Accommodation of GDP-Linked Sugars in the Active Site of GDP-Perosamine Synthase†,‡

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ABSTRACT: Perosamine (4-amino-4,6-dideoxy-d-mannose), or its N-acetylated form, is one of several dideoxy sugars found in the O-antigens of such infamous Gram-negative bacteria as Vibrio cholerae O1 and Escherichia coli O157:H7. It is added to the bacterial O-antigen via a nucleotide-linked version, namely GDP-perosamine. Three enzymes are required for the biosynthesis of GDP-perosamine starting from mannose 1-phosphate. The focus of this investigation is GDP-perosamine synthase from Caulobacter crescentus, which catalyzes the final step in GDP-perosamine synthesis, the conversion of GDP-4-keto-6-deoxymannose to GDP-perosamine. The enzyme is PLP-dependent and belongs to the aspartate aminotransferase superfamily. It contains the typically conserved active site lysine residue, which forms a Schiff base with the PLP cofactor. Two crystal structures were determined for this investigation: a site-directed mutant protein (K186A) complexed with GDP-perosamine and the wild-type enzyme complexed with an unnatural ligand, GDP-3-deoxyperosamine. These structures, determined to 1.6 and 1.7 Å resolution, respectively, revealed the manner in which products, and presumably substrates, are accommodated within the active site pocket of GDP-perosamine synthase. Additional kinetic analyses using both the natural and unnatural substrates revealed that the $K_m$ for the unnatural substrate was unperturbed relative to that of the natural substrate, but the $k_{cat}$ was lowered by a factor of approximately 200. Taken together, these studies shed light on why GDP-perosamine synthase functions as an aminotransferase whereas another very similar PLP-dependent enzyme, GDP-4-keto-6-deoxy-d-mannose 3-dehydratase or ColD, catalyzes a dehydration reaction using the same substrate.

The lipopolysaccharide (LPS) is a major component of the outer membrane of Gram-negative bacteria. It is a complex entity composed of three parts: the outermost O-specific polysaccharide, the middle core polysaccharide, and the innermost portion termed Lipid A. The O-specific polysaccharide, or O-antigen, typically contains repeating units of three to five sugars. It varies among bacterial strains in its sugar composition, linkage, and sequence. Often, the LPS contains unusual dideoxy sugars such as D-perosamine, D-tyvelose, or L-colitose, among others. In the O-specific polysaccharide, the middle core polysaccharide, and the innermost portion termed Lipid A, the O-specific polysaccharide is oxidized to a keto functionality, yielding NDP-4-keto-6-deoxymannose. This reaction is catalyzed by NDP-hexose 4,6-dehydratase. Both the nucleotidyltransferases and the dehydratases have been well characterized with respect to structure and/or function. Importantly, the NDP-4-keto-6-deoxymannose intermediate represents the branching point for all of the subsequent enzymatic reactions that ultimately lead to the production of unusual sugars such as perosamine.

The focus of this investigation is GDP-perosamine synthase (Scheme 1), which catalyzes the formation of GDP-d-perosamine from GDP-4-keto-6-deoxymannose. Interest in this enzyme grew from our previous structural investigation of GDP-4-keto-6-deoxy-d-mannose-3-dehydratase, an enzyme involved in colitose production and hereafter referred to as ColD (8–10). Like GDP-perosamine synthase, ColD utilizes GDP-4-keto-6-deoxymannose as a substrate (Scheme 1). However, rather than catalyzing the amination of the sugar C-4′ atom, ColD functions to remove the hydroxyl group at C-3′. ColD and GDP-perosamine synthase display an amino acid sequence identity of ~23%, and both of these enzymes require pyridoxal 5′-phosphate (PLP) and L-glutamate for activity. Whereas GDP-perosamine synthase contains the typically conserved lysine residue responsible for anchoring the PLP cofactor to the

NMP moiety via a nucleotidyltransferase. Subsequently, the C-6′ hydroxyl group is removed, and the C-4′ hydroxyl group is oxidized to a keto functionality, yielding NDP-4-keto-6-deoxymannose. This reaction is catalyzed by NDP-hexose 4,6-dehydratase. Both the nucleotidyltransferases and the dehydratases have been well characterized with respect to structure and/or function. Importantly, the NDP-4-keto-6-deoxymannose intermediate represents the branching point for all of the subsequent enzymatic reactions that ultimately lead to the production of unusual sugars such as perosamine.

1 This research was supported in part by NIH Grant DK47814 to H.M.H.
2 X-ray coordinates have been deposited in the Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, N. J. (accession nos. 3DR4 and 3DR7).
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4 Abbreviations: GDP, guanosine 5′-diphosphate; HEPES, N-(2-hydroxyethyl)piperazine- N′-2-ethanesulfonic acid; MES, 2-(N-morpholino)ethanesulfonic acid; NADP⁺, nicotinamide adenine dinucleotide phosphate; NMP, nucleotide monophosphate; PLP, pyridoxal 5′-phosphate; PMP, pyridoxamine 5′-phosphate; Tris, tris(hydroxymethyl)- aminomethane.
protein, CoID contains a histidine residue, thereby precluding covalent bond formation between the protein and the cofactor.

We recently determined the structure of GDP-perosamine synthase from *Caulobacter crescentus* CB15 with crystals grown in the presence of α-ketoglutarate (11). From this initial investigation of the *C. crescentus* GDP-perosamine synthase, we were able to produce a novel GDP-linked sugar, GDP-4-amino-3,4,6-trideoxy-D-mannose, hereafter termed GDP-3-deoxyperosamine (11). This was accomplished by reacting the CoID product with GDP-perosamine synthase in the presence of L-glutamate (Scheme 1). This study also revealed that GDP-perosamine synthase is a dimer with two regions, delineated by Arg 19—Ser 33 and Tyr 221—Gln 236, providing extensive subunit—subunit contacts. These two regions from one subunit project toward the active site of the second subunit (and vice versa). The individual subunits of GDP-perosamine synthase are characterized by a seven-stranded mixed β-sheet, a two-stranded antiparallel β-sheet, and 12 α-helices as shown in Figure 1a. The overall fold of the enzyme places it into the well-characterized aspartate aminotransferase superfamily (12–14). Shown in Figure 1b is a close-up view of the interactions between the protein and the α-ketoglutarate ligand. A key electrostatic interaction between Arg 231 from the second subunit of the dimer and α-ketoglutarate serves to anchor the ligand into the active site region.

Perhaps one of the most intriguing biochemical questions regarding CoID and GDP-perosamine synthase is why one functions as a dehydratase and the other acts as an aminotransferase. Is it simply a result of CoID containing a histidine rather than the lysine residue normally found in PLP-dependent enzymes, or are there other factors involved? To further address this fascinating issue, we report here the structural analysis of GDP-perosamine synthase determined in the presence of either GDP-perosamine, its natural product, or GDP-3-deoxyperosamine. Taken together, the high-resolution structures described herein provide a more detailed understanding of the active site geometry of GDP-perosamine synthase and allow for its comparison with the substrate binding pocket of CoID.

**MATERIALS AND METHODS**

Cloning, Site-Directed Mutagenesis, Expression, and Purification of GDP-Perosamine Synthase. The gene encoding GDP-perosamine synthase was cloned from *C. crescentus* as previously described (11) and used to produce the pET28t-Per plasmid required for this investigation. Lys 186 was changed to an alanine through site-directed mutagenesis using the QuikChange mutagenesis kit (Stratagene). Mutagenesis was confirmed by DNA sequence analysis. The protein was expressed and purified as previously described (11), and subsequently dialyzed against 25 mM Tris-HCl (pH 8.0) and 100 mM NaCl. For crystallization experiments, the protein was concentrated to 25 mg/mL as estimated by the absorbance at 280 nm using an extinction coefficient of 1.40 (mg/mL)^-1^ cm^-1^.

Wild-type enzyme required for the structural and functional studies was prepared as previously reported (11).

Crystallization of GDP-Perosamine Synthase. Crystallization conditions were first surveyed by the hanging drop method of vapor diffusion using a sparse matrix screen developed in the laboratory. Large single crystals of the His-tagged GDP-perosamine synthase K186A mutant protein were subsequently grown via batch methods by mixing 20 µL of the protein solution (at 25 mg/mL) with 20 µL of a precipitant solution containing 100 mM MES (pH 6.5), 20% poly(ethylene glycol) 8000, 2 mM PLP, and 2 mM glutamate. The crystals grew to maximum dimensions of ~0.1 mm × ~0.3 mm × ~0.6 mm in 1 week. They belonged to space group P2_1 with two dimers in the asymmetric unit. Crystals of wild-type GDP-perosamine synthase were grown as previously reported (11).

Structural Analysis of GDP-Perosamine Synthase. GDP-perosamine and GDP-3-deoxyperosamine were produced as described previously (11). Crystals of the wild-type GDP-perosamine synthase were transferred to a synthetic mother liquor consisting of 100 mM MES (pH 6.5), 50 mM NaCl, 26% poly(ethylene glycol) 8000, 2 mM PLP, 2 mM α-ketoglutarate, and 20 mM GDP-3-deoxyperosamine. The K186A GDP-perosamine synthase crystals were transferred to the same synthetic solution, but with 20 mM GDP-perosamine replacing the GDP-3-deoxyperosamine. Crystals were incubated at room temperature in these solutions for more than 24 h. α-Ketoglutarate was used in the soaking solutions to ensure that the cofactor remained in the PLP state, thereby allowing the amino sugar to react with the cofactor.

Single crystals were subsequently transferred to cryoprotectant solutions containing 100 mM MES (pH 6.5), 150 mM NaCl, 26% poly(ethylene glycol) 8000, 15% ethylene glycol, 2 mM PLP, and 2 mM α-ketoglutarate, with the addition of either 30 mM GDP-perosamine (for the K186A crystals) or 30 mM GDP-3-deoxyperosamine (for the wild-type crystals). X-ray data were collected from the K186A crystal at SBC beamline 19-BM (Advanced Photon Source, Argonne National Laboratory, Argonne, IL). The data set was processed.
with HKL2000 (15) and scaled with SCALEPACK (15).

X-ray data were collected from the wild-type crystal on a Bruker Proteum CCD detector system. The X-ray source was Cu Kα radiation from a Rigaku RU200 X-ray generator operated at 50 kV and 90 mA. These X-ray data were processed with SAINT version 7.06A (Bruker AXS Inc.) and internally scaled with SADABS version 2005/1 (Bruker AXS Inc.). Relevant X-ray data collection statistics are presented in Table 1.

The structure of the K186A GDP-perosamine synthase containing GDP-perosamine was determined via molecular replacement with EPMR (16), employing the wild-type GDP-perosamine synthase determined in this laboratory as a search model. All solvent molecules, Lys 186, and the coordinates for the PLP cofactor were omitted from the search probe. The resulting model had an initial R-factor of 38%. Alternate cycles of manual model building using Coot (17) and least-squares refinement of the model with TNT (18) reduced the overall R-factor to 19.9% for all measured X-ray data to 1.7 Å resolution. Relevant refinement statistics are presented in Table 2. Ramachandran statistics indicate that 87.8% of the residues adopt φ, ψ angles in the “most favored”, whereas 12.2% lie in the “additionally allowed” regions.

Enzymatic Assays. GDP-mannose-4,6-dehydratase was purified as previously described (10). GDP-4-keto-6-deoxy-mannose was produced by reacting 5 mM GDP-mannose (Sigma), 0.1 mM NADP⁺, and 3 μM GDP-mannose-4,6-dehydratase in buffer A [20 mM HEPES (pH 7.5), 50 mM NaCl, and 10 mM MgCl₂] for 2 h at room temperature. The reaction mixture was filtered through a 30 kDa cutoff filter to remove the protein. The flow-through was purified on an AKTA Purifier HPLC system (GE Healthcare) equipped with a Resource-Q 6 mL anion exchange column (GE Healthcare). The column was equilibrated with buffer A, after which the reaction mixture was loaded onto the column, washed, and eluted with a linear gradient to 30% buffer B [20 mM HEPES (pH 7.5), 1 M NaCl, and 10 mM MgCl₂]. GDP-4-keto-6-deoxy-mannose eluted in 110 mM NaCl.

The activity of wild-type GDP-perosamine synthase was assessed using an HPLC-based assay in which the decrease

![Figure 1: Molecular architecture of C. crescentus GDP-perosamine synthase. (a) Ribbon representation of the dimer. The initial structure that was determined contained PLP attached to Lys 186 via a Schiff base (indicated in sphere representative). The local 2-fold rotational axis relating one subunit to the other lies in the plane of the figure as indicated by the black arrow. (b) Close-up view of the active site. Crystals used for the initial structural analysis of GDP-perosamine synthase were grown in the presence of α-ketoglutarate, and the structure was determined to 1.8 Å resolution. The Lys 186/PLP internal aldimine and α-ketoglutarate are highlighted with slate and green bonds, respectively. Possible hydrogen bonding interactions are depicted as dashed lines. Figures 1–4 were prepared with PyMOL (21).](image-url)
in the level of GDP-4-keto-6-deoxymannose was measured over time. Reactions were carried out at room temperature, and mixtures consisted of varying concentrations of GDP-4-keto-6-deoxymannose (0.010–0.200 mM) or GDP-4-keto-3,6-dideoxymannose (0.006–0.080 mM), L-glutamate (0.005–1.50 mM), 100 µM PLP, and 1.9 µg/mL GDP-perosamine synthase (for the GDP-4-keto-6-deoxymannose reaction) or 19 µg/mL GDP-perosamine synthase (for the GDP-4-keto-3,6-dideoxymannose reaction) in buffer C [20 mM HEPES (pH 7.5), 110 mM NaCl, and 10 mM MgCl2]. At appropriate time intervals, samples of the reaction mixture were removed and treated with HCl to 80 mM. Carbon tetrachloride was added to the sample, which was subsequently mixed. The sample was then centrifuged, and the aqueous fraction was removed and diluted 10-fold with buffer A. The diluted sample was loaded onto the HPLC system equipped with a 1 m L Resource-Q anion exchange column equilibrated with buffer A. The sample was eluted with buffer B, and the concentration of GDP-4-keto-6-deoxymannose in the sample was determined by comparing the observed GDP-4-keto-6-deoxymannose peak integration to that of a standard sample.

Reaction rates determined from the assay using the natural substrate were then fitted to eq 1, whereas the rates from the assay using the unnatural substrate were fitted to eq 2.

<table>
<thead>
<tr>
<th>Table 1: X-ray Data Collection Statistics</th>
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<tbody>
<tr>
<td>enzyme complexed with GDP-perosamine</td>
</tr>
<tr>
<td>space group</td>
</tr>
<tr>
<td>unit cell</td>
</tr>
<tr>
<td>dimensions</td>
</tr>
<tr>
<td>resolution limits</td>
</tr>
<tr>
<td>no. of independent reflections</td>
</tr>
<tr>
<td>completeness (%)</td>
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<tr>
<td>redundancy</td>
</tr>
<tr>
<td>avg I/avg σ(I)</td>
</tr>
<tr>
<td>Rsym (%)</td>
</tr>
</tbody>
</table>

\[ R_{sym} = (\sum |I_I - I_J|/\sum I_I) \times 100. \]

Statistics for the highest-resolution bin.

in the level of GDP-4-keto-6-deoxymannose was measured over time. Reactions were carried out at room temperature, and mixtures consisted of varying concentrations of GDP-4-keto-6-deoxymannose (0.010–0.200 mM) or GDP-4-keto-3,6-dideoxymannose (0.006–0.080 mM), L-glutamate (0.005–1.50 mM), 100 µM PLP, and 1.9 µg/mL GDP-perosamine synthase (for the GDP-4-keto-6-deoxymannose reaction) or 19 µg/mL GDP-perosamine synthase (for the GDP-4-keto-3,6-dideoxymannose reaction) in buffer C [20 mM HEPES (pH 7.5), 110 mM NaCl, and 10 mM MgCl2]. At appropriate time intervals, samples of the reaction mixture were removed and treated with HCl to 80 mM. Carbon tetrachloride was added to the sample, which was subsequently mixed. The sample was then centrifuged, and the aqueous fraction was removed and diluted 10-fold with buffer A. The diluted sample was loaded onto the HPLC system equipped with a 1 mL Resource-Q anion exchange column equilibrated with buffer A. The sample was eluted with buffer B, and the concentration of GDP-4-keto-6-deoxymannose in the sample was determined by comparing the observed GDP-4-keto-6-deoxymannose peak integration to that of a standard sample.

Reaction rates determined from the assay using the natural substrate were then fitted to eq 1, whereas the rates from the assay using the unnatural substrate were fitted to eq 2.
can utilize only L-glutamate as its amino donor, whereas the

\[ v_0 = \frac{V_{\text{max}}AB}{K_B + K_A + AB} \]  \hspace{1cm} (1)

\[ v_0 = \frac{V_{\text{max}}AB}{K_B(1 + B/K_i) + K_A + AB} \]  \hspace{1cm} (2)

The aminotransferase activity of K186A GDP-perosamine synthase was determined by monitoring the production of GDP-perosamine as described previously (11). The mutant enzyme was incapable of producing GDP-perosamine, further supporting the role of Lys 186 as the catalytic acid or base.

**RESULTS AND DISCUSSION**

**Amino Donor Assay.** Solutions containing 0.020 mM GPD-4-keto-6-deoxymannose, 0.100 mM PLP, 2 µg/mL GDP-perosamine synthase, and an amino donor (L-glutamate, L-glutamine, L-serine, L-aspartate, L-asparagine, L-alanine, L-glycine, or L-arginine) at 10 mM were incubated in buffer C at room temperature. The reactions were terminated after 30 min and analyzed by HPLC as described above.

**Table 2: Least-Squares Refinement Statistics**

<table>
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<th>enzyme complexed with GDP-perosamine</th>
<th>enzyme complexed with GDP-3-deoxyperosamine</th>
</tr>
</thead>
<tbody>
<tr>
<td>resolution limits (Å)</td>
<td>30–1.6</td>
<td>30–1.7</td>
</tr>
<tr>
<td>R-factor(^a) (overall) (%)/no. of reflections</td>
<td>16.9/186447</td>
<td>19.9/161554</td>
</tr>
<tr>
<td>R-factor (working) (%)/no. of reflections</td>
<td>16.6/167517</td>
<td>19.7/153306</td>
</tr>
<tr>
<td>R-factor (free) (%)/no. of reflections</td>
<td>23.8/18930</td>
<td>26.3/16248</td>
</tr>
<tr>
<td>no. of protein atoms(^b)</td>
<td>11299</td>
<td>11408</td>
</tr>
<tr>
<td>no. of heteroatoms(^c)</td>
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<td>1195</td>
</tr>
<tr>
<td>average B value (Å(^2))</td>
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<td>21.5</td>
</tr>
<tr>
<td>protein atoms</td>
<td>28.7</td>
<td>42.8</td>
</tr>
<tr>
<td>ligands</td>
<td>39.1</td>
<td>26.8</td>
</tr>
<tr>
<td>solvent</td>
<td></td>
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<tr>
<td>weighted rms deviations from ideality</td>
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</tr>
<tr>
<td>bond lengths (Å)</td>
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<td>0.014</td>
</tr>
<tr>
<td>bond angles (deg)</td>
<td>2.01</td>
<td>2.10</td>
</tr>
<tr>
<td>trigonal planes (Å)</td>
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<td>0.009</td>
</tr>
<tr>
<td>general planes (Å)</td>
<td>0.018</td>
<td>0.018</td>
</tr>
<tr>
<td>torsional angles (deg)</td>
<td>16.4</td>
<td>16.7</td>
</tr>
</tbody>
</table>

\(^a\) R-factor = \(\sum F_o - |F_c|/\sum|F_o| \times 100\), where \(F_o\) is the observed structure factor amplitude and \(F_c\) is the calculated structure factor amplitude.

\(^b\) These include multiple conformations for E145, V162, R231, Q236, and E313 in subunit 2, K43 and I118 in subunit 3, and I188 in subunit 4 in the enzyme–GDP-perosamine complex and S32 and V39 in subunit 1 and S32, V39, V162, I225, R231, and I132 in subunit 2 of the enzyme–GDP-3-deoxyperosamine complex.

\(^c\) Heteroatoms include 1259 water molecules, four PLP-linked sugar ligands, and three ethylene glycols for the enzyme–GDP-perosamine complex and 1039 water molecules, four PLPs, four GDP-3-deoxyperosamine ligands, and two ethylene glycols for the enzyme–GDP-3-deoxyperosamine complex. \(^d\) The torsional angles were not restrained during the refinement.

Although the \(K_m\) for the unnatural substrate is unperturbed relative to that of the natural substrate, the \(k_{\text{cat}}\) is lower by a factor of approximately 200. These parameters suggest that the unnatural substrate is accommodated in the active site nearly as well as the natural substrate, but the absence of the 3′-OH group affects the alignment of the sugar moiety and, thus, is not properly positioned for efficient catalysis. As a result of the decrease in \(k_{\text{cat}}\), less glutamate is required to saturate the enzyme, which leads to a decline in the \(K_m\) for glutamate. It is this decrease in the glutamate \(K_m\) that eliminates the substrate inhibition phenomenon observed with the natural substrate.

**Active Site of K186A GDP-Perosamine Synthase in Complex with GDP-Perosamine.** The crystals used in this investigation contained two dimers in the asymmetric unit. Given that the \(\alpha\)-carbon positions for all four subunits in the asymmetric unit are very similar with root-mean-square deviations between the individual chains ranging from 0.17 to 0.26 Å, the following discussion refers only to the second dimer in the X-ray coordinate file.

Initially, crystals of the wild-type enzyme were soaked in
GDP-perosamine, but the corresponding electron density maps never demonstrated convincing density for the nucleotide-linked sugar. Thus, to trap GDP-perosamine in the catalytic cleft of the enzyme, we constructed a site-directed mutant protein in which the active site Lys 186 was changed to an alanine residue. Note that the\( R\)-carbons for the subunits of the wild-type enzyme and the K186A protein superimpose with a root-mean-square deviation of \(\sim 0.15\) Å, thereby indicating no major structural perturbations resulting from the site-directed mutation.

Shown in Figure 2a is the electron density for the PLP cofactor and GDP-perosamine. As can be seen, a Schiff base between the PLP cofactor and the nucleotide-linked sugar has been trapped in the active site region. This is termed the external aldimine. The pyranosyl moiety of GDP-perosamine adopts the\( B_{2,5}\) boat rather than the more typical chair conformation. Those amino acid residues located within \(\sim 3.5\) Å of the external aldimine are shown in Figure 2b. The phosphate of the PLP moiety is anchored to the protein via the side chains of Thr 61 and Ser 181 from subunit 3 and Asn 229 from subunit 4. Additional hydrogen bonds to the phosphoryl group are provided by the backbone amide nitrogens of Gly 60, Thr 61, and two water molecules. The pyranosyl group of GDP-perosamine is positioned into the active site of the enzyme via a hydrogen bond between the ring oxygen and

### Table 3: Kinetic Parameters for *C. crescentus* GDP-Perosamine Synthase

<table>
<thead>
<tr>
<th></th>
<th>(K_m) (sugar) (mM)</th>
<th>(K_m) (glutamate) (mM)</th>
<th>(K_i) (sugar) (mM)</th>
<th>(k_{cat}) (s(^{-1}))</th>
<th>(k_{cat}/K_m) (s(^{-1}) M(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>GDP-4-Keto-6-deoxymannose Reaction</strong></td>
<td>0.013 ± 0.006</td>
<td>4.6 ± 1.4</td>
<td>0.089 ± 0.034</td>
<td>2.7 ± 0.6</td>
<td>((2.1 ± 1.1) \times 10^5)</td>
</tr>
<tr>
<td><strong>GDP-4-Keto-3,6-dideoxymannose Reaction</strong></td>
<td>0.016 ± 0.003</td>
<td>0.13 ± 0.02</td>
<td>N/A</td>
<td>0.015 ± 0.001</td>
<td>((9.4 ± 1.8) \times 10^2)</td>
</tr>
</tbody>
</table>

FIGURE 3: Structure of wild-type GDP-perosamine synthase in complex with GDP-3-deoxyperosamine. (a) Electron density corresponding to the bound nucleotide-linked sugar. The map was calculated with coefficients of the form \(F_o - F_c\), where \(F_o\) was the native structure factor amplitude and \(F_c\) was the calculated structure factor amplitude. Atoms corresponding to the GDP-3-deoxyperosamine ligand were excluded from the coordinate file. The map was contoured at 3\(\sigma\). (b) Close-up view of the active site with bound GDP-3-deoxyperosamine. Amino acids lying within \(\sim 3.5\) Å of the PLP–GDP-3-deoxyperosamine complex are shown. Those residues highlighted in slate correspond to subunit 2 in the X-ray coordinate file, whereas those displayed in green belong to subunit 1. Residue labels ending with an asterisk correspond to subunit 1. GDP-3-deoxyperosamine is highlighted with gold bonds. Potential hydrogen bonds are represented as dashed lines. Ser 32\(^*\) adopts two conformations, one of which is in the proper orientation for hydrogen bonding to the guanine base of the ligand.
between the 2'-OH group and a water. The side chain of Phe 183 (subunit 3) abuts the pyranosyl ring. Arg 315 from subunit 3 and Arg 220 and Tyr 221 from subunit 4 provide most of the hydrogen bonding and/or electrostatic interactions between the protein and the pyrophosphoryl group of GDP-perosamine. The nucleotide ribose adopts the C_2-endo pucker, and its 3-OH group hydrogen bonds to the side chain of Glu 313. Finally, the guanine base interacts extensively with the protein via the side chains of Thr 29 and Ser 32, the backbone amide nitrogens of Ile 31 and Ser 32, and the carbonyl group of Thr 29, all of which are contributed by subunit 4. In addition to these hydrogen bonds, the indole side chain of Trp 30 from subunit 4 forms a parallel stacking interaction with the guanine base.

Active Site of Wild-Type GDP-Perosamine Synthase in Complex with GDP-3-Deoxyperosamine. Like that for the enzyme–GDP-perosamine complex, the crystals used to examine the enzyme–GDP-3-deoxyperosamine structure contained two dimers in the asymmetric unit. Given that the α-carbons for the four subunits superimpose with root-mean-square deviations of 0.22–0.30 Å, the following discussion refers to only the first dimer in the X-ray coordinate file. For this complex structure, it was not necessary to employ the K186A mutant form to trap the ligand in the active site. Shown in Figure 3a is the electron density observed in the active site cleft, which is consistent with GDP-3,6-dideoxymannose. Given the resolution of the X-ray data (1.7 Å), however, it is not possible to completely rule out the possibility that a small fraction of the density corresponds to GDP-4-keto-3,6-dideoxymannose or possibly the external aldime. Again, the pyranosyl moiety of GDP-3-deoxyperosamine adopts the B_2.5 boat conformation. A close-up view of the active site is presented in Figure 3b. As can be seen, the hydrogen bonding interactions that serve to position GDP-3-deoxyperosamine into the active site are virtually identical to those observed for the enzyme–GDP-perosamine complex.

One immediate question is why it was possible to observe the binding of GDP-3-deoxyperosamine to the wild-type enzyme in the crystalline lattice, but not GDP-perosamine, given the strikingly similar manners in which these two ligands are accommodated within the active site clefts of their respective proteins (Figures 2b and 3b). An answer to this comes from the kinetic assays, described above, that probed the forward reactions for the enzyme, namely the formation of GDP-perosamine from GDP-4-keto-6-deoxymannose (or GDP-3-deoxyperosamine from GDP-4-keto-3,6-dideoxymannose). Clearly, from the results presented in Table 3, the loss of the 3'-OH group from the substrate perturbs the catalytic efficiency of the enzyme by approximately 2 orders of magnitude. Given the caveat that the X-ray crystallographic analyses exploit the back reaction (Scheme 1), it still is possible to use the kinetic results to provide a possible answer to this question. In the case of GDP-perosamine binding which resulted in the formation of the external aldime, the X-ray results clearly indicate the presence of a hydrogen bond between the pyranosyl 3'-OH group and a PLP phosphoryl oxygen (Figure 2b). Presumably, this interaction also occurs when the substrate GDP-4-keto-6-deoxymannose binds. Such an interaction cannot occur when GDP-4-keto-3,6-dideoxymannose is the substrate. This perturbation of the hydrogen bonding network defining the active site results in the unnatural substrate binding to the catalytic pocket in a manner that simply is not conducive for efficient catalysis, and this would explain why we observed GDP-3,6-dideoxymannose but not the natural product GDP-perosamine in the wild-type crystal soaking experiments. Most likely, GDP-perosamine reacted quickly with the enzyme in the crystalline lattice to form the GDP-4-keto-6-deoxymannose, which then dissociated from the enzyme.

Comparison of GDP-Perosamine Synthase with ColD. ColD catalyzes a dehydration reaction at C-3', whereas GDP-perosamine synthase catalyzes an amination reaction at C-4' (Scheme 1). These are two very different chemical outcomes, yet the overall three-dimensional architectures of ColD and GDP-perosamine synthase are strikingly similar to one another.
similar such that their subunits superimpose with a root-mean-square deviation of 1.5 Å for 327 structurally equivalent α-carbons. What then are the structural features that allow for these different reactivities with the same substrate? A superposition of the ligand binding sites for GDP-perosamine synthase and ColD is presented in Figure 4. Note that the PLP and pyranosyl moieties of the ligands are accommodated in the respective active sites of GDP-perosamine synthase and ColD in nearly identical manners. The amino acids surrounding these groups are either identical or chemically similar, the one exception being Asn 185 in GDP-perosamine synthase which forms a hydrogen bond with the ring oxygen. This does not occur in ColD due to its replacement with Ser 187. The major differences in ligand binding between these two proteins begin at the pyrophosphoryl groups and are propagated to the ribosyl moieties and guanine bases (Figure 4).

There are three major insertions in ColD relative to GDP-perosamine synthase, but only one of these is near the active site cleft. Specifically, there is a large 17-residue insertion after Arg 220 in ColD, which results in an additional α-helical turn and the insertion of Arg 219 into the active site (Figure 4). The guanidinium group of Arg 219 participates in electrostatic interactions with the phosphoryl groups of the nucleotide-linked sugar with the net effect of shifting the pyrophosphoryl moiety of the ligand more toward the interior of the pocket relative to that observed in GDP-perosamine synthase. A strictly homologous residue for Arg 219 does not exist in GDP-perosamine synthase due to the lack of this 17-residue insertion. However, in GDP-perosamine synthase, there is an arginine located at position 220 that forms electrostatic interactions with the pyrophosphoryl groups of the nucleotide-linked sugar. Notably in ColD, the homologous residue for this arginine is Ser 239 (Figure 4).

It was originally suggested in the initial study of ColD that perhaps the observed boat conformation of the sugar favored a dehydration rather than an amination reaction (10). The structures described in this report of the nucleotide-linked sugars bound to GDP-perosamine synthase, however, also show the boat conformations. As such, it can be concluded that the pyranosyl conformation has nothing to do with whether an enzyme functions as a dehydratase or an aminotransferase. Are the different reactions catalyzed by these two enzymes then simply the result of ColD having an active site histidine and GDP-perosamine synthase containing an active site lysine? Recent investigations from our laboratory demonstrate that this is also not the case. First, a site-directed mutant protein of ColD was constructed in which His 188 was replaced with a lysine (as in GDP-perosamine synthase) (9). Functional assays demonstrated that this mutant form of ColD could not catalyze the dehydratase reaction or function to aminate the nucleotide-linked sugar. In a parallel study, a site-directed mutant protein of GDP-perosamine synthase was constructed in which Lys 186 was replaced with a histidine. Again, the catalytic activity of the K186H mutant protein was below the detection limit of the assay.

From the investigations presented here, it is clear that a combination of subtle factors determines whether a particular enzyme functions as a simple aminotransferase or a more complicated dehydratase. These studies highlight the fact that the enzymatic differences between CoID and GDP-perosamine synthase are not simply confined to the manner in which the pyranose is accommodated in the active site pocket. Likewise, our experiments confirm that it is not simply the replacement of an active site lysine with a histidine that results in ColD being a dehydratase. The explanations for the differences in reactions catalyzed by these two enzymes most likely extend from the immediate active site to additional residues involved in accommodating the pyrophosphoryl and nucleotide moieties of the substrates (Figure 4). Site-directed mutagenesis experiments designed to test these hypotheses are underway.

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