283} \) is a similar but shorter Asn\(^{282} \). Likewise, the \( R. \) \( \text{rubrum} \) equivalents of Lys\(^{282} \) and Arg\(^{99} \) are Asp\(^{281} \) and Lys\(^{98} \), respectively (Fig. 10C). In place of the four hydrogen bonds in \( R. \) \( \text{palustris} \), only a single hydrogen bond would be formed between the side chains of Asp\(^{281} \) and Lys\(^{98} \) if the \( R. \) \( \text{rubrum} \) subunits assembled into a higher order oligomer. This is further substantiated by additional surface analysis. Subunits A and F from the CABB-bound \( R. \) \( \text{palustris} \) structure that contribute to one of the three interdimer interfaces were used as the reference (4LF1_A-F). Subunit A from the \( R. \) \( \text{rubrum} \) dimer structure (Protein Data Bank code 9RUB) was aligned with 4LF1_A and then combined with another 9RUB subunit, either chain A or chain B, which was aligned with 4LF1_F. This generated two models (9RUB_A-A and 9RUB_A-B) that resemble the interdimer interface in 4LF1_A-F. Shape complementarity analysis gave a value of 0.538 for the 4LF1_A-F interface and much lower values of 0.231 and 0.299 for 9RUB_A-A and 9RUB_A-B interfaces, respectively. Higher values are indicative of higher interface-shape complementarity with most highly complementary protein interfaces receiving on average a value of 0.53 (42). Protein Interfaces, Surfaces and Assemblies (PISA) analyses estimated an interface solvation free energy (\( \Delta G \)) of \(-4.7 \text{ kcal/mol} \) for the model 4LF1_A-F and 0.88 and 0.78 kcal/mol for models 9RUB_A-A and 9RUB_A-B, respectively. Higher interface affinities typically result in more negative \( \Delta G \) values. Positive values obtained for the \( R. \) \( \text{rubrum} \) models indicate that the interactions between the constituent subunit surfaces are too weak to allow for stable interface formation.

**R. palustris Rubisco as a Model Enzyme for Carrying Out Structure-Function Analyses**—The form II Rubisco from \( R. \) \( \text{rubrum} \) has been the preferred model enzyme for discerning the role of several active site residues in Rubisco catalysis (10–13). The activated CABB-bound structure of the \( R. \) \( \text{palustris} \) form II Rubisco now allows for some of these studies to be carried out with this enzyme. Moreover, the gene encoding the \( R. \) \( \text{palustris} \) Rubisco is conveniently expressed in the \( R. \) \( \text{capsulatus} \) strain SB I/II\(^-\) host, allowing one to combine biological selection methodologies with protein structure-function studies. We report here one such analysis in which divergent residues near the active site and the carboxyl termini have both been targeted. Most substitutions at Ile\(^{165} \) and Met\(^{311} \) of the \( R. \) \( \text{palustris} \) Rubisco result in detrimental changes to both the enzymatic properties (Table 2) and the ability to complement for Rubisco function (e.g. \( \text{CO}_2 \)-dependent growth) in \( R. \) \( \text{capsulatus} \) strain SB I/II\(^-\) (Fig. 7). Although these two residues have not been ascribed direct roles in the reaction mechanism (52), the results presented here underline their importance for catalysis in the context of the form II active site. This has been partially demonstrated in previous studies in which the \( R. \) \( \text{rubrum} \) equivalent of Ile\(^{165} \) was changed to Thr, Asn, or Asp. All substitutions led to substantial decreases in \( k_{\text{cat}} \) and increases in

### Table 2: Kinetic properties of recombinant \( R. \) \( \text{palustris} \) Rubisco enzymes purified from either autotrophically grown \( R. \) \( \text{capsulatus} \) or E. coli

| Enzyme      | \( k_{\text{cat}}^a \) | \( K_a^a \) | \( K_i^a \) | \( K_i/K_a^a \) | \( K_{\text{RUBP}}^a \) |
|-------------|---------------------|------------|------------|----------------|------------------|
| Wild type   | 4.9 ± 0.2           | 4.1 ± 0.2  | 277 ± 15   | 2.7            | 10 ± 2           |
| A47V       | 0.9 ± 0.2           | 333 ± 61   | 82 ± 31    | 0.8            | 6 ± 2            |
| A47V/M331A | 0.2 ± 0.1           | 178 ± 47   | 391 ± 45   | 2.2            | 35 ± 9           |
| I165V      | 1.3 ± 0.2           | 111 ± 28   | 270 ± 71   | 2.4            | 8 ± 2            |
| Rtc        | 6.7 ± 0.7           | 125 ± 12   | 143 ± 3    | 1.1            | 5 ± 2            |
| RcpbM WT   | 4.3 ± 0.5           |           |           |                |                  |

*Values are the means ± S.D. (n = 3) of at least three separate enzyme preparations.

*Calculated values.

**FIGURE 9.** Time courses for Rubisco activation measured from purified recombinant wild-type and mutant I165A and A47V/M331A enzymes. Percentages of maximal specific activity (measured with respective enzymes that had been fully activated) were plotted against the time allowed for activation after being inactivated by the formation of the enzyme-inhibitor complex. The maximal specific activity values measured were 2.2 \( \mu \)mol/min/mg for the wild type, 0.24 \( \mu \)mol/min/mg for mutant I165A, and 0.05 \( \mu \)mol/min/mg for mutant A47V/M331A. Data are representative of two identical experiments carried out with two separate enzyme preparations, which gave similar results.
but there was no convenient way to assess the functionality of these mutants in vivo (13). The reduced catalytic efficiency of *R. palustris* mutant I165T enzyme described in this study is consistent with what was measured for the *R. rubrum* mutant I164T enzyme. However, unlike the *R. rubrum* equivalent, there is no change in the specificity factor of the mutant I165T relative to the wild type (Table 2). In another study, mutant screening in *Chlamydomonas* identified a missense mutation at the equivalent position (residue 173) resulting in a Thr to Ile change. This led to complete loss of carboxylase activity (25). Because reciprocal substitutions at this residue in form I (*Chlamydomonas*; from Thr to Ile) and form II (*R. palustris*; Ile to Thr) led to loss of function, it must be concluded that other divergent residues complement for different requirements in the different enzyme forms. It appears that in form II enzymes an Ile at this position helps create a hydrophobic environment, which allows for seamless electron transfers mediated by the neighboring active site residues (Fig. 6). A Val substitution is thus expected to retain a similar steric and hydrophobic environment, leading to a milder impact on the catalytic activity as seen by the fact that this mutant enzyme retained functional competency. Nevertheless, the negative changes to the enzymatic properties measured for the mutant I165V enzyme suggest that the side-chain length in Ile may be optimal for this enzyme. Conversely, substitution with an Ala would create a void that may be large enough to allow an extra water molecule, thereby disrupting the balanced electrostatic environment. It is thus not surprising that the mutant I165A enzyme lacks functional competency.

It was surmised that Met331, another divergent residue positioned on the other side of CABP (Fig. 6), could compensate for a change at Ile165. However, neither the M331L single mutant nor the I165T/M331L double mutant enzyme was able to complement for function in vivo. Clearly, there must be other subtle differences in the form I active site that would allow Thr173 (equivalent to Ile165) and Leu335 (equivalent to Met331) to function optimally at their respective positions. The substitution analogous to M331L in the *R. rubrum* enzyme (M330L) also resulted in similar changes to the kinetic constants, i.e., no change in *K₉₀₂₄*, reduced *kₖₐₜ*, and higher *K₉₃₅* values for RuBP, CO₂, and O₂ (24). Lys330 and Met331 are located in loop 6 (residues Gly324–Gly333), which closes or “plugs” the active site in conjunction with residues from a neighboring subunit of the homodimer (26) (supplemental Movie S2). This is also evident from various Rubisco structures in which the entire loop is disordered in the absence of an active site ligand. However, these two residues are part of the most conformationally stable region of this flexible loop in the closed conformation as indicated by the temperature factors in the crystal structures. Lys330 plays a direct role in catalysis, and hence no substitutions are tolerated at this position (26). Conformational stabilization of Met331 is achieved via van der Waals interactions with residues Gly324–Gly333, which closes or “plugs” the active site in conjunction with residues from a neighboring subunit (Fig. 6). The form I equivalent, Leu335, in the *Synechococcus* 6301 Rubisco is likewise stabilized by interactions with Gly126, Phe127, and Lys128. Lys128, which is conserved in both forms I and III, is believed to stabilize the closed conformation by hydrogen bonding to the main chain carbonyl groups of residues in both loop 6 and the carboxyl terminus (Fig. 6). It has been shown before that activation and ligand binding in form I Rubisco cause decreases in the affinity for RuBP and *kₖₐₜ* and resulted in increased synthesis of a side reaction product (9). It has been shown before that activation and ligand binding in form I Rubisco cause decreases in the affinity for RuBP and *kₖₐₜ* and resulted in increased synthesis of a side reaction product (9).

**TABLE 3**

Comparison of buried intradimer (highlighted in green color) and interdimer (highlighted in salmon color) interaction surface areas calculated from available structures of various Rubisco enzymes in the Protein Data Bank

| PDB ID   | Source organism | Organism type | Form | Biological arrangement | Relevant ligands | Accessible surface area buried in the interface between subunits in a dimer (Å² per dimer) | Accessible surface area buried in the interface between a dimer and its neighboring subunit (Å² per dimer) |
|----------|-----------------|---------------|------|------------------------|------------------|-----------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------|
| 1BWV     | Galdieria partita | Red alga      | I    | (L₅S₅)₆                | CABP, Mg²⁺       | 4761                                                                                           | 7590                                                                                           |
| 8SRU     | Spinacia oleracea | Plant         | I    | (L₅S₅)₆                | CABP, Mg²⁺       | 4757                                                                                           | 8858                                                                                           |
| 1LRC     | Nicotiana tabacum | Plant         | I    | (L₅S₅)₆                | CABP             | 3080                                                                                           | 8048                                                                                           |
| 4HHH     | Pismum sativum   | Plant         | I    | (L₅S₅)₆                | RuBP             | 4684                                                                                           | 9051                                                                                           |
| 1WDD     | Orzya sativa japonica | Plant        | I    | (L₅S₅)₆                | CABP, Mg²⁺       | 4713                                                                                           | 8619                                                                                           |
| 1BXN     | Ralstonia eutropha | Bacteria      | I    | (L₅S₅)₆                | Phosphate        | 3834                                                                                           | 7686                                                                                           |
| 1GK8     | Chlamydomonas reinhardtii | Green alga | I    | (L₅S₅)₆                | CABP, Mg²⁺       | 4756                                                                                           | 9133                                                                                           |
| 1RLB     | Synechococcus elongatus | Bacteria    | I    | (L₅S₅)₆                | CABP, Mg²⁺       | 4815                                                                                           | 7452                                                                                           |
| 3ZXW     | Thermosynechococcus elongatus | Bacteria | I    | (L₅S₅)₆                | CABP, Mg²⁺       | 4793                                                                                           | 6229                                                                                           |
| 4LF1     | Rhodospseudomonas palustris | Bacteria | I    | (L₅S₅)₆                | CABP             | 4232                                                                                           | 2131                                                                                           |
| 9RUB     | Rhodospirillum rubrum | Bacteria   | I    | (L₅S₅)₆                | RuBP, Mg²⁺       | 3461                                                                                           | Not applicable for a free dimer                                                                |
| 2D69     | Pyrococcus horikoshii | Archaea     | III  | (L₅S₅)₆                | Sulfate          | 3518                                                                                           | 2691                                                                                           |
| 3A12     | Thermococcus kodakaeensis | Archaea    | III  | (L₅S₅)₆                | CABP, Mg²⁺       | 4387                                                                                           | 2085                                                                                           |
Leu, it was possibly easier to obtain a suppressor for M331A and not M331L. The α-carbon of Ala47 is ~15 Å away from Met331 in the crystal structure. Its side chain is in a hydrophobic environment within 4 Å of Ala70 and Val72. Substitution with a bulkier Val would be expected to push Ala47 with A47V toward the active site. Glu49 of A47V functions to stabilize Lys330 and assists in loop 6 closure during catalysis, and Thr54 in the adjoining loop is involved in binding the P1 phosphate of the substrate (53). Hence, the movement of αβ likely nudges these catalytically important residues along with Asn112 closer to the active site (Fig. 6). This could also restore steric shielding around the active site that was lost when Met331 was substituted with a smaller Ala (supplemental Movie S2). Together, these structural changes likely account for the increased affinity for RuBP as well as the lower $K_c$ value caused by an A47V substitution, which resulted in suppressing the negative effect of M331A in vivo (Table 2). This was further confirmed by in silico molecular dynamics simulations with a modeled structure of the A47V/M331A double mutant, which showed an increased distance between Ala70 and Val47 and decreased distances between the active site residues Thr54 and Asn112 from the side chain of Lys330 (data not shown).

Analysis of Carboxyl-terminal Chimera Accentuates the Importance of Carboxyl Terminus as a Potential Target for Enhancing Rubisco Catalysis—The most distinctive feature of the $R. palustris$ form II-CABP structure is the position of its carboxyl terminus.
carboxyl terminus, i.e. pointed away from the active site region and with no contacts to the loop 6 residues (Fig. 6). When CABP is bound to the active site, the carboxyl terminus is folded over loop 6 in all other Rubisco structures available in the database. In form I Rubisco, the loop 6/carboxyl terminus arrangement has been linked to the regulation of active site opening and closure and is thus believed to be critical for optimal rates of catalysis (53). However, disruption of these interactions via substitutions at a conserved carboxyl-terminal residue could still be tolerated for the function of the enzyme in vivo in Chlamydomonas (49). When the active site is unliganded or occupied by loosely bound substrates, the carboxyl terminus is either disordered or pointed away from loop 6 (3, 7). The presence of α1 in both R. palustris and R. rubrum form II enzymes and α1 in the former likely constrains this region from taking up an extended conformation on top of loop 6 (Figs. 3B and 6). The ability of the RrCt chimera to support wild-type levels of growth likely reflects the similarities and common structural requirements in R. palustris and R. rubrum form II enzymes. However, a deeper truncation of the native terminus and replacement with shorter sequences from either Synechococcus 6301 (SyCt) or T. kodakaraensis (TkCt) Rubiscos both resulted in negative phenotypes in vivo correlated with a complete loss of in vitro activity. This suggests that the terminal α helix I may play a critical role in the form II enzyme (Fig. 2B), which is also consistent with results from previous studies in which deeper truncations to the R. rubrum carboxyl terminus caused severe perturbations to both structure and function (54, 55). Furthermore, it is likely that the terminal residues introduced by substitutions in the SyCt and TkCt enzymes may not have sterical compatibility with residues from loop 6 and the neighboring large subunit (Fig. 6) when these proteins adopt a form I- or form III-like closed conformation. The significant increase in the $k_{\text{cat}}$ of the chimeric RrCt enzyme indicates that a longer form II carboxyl terminus might be beneficial for the R. palustris enzyme (Table 2). Systematic future analyses will be needed to test this hypothesis, and such studies bode well for enhancing Rubisco efficiency.

Conclusions—The structure-function analyses presented here with the unique hexameric form II enzyme from R. palustris has identified a region in the amino terminus that has been proposed to undergo major conformational changes accompanying the reaction of the enzyme with its substrate RuBP. Tertiary structure alignments show this region to be highly variable among different forms of Rubisco, making it a promising target for future genetic engineering. Two substitutions created in the vicinity of the active site of R. palustris form II enzyme to mimic the arrangement in form I enzymes resulted in a negative phenotype. It should thus be concluded that other divergent residues elsewhere in the structure must contribute differently to catalysis in form II and form I enzymes. One such region could be the carboxyl terminus where chimeric replacements resulted in opposing changes to the enzymatic properties. The high resolution structure of the R. palustris form II Rubisco complexed with the transition-state analog CABP now allows for comparisons with similar form I and form III CABP structures, potentially leading to rational design of future experimental approaches to understand functional differences between these structurally distinct enzyme forms.

Acknowledgments—We thank Prof. David Eisenberg for invaluable and generous support for the project and Dr. Michael Sawaya for helpful advice regarding various aspects of x-ray crystallography and structural analyses. Dr. Mark Arbing and Dan E. McNamara helped with the determination of protein molecular weights using static light scattering and provided critical feedback on the manuscript. X-ray data collection was facilitated by the UCLA-DOE X-ray Crystallography Core Facility and Lawrence Berkeley National Laboratory Advanced Light Source. We also acknowledge Richa Patel and Katherine Huening for help with cloning and initial mutagenesis experiments. Comments from Drs. Daulio Cascio, Michael Sawaya, and Lorena Saeldices were of great help in improving the quality of the manuscript, and we are grateful for that.

REFERENCES

1. Tabita, F. R., Hanson, T. E., Satagopan, S., Witte, B. H., and Kree, N. E. (2008) Phylogenetic and evolutionary relationships of Rubisco and the Rubisco-like proteins and the functional lessons provided by diverse molecular forms. Philos. Trans. R. Soc. Lond. B Biol. Sci. 363, 2629–2640

2. Peterhansel, C., and Offermann, S. (2012) Re-engineering of carbon fixation in plants—challenges for plant biotechnology to improve yields in a high-CO2 world. Curr. Opin. Biotechnol. 23, 204–208

3. Andersson, I. (2008) Catalysis and regulation in Rubisco. J. Exp. Bot. 59, 1555–1568

4. Parry, M. A., Andralojc, P. J., Scales, J. C., Salvucci, M. E., Carmona-Silva, A. E., Alonso, H., and Whitney, S. M. (2013) Rubisco activity and regulation as targets for crop improvement. J. Exp. Bot. 64, 717–730

5. Guadalupe-Medina, V., Wisselink, H. W., Luttik, M. A., de Hulster, E., Daran, J. M., Pronk, J. T., and van Maris, A. J. (2013) Carbon dioxide fixation by Calvin-Cycle enzymes improves ethanol yield in yeast. Biotechnol. Biofuels 6, 125

6. Tabita, F. R., Satagopan, S., Hanson, T. E., Kree, N. E., and Scott, S. S. (2008) Distinct form I, II, III, and IV Rubisco proteins from the three kingdoms of life provide clues about Rubisco evolution and structure/function relationships. J. Exp. Bot. 59, 1515–1524

7. Andersson, I., and Backlund, A. (2008) Structure and function of Rubisco. Plant Physiol. Biochem. 46, 275–291

8. Tabita, F. R., Hanson, T. E., Li, H., Satagopan, S., Singh, J., and Chan, S. (2007) Function, structure and evolution of the Rubisco-like proteins and their Rubisco homologs. Microbiol. Mol. Biol. Rev. 71, 576–599

9. van Lun, M., van der Spoel, D., and Andersson, I. (2011) Subunit interface dynamics in hexadecameric rubisco. J. Mol. Biol. 411, 1083–1098

10. Nakano, T., Ashida, H., Mizohata, E., Matsumura, H., and Yokota, A. (2010) An evolutionally conserved Lys112 is essential for function in Rhodospirillum rubrum bona fide Rubisco and Bacillus subtilis Rubisco-like protein. Biochem. Biophys. Res. Commun. 392, 212–216

11. Liggins, J. R., and Gready, J. E. (2009) Putative functional role for the invarient aspartate 263 residue of Rhodospirillum rubrum Rubisco. Biosci. Biotech. Biochem. 48, 2226–2236

12. Harpel, M. R., Larimer, F. W., and Hartman, F. C. (1998) Multiple catalytic roles of His287 of Rhodospirillum rubrum ribulose 1,5-bisphosphate carboxylase/oxygenase. Protein Sci. 7, 730–738

13. Chêne, P., Day, A. G., and Fersht, A. R. (1997) Role of isoleucine-164 at the active site of rubisco from Rhodospirillum rubrum. Biochem. Biophys. Res. Commun. 232, 482–486

14. Andersson, I. (1996) Structure and function relationships. Philos. Trans. R. Soc. Lond. B Biol. Sci. 351, 201–221

15. Paoli, G. C., and Tabita, F. R. (1998) Aerobic chemolithoautotrophic growth and Rubisco function in Rhodobacter capsulatus and a spontaneous gain of function mutant of Rhodobacter sphaeroides. Arch. Microbiol. 2226–2236

16. Harpel, M. R., Larimer, F. W., and Hartman, F. C. (1998) Multiple catalytic roles of His287 of Rhodospirillum rubrum ribulose 1,5-bisphosphate carboxylase/oxygenase. Protein Sci. 7, 730–738

17. Chêne, P., Day, A. G., and Fersht, A. R. (1997) Role of isoleucine-164 at the active site of rubisco from Rhodospirillum rubrum. Biochem. Biophys. Res. Commun. 232, 482–486

18. Andersson, I. (1996) Large structural changes at high resolution: the 1.6 Å crystal structure of spinach ribulose-1,5-bisphosphate carboxylase/oxygenase complexed with 2-carboxyarabinitol bisphosphate. J. Mol. Biol. 259, 160–174

19. Paoli, G. C., and Tabita, F. R. (1998) Aerobic chemolithoautotrophic growth and Rubisco function in Rhodobacter capsulatus and a spontaneous gain of function mutant of Rhodobacter sphaeroides. Arch. Microbiol.
Structure-Function Analysis of a Hexameric Form II Rubisco

biol. 170, 8–17

16. Larimer, F. W., Chain, P., Hauser, L., Lamerdin, J., Malfatti, S., Do, L., Land, M. L., Pelletier, D. A., Beatty, J. T., Lang, A. S., Tabita, F. R., Gibson, J. L., Hanson, T. E., Bobst, C., Torres, J. L., Peres, C., Harrison, F. H., Gibson, J., and Harwood, C. S. (2004) Complete genome sequence of the metabolically versatile photosynthetic bacterium Rhodopseudomonas palustris. Nat. Biotechnol. 22, 55–61

17. Romagnoli, S., and Tabita, F. R. (2006) A novel three-protein two-component system provides a regulatory twist on an established circuit to modulate expression of the cbbI region of Rhodopseudomonas palustris CGA010. J. Bacteriol. 188, 2780–2791

18. Joshi, G. S., Zianni, M., Bobst, C. E., and Tabita, F. R. (2013) Regulatory twist and synergistic role of metabolic coinducer- and response regulator-mediated CbbR-cbbI interactions in Rhodopseudomonas palustris CGA010. J. Bacteriol. 195, 1381–1388

19. Spreitzer, R. J., and Salvucci, M. E. (2002) Rubisco: structure, regulatory interactions, and possibilities for a better enzyme. Annu. Rev. Plant. Biol. 53, 449–475

20. Satagopan, S., Scott, S. S., Smith, T. G., and Tabita, F. R. (2009) A Rubisco mutant that confers growth under a normally “inhibitory” oxygen concentration. Biochemistry 48, 9076–9083

21. Mueller-Cajar, O., and Whitney, S. M. (2008) Directing the evolution of Rubisco and Rubisco activase: first impressions of a new tool for photosynthesis research. Photosynth. Res. 98, 667–675

22. Smith, S. A., and Tabita, F. R. (2004) Glycine 176 affects catalytic properties and stability of the Synechococcus sp. strain PCC6301 ribulose-1,5-bisphosphate carboxylase/oxygenase. J. Biol. Chem. 279, 25632–25637

23. Smith, S. A., and Tabita, F. R. (2003) Positive and negative selection of mutant forms of prokaryotic (cyanobacterial) ribulose-1,5-bisphosphate carboxylase/oxygenase. J. Biol. Chem. 331, 557–569

24. Terzaghi, B. E., Laing, W. A., Christeller, J. T., Petersen, G. B., and Hill, D. F. (1986) Ribulose 1,5-bisphosphate carboxylase. Effect on the catalytic properties of changing methionine-330 to leucine in the RpH1 ribuliprotein enzyme. Biochem. J. 235, 839–846

25. Spreitzer, R. J., Brown, T., Chen, Z., Zhang, D., and Al-Abed, S. R. (1988) Missense mutation in the Chlamydomonas chloroplast gene that encodes the Rubisco large subunit. Plant Physiol. 86, 887–898

26. Lee, G. J., McDonald, K. A., and McFadden, B. A. (1993) Leucine 332 mutation of the Rubisco large subunit. J. Biol. Chem. 268, 909–912

27. Li, L. A., and Tabita, F. R. (1997) Maximum activity of recombinant ribulose 1,5-bisphosphate carboxylase/oxygenase of Anacystis nidulans. Protein Sci. 6, 1147–1154

28. Whitney, S. M., von Caemmerer, S., Hudson, G. S., and Andrews, T. J. (1999) Directed mutation of the Rubisco large subunit of tobacco influences photosynthesis and growth. Plant Physiol. 121, 579–588

29. Whitman, W., and Tabita, F. R. (1976) Inhibition of D-ribulose 1,5-bisphosphate carboxylase by pyridoxal 5’-phosphate. Biochem. Biophys. Res. Commun. 71, 1034–1039

30. Bradford, M. M. (1976) A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of protein-dye binding. Anal. Biochem. 72, 248–254

31. Kuehn, G. D., and Hsu, T. C. (1978) Preparative-scale enzymatic synthesis of D-1-[14C]ribulose 1,5-bisphosphate. Biochem. J. 175, 909–912

32. Li, L. A., and Tabita, F. R. (1997) Maximum activity of recombinant ribulose 1,5-bisphosphate carboxylase/oxygenase of Anabaena sp. strain CA requires the product of the rbcX gene. J. Bacteriol. 179, 3793–3796

33. Zhu, G., and Jensen, R. G. (1990) Status of the substrate binding sites of ribulose bisphosphate carboxylase as determined with 2-C-carboxyaryl-nitro 1,5-bisphosphate. Plant Physiol. 93, 244–249

34. Otwinowski, Z., and Minor, W. (1997) Processing of X-ray diffraction data collected in oscillation mode. Methods Enzymol. 276, 307–326

35. Adams, P. D., Afonine, P. V., Bunkóczi, G., Chen, V. B., Davis, I. W., Echols, N., Headd, J. J., Hung, L. W., Kapral, G. J., Grosse-Kunstleve, R. W., McCoy, A. J., Moriarty, N. W., Oeffner, R., Read, R. J., Richardson, D. C., Richardson, J. S., Terwilliger, T. C., and Zwart, P. H. (2010) PHENIX: a comprehensive Python-based system for macromolecular structure solution. Acta Crystallogr. D Biol. Crystallogr. 66, 213–221

36. Emsley, P., Lohkamp, B., Scott, W. G., and Cowtan, K. (2010) Features and development of Coot. Acta Crystallogr. D Biol. Crystallogr. 66, 486–501

37. Lee, B., and Richards, F. M. (1971) The interpretation of protein structures: estimation of static accessibility. J. Mol. Biol. 55, 379–400

38. Satagopan, S., and Kuijlaars, A. B. (1997) Distributing many points on a sphere. Math. Intell. 19, 5–11

39. Yeates, T. O. (1995) Algorithms for evaluating the long-range accessibility of protein surfaces. J. Mol. Biol. 249, 804–815

40. Winz, M. D., Isupov, M. N., and Murshudov, G. N. (2001) Use of TLS parameters to model anisotropic displacements in macromolecular refinement. Acta Crystallogr. D Biol. Crystallogr. 57, 122–133

41. Fabiola, F., Bertram, R., Korostelev, A., and Chapman, M. S. (2002) An improved hydrogen bond potential: impact on medium resolution protein structures. Protein Sci. 11, 1415–1423

42. Lawrence, M. C., and Colman, P. M. (1993) Shape complementarity at protein/protein interfaces. J. Mol. Biol. 234, 946–950

43. Krissinel, E., and Henrick, K. (2007) Inference of macromolecular assemblies from crystalline state. J. Mol. Biol. 372, 774–797

44. Sugawara, H., Yamamoto, H., Shibata, N., Inoue, T., Okada, S., Miyake, C., Yokota, A., and Kai, Y. (1999) Crystal structure of carboxylase reaction-oriented ribulose 1,5-bisphosphate carboxylase/oxygenase from a thermophilic red alga, Gaetviera partita. J. Biol. Chem. 274, 15655–15661

45. Schneider, G., Lindqvist, Y., and Lundqvist, T. (1990) Crystallographic refinement and structure of ribulose-1,5-bisphosphate carboxylase from Rhodospirillum rubrum at 1.7 Å resolution. J. Mol. Biol. 211, 989–1008

46. Parry, M. A., Keys, A. J., Madgwick, P. J., Carmo-Silva, A. E., and Andralojc, P. I. (2008) Rubisco regulation: a role for inhibitors. J. Exp. Bot. 59, 1569–1580

47. Clandon, W. W., Andrews, T. J., Gutteridge, S., Hartman, F. C., and Lorimer, G. H. (1998) Mechanism of rubisco: the carbamate as general base. Chem. Rev. 98, 549–562

48. Taylor, T. C., and Andersson, I. (1996) Structural transitions during activation and ligand binding in hexadecameric Rubisco inferred from the crystal structure of the activated unliganded spinach enzyme. Nat. Struct. Biol. 3, 95–101

49. Lohkamp, B., Scott, W. G., and Cowtan, K. (2010) Carboxylterminal deletion mutants of ribulosebisphosphate carboxylase from Rhodospirillum rubrum. FEBS Lett. 265, 41–45

50. Ranty, B., Lundqvist, T., Schneider, G., Middan, M., Howard, R., and Lorimer, G. (1990) Truncation of ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco) from Rhodospirillum rubrum affects the holoenzyme assembly and activity. EMBO J. 9, 1365–1373