Molecular Architecture of the Glucose 1-Phosphate Site in ADP-glucose Pyrophosphorylase*3

Clarisa Maria Bejar, Xiangshu Jin1, Miguel Angel Ballicora2, and Jack Preiss3
From the Department of Biochemistry and Molecular Biology, Michigan State University, East Lansing, Michigan 48824

ADP-Glc pyrophosphorylase (PPase), a key regulatory enzyme in the biosynthetic pathway of starch and bacterial glycogen, catalyzes the synthesis of ADP-Glc from Glc-1-P and ATP. A homology model of the three-dimensional structure of the Escherichia coli enzyme complexed with ADP-Glc has been generated to study the substrate-binding site in detail. A set of amino acids in the model has been identified to be in close proximity to the glucose moiety of the ADP-Glc ligand. The role of these amino acids (Glu194, Ser212, Tyr216, Asp239, Phe240, Trp274, and Asp276) was studied by site-directed mutagenesis through the characterization of the kinetic properties and thermal stability of the designed mutants. All purified alanine mutants had 1 or 2 orders of magnitude lower apparent affinity for Glc-1-P compared with the wild type, indicating that the selected set of amino acids plays an important role in their interaction with the substrate. These amino acids, which are conserved within the ADP-Glc PPase family, were replaced with other residues to investigate the effect of size, hydrophobicity, polarity, aromaticity, or charge on the affinity for Glc-1-P. In this study, the architecture of the Glc-1-P-binding site is characterized. The model overlaps with the Glc-1-P site of other PPases such as Pseudomonas aeruginosa DTDP-Glc PPase and Salmonella typhi CDP-Glc PPase. Therefore, the data reported here may have implications for other members of the nucleotide-diphosphoglucose PPase family.

The biosynthetic pathways of starch and bacterial glycogen are very similar (1). The initial and key regulatory step is the formation of the glucosyl donor molecule ADP-Glc from ATP and Glc-1-P via a reaction catalyzed by ADP-glucose pyrophosphorylase (PPase4; glucose-1-phosphate adenylyltransferase, EC 2.7.7.27), with the requirement of a divalent cation (Mg2+): ATP + Glc-1-P ↔ Mg2+ + ADP-Glc + PPi.

Most ADP-Glc PPases are allosterically regulated by small effector molecules. Although these vary according to the source, they are all intermediates of the principal carbon assimilation pathway in the respective cell (2–6). Thus, bacterial glycogen and plant starch syntheses are not modulated only by the availability of ATP but also by the accumulation of key metabolites that represent the carbon and energy balance within the cell. For instance, the enzymes from heterotrophic bacteria such as Escherichia coli are regulated by intermediates of the glycolytic pathway, with Fru-1,6-P2 as the main activator and AMP as the main inhibitor. On another hand, the ADP-Glc PPases from cells performing oxygenic photosynthesis and assimilating atmospheric CO2 through the reductive pentose phosphate pathway or the Calvin cycle (specifically cyano bacteria, green algae, and photosynthetic tissues from higher plants) are activated by 3-phosphoglycerate and inhibited by P4 (6).

Except for some Bacillus species (5–7), prokaryotic ADP-Glc PPases are homotetramers, with the monomer being ~50 kDa (2, 5, 8, 9). Characterized ADP-Glc PPases from higher plants are heterotetramers of two different but homologous subunits (2–6), the “small” or catalytic subunit (50–54 kDa) and the “large” or regulatory subunit (51–60 kDa) (10). The small subunits are highly homologous (85–95% identity), whereas the large subunits have greater divergence (50–60% identity between them) and share ~50% identity with the small subunits (11, 12). Cyanobacterial ADP-Glc PPase shares features of both bacterial and plant enzymes. The native enzyme is a homotetramer, similar to the bacterial enzyme, but is regulated by 3-phosphoglycerate and P4, like the plant enzyme, and is also immunologically more related to the plant enzyme (13).

The first ADP-Glc PPase crystal structure became recently available when Jin et al. (14) solved that of the homotetrameric Solanum tuberosum (potato tuber) small subunit in its allosterically inhibited form at a resolution of 2.1 Å. They also reported the structural determination of the enzyme complexed with either ATP or ADP-Glc at 2.6 and 2.2 Å, respectively. Attempts to obtain information on the E. coli enzyme structure through x-ray crystallography were unsuccessful. The potato tuber small subunit has only ~33% sequence identity to the E. coli enzyme, but the similar predicted secondary structure profile, together with available biochemical data, suggests that they share a common three-dimensional fold (5).

Previous chemical modification (15) and site-directed mutagenesis (16) studies on E. coli ADP-Glc PPase identified Lys195 as an important residue for Glc-1-P interaction. Replace-
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ment with other amino acids generated 100–10,000-fold increases in the S0.5 for this substrate, with all other kinetic constants at wild-type levels. Later, Fu et al. (17) reported similar results from analysis of the homologous Lys198 in the potato tuber catalytic subunit. The proposed role of this amino acid is to form an ionic bond between the ε-amino group and the negatively charged phosphate of Glc-1-P. Results with Hex-1-P analogs, which differ from Glc-1-P in their hydroxyl groups, show that other residues in the active site participate in substrate binding.

Our aim was to obtain structural information on E. coli ADP-Glc PPase by building a homology model and to probe a set of highly conserved residues in the N-terminal domain possibly involved in Glc-1-P binding. We studied the role of Glu194, Ser212, Tyr216, Asp239, Phe240, Trp274, and Asp276 by site-directed mutagenesis and kinetic characterization of the mutant enzymes and their thermal stability. All residues were replaced with alanine and other amino acids to evaluate the importance of size, charge, or hydrophobicity on the effects observed in substrate interaction.

Because these residues are highly conserved among ADP-Glc PPases, it was of interest to investigate whether they are present in other PPases that use Glc-1-P as a substrate. The observations made by comparison of the putative Glc-1-P site from our E. coli ADP-Glc PPase model and the reported crystal structures of two pyrophosphorylases, the Pseudomonas aeruginosa dTDP-Glc PPase RmA1 (18) and Salmonella typhi CDP-Glc PPase (19), have led us to propose that the results presented here have implications beyond the family of ADP-Glc PPases.

EXPERIMENTAL PROCEDURES

Materials

Oligonucleotides were synthesized and purified at the Macromolecular Facility of Michigan State University. [32P]PPP was purchased from PerkinElmer Life Sciences, and [14C]Glc-1-P from ICN Pharmaceuticals, Inc. Sodium PPI, ATP, ADP-Glc, AMP, and inorganic pyrophosphatase were purchased from Sigma. Pfu DNA polymerase was purchased from Stratagene (La Jolla, CA). All other reagents were of the highest quality available.

Homology Modeling

Comparative (homology) modeling of E. coli ADP-Glc PPase (residues 12–431) was carried out with MODELLER6 Version 1 (20–22) using the atomic coordinates of Salmonella typhi CDP-Glc PPase (small subunit; AAF75832) (37). Site-directed mutagenesis was performed by overlap extension PCR (38). Plasmid pMB3 containing the E. coli ADP-Glc PPase gene between Ndel and SacI sites, previously obtained in our laboratory,5 was used as a template. The flanking primers, which annealed with the T7 promoter and the SacI site (underlined) were 5’-TAATACGACTCACTATAGG-3’ and 5’-GATATCTGAATTCGAGCTC-3’. The overlapping primers for each mutant are depicted in supplemental Table 1. The final PCR products were gel-purified, digested with Ndel and SacI, and subcloned to obtain the different pMB3-single mutant plasmids. Plasmid pETEC-NA15-D276N was obtained using pETEC-NA15 (39) as a template, with the T7 promoter and T7 terminator as flanking primers and the same mutated overlapping primers used for pMB3-D276N (supplemental Table 1). All plasmids were sequenced at the Genomics Facility of Michigan State University to confirm incorporation of only the desired mutation.

Bacterial Strains and Expression of Recombinant ADP-Glc PPases

E. coli AC70R1-504 cells lacking endogenous ADP-Glc PPase activity were used for expression of the wild-type and pMB3-mutant enzymes as described previously for pML10 (35). EcNΔ15-D276N was expressed as EcNΔ15 (39).

Purification of pMB3-Single Mutant Plasmids

One-liter cultures of AC70R1-504 cells transformed with the pMB3-single mutant plasmids or BL21(DE3) cells transformed with pETEC-NA15-D276N were grown in 25 μg/ml kanamycin/Luria broth (1 liter) at 37 °C up to A600 = 0.8. Induction was initiated by the addition of isopropyl β-D-thiogalactopyranoside (1 mM final concentration), with subsequent incubation at 25 °C for 16 h. After induction, cells were harvested, and crude extracts were obtained as described previously (35). After centrifugation, the precipitate was resuspended in buffer A (50 mM Heps [pH 8.0], 5 mM MgCl2, 0.1 mM EDTA, and 10% sucrose). The samples were individually applied to a DEAE-Fractogel column (EMD Biosciences) and eluted with a linear gradient of 0–0.5 M NaCl. The active fractions were pooled and desalted. After this step, samples were 60–70% pure and suitable for performing kinetic analysis. Mutants E194A/Q/D, D276N (supplemental Table 1). All plasmids were sequenced at

5 M. A. Ballicora and J. Preiss, unpublished data.
D276A/N, W274A, Y216F, and D239N were resuspended in buffer B (buffer A plus 1.2 M ammonium sulfate); applied to a phenyl-Superase fast protein liquid chromatography column (GE Healthcare) equilibrated with buffer B; and eluted with a linear gradient of 1.2 to 0.001 M ammonium sulfate. Further purification of the rest of the mutants and the wild type was performed by applying the DEAE pool samples to a Matrex™ gel green A affinity chromatography column (Amicon Corp.) and eluting with a linear gradient of 0–2 M NaCl. The purest fractions of each enzyme were pooled, desalted, and concentrated; and after these steps, the proteins were >95% pure as assessed by SDS-PAGE (data not shown). 

**Protein Methods**

Protein assay, electrophoresis (SDS-PAGE), and immunoblotting were performed following protocols described previously (40). Samples were desalted and concentrated with Centricon-30 devices (Amicon Corp.).

**Enzyme Assays**

**Assay A: Pyrophosphorolysis Direction**—Formation of [32P]ATP from [32P]PP, was determined by the method of Morell et al. (41). The reaction was carried out for 10 min at 37 °C in a mixture containing 50 mM HEPES (pH 8.0), 10 mM MgCl2, 1.5 mM [32P]PP (1500–2500 dpm/nmol), 4 mM ADP-Glc, 4 mM NaF, 2 mM Fru-1,6-P2, and 0.05 mg/ml bovine serum albumin plus enzyme in a total volume of 0.25 ml.

**Assay B: Synthesis Direction**—Formation of ADP-[14C]Glc from [14C]Glc-1-P was determined by the method of Yep et al. (42). The reaction was carried out for 10 min at 37 °C in a mixture containing [14C]Glc-1-P (~400 dpm/nmol), ATP, MgCl2, and Fru-1,6-P2 at varying concentrations according to the mutant enzyme assay; 50 mM HEPES (pH 8.0); 1.5 units/ml pyrophosphatase; and 0.2 mg/ml bovine serum albumin plus enzyme in a total volume of 0.20 ml. One unit of enzyme activity is 1 μmol of product (either [32P]ATP or ADP-[14C]Glc) formed per min at 37 °C.

**Kinetic characterization**

Kinetic data were plotted as specific activity (units/mg) versus substrate or effector concentration. Kinetic constants were acquired by fitting the data to the Hill equation with a nonlinear least-square formula using Origin™ Version 5.0. Hill plots were used to calculate the Hill coefficient and the kinetic constants corresponding to the substrate or activator concentrations giving 50% of the maximal velocity (S0.5) or activation (A0.5).

**Thermal Stability**

Enzyme samples were in buffer A supplemented with bovine serum albumin to 1 mg/ml in a final volume of 100 μl. Half of the sample (50 μl) was incubated in a water bath equilibrated at 60 °C for 5 min and placed on ice immediately after. The remaining half (50 μl) was kept on ice as a control. The enzyme activities for both the heat-treated and control samples were determined in the ADP-Glc synthesis direction as described for Assay B.

**RESULTS**

**Homology Modeling**—We obtained a three-dimensional model of E. coli ADP-Glc PPase by comparative modeling using the coordinates of the recently solved crystal structure of the potato tuber ADP-Glc PPase small subunit (Protein Data Bank code 1YP2) as a template as described under “Experimental Procedures” (Fig. 1A). Although modeling is generally guaranteed to be successful if residue identity is >40%, for lower percentages, errors can be reduced employing an accurate sequence alignment (43–45). Our two enzymes shared only 33% residue identity; therefore, the alignment was manually edited, incorporating information such as conservation of functional residues and prediction of secondary structures.

Using MODELLER6 Version 1, we generated 143 models after several iterative refinements of the alignment to accommodate gaps, deletions, and insertions of the query sequence with respect to the template in the best possible way. We assessed the models with the program VERIFY_3D (23, 24) as described under “Experimental Procedures,” which evaluates the compatibility of a given residue (1D) in a certain environment (3D). A score below zero for a given residue means that the conformation adopted by that residue in the model is not compatible with its surrounding environment. In our study, we considered only those models with all 1D–3D averaged scores above zero; and among them, we chose the one with a profile most similar to that generated by the template crystal structure (Fig. 1B). The two profiles followed the same general trend except for two specific regions, both corresponding to residues located in or adjacent to loops not present in the template structure (indicated by arrows in Fig. 1A). The first, encompassing Phe90–Glu97 in the E. coli enzyme, aligns with a region in the potato tuber enzyme that is disordered in the crystal structure. The second loop, containing Lys259–Pro271, is an insertion in the bacterial enzyme. Therefore, the final conformation of these two loops in the model, which might also affect immediately adjacent secondary structures, accounted for the differences from the template structure profile. According to the model, these loops are not part of the active site, and they do not contain important conserved residues.

In agreement with our previous biochemical results (46), the modeled monomer shows a two-domain structural organization (Fig. 1A). The N terminus of ~300 residues presents a β-α-β motif arranged in an open twisted β-sheet surrounded by α-helices. It resembles the Rossmann fold, typically present in nucleotide-binding domains (47). Residues important for catalysis, Asp142 (40), and for substrate binding, Tyr114 for ATP (48) and Lys195 for Glc-1-P (16), are located in the active-site pocket in close proximity to the ADP-Glc molecule (Fig. 1C), observations that further validate the quality of our model. The C terminus is a separate domain folded as a β-helix and linked to the N terminus by a long loop. The two domains are in intimate interaction through extensive hydrophobic contacts, supporting the requirement of a full-length polypeptide to obtain normal enzyme activity and regulation (46).

**Selection of Residues for Analysis**—The three-dimensional model complexed with ADP-Glc shows the ligand placed in a
well defined pocket in the active site (Fig. 1A), and several direct interactions between the ligand and the enzyme are evident (Fig. 2). Lys^{195} makes a salt bridge with the glucose phosphate, an interaction that has been biochemically probed by Hill et al. (16) in *E. coli* ADP-Glc PPase and by Fu et al. (17) in their analysis of the homologous residue (Lys^{198}) in the potato tuber enzyme. Additionally, the hydroxyl groups of the glucosyl moiety of the ligand are involved in a complex network of hydrogen bonds with the enzyme. The side chains of Glu^{194}, Asp^{276}, and Ser^{212} and the backbone of the latter participate in such interactions.

We performed a multiple sequence alignment using the catalytic subunits of 15 ADP-Glc PPases from several sources, each of them representative of a different taxonomic group. Fig. 3 depicts part of the aligned sequences, comprising residues located in and around the putative Glc-1-P-binding domain in the N terminus of the protein. The residues that, in the model, appear to interact through hydrogen bonds with the glucosyl moiety of the ligand are absolutely conserved among all ADP-Glc PPases analyzed, suggesting that they are involved in a conserved role such as substrate binding. According to our structural model, other conserved residues in this region are also located in and around the substrate-binding pocket. Based on our observations, we selected Tyr^{216}, Asp^{239}, Phe^{240}, and Trp^{274} to be characterized together with Glu^{194}, Ser^{212}, and Asp^{276}.

Expression and Purification of pMAB3-Single Mutant Plasmids—All selected amino acids were mutated to alanine to analyze their potential role in Glc-1-P interaction. We created additional mutations to investigate whether the observed effect

![FIGURE 1. Structural model of *E. coli* ADP-Glc PPase. A, schematic representation of the monomer. The N terminus presents a Rossmann-like fold and holds the ADP-Glc molecule in the active site. Carbon atoms are shown in green, and all other atoms are colored by type. The C terminus adopts a β-helix fold and is connected to the N terminus by a long loop. Loops of low structural reliability, comprising Phe^{90}–Glu^{97} and Lys^{259}–Pro^{271}, are shown in black and indicated by arrows. B, Verify_3D profile obtained from assessment of the *E. coli* ADP-Glc PPase structural model. The profile shown in gray corresponds to our model, and that shown in black corresponds to the template crystal structure. Gaps in the template profile correspond to gaps in the sequence alignment with the *E. coli* enzyme and to stretches of amino acids not solved in the crystal structure. The two big depressions in the *E. coli* profile indicated by arrows are regions of low structural reliability and correspond to the Phe^{90}–Glu^{97} and Lys^{259}–Pro^{271} loops. C, close-up view of the modeled active site, with a bound ADP-Glc molecule. Carbon atoms are shown in green. Asp^{142}, Tyr^{114}, and Lys^{195} (white carbons), which are involved in catalysis (40) and in the binding of ATP (48) and Glc-1-P (16), respectively, are in the active site and close to the ligand.

![FIGURE 2. *E. coli* ADP-Glc PPase-substrate interaction. The stereo representation of the putative Glc-1-P-binding site shows the residues studied in this work (white carbons) and their proposed hydrogen bond interactions (dashed blue lines) with the bound ADP-Glc molecule (green carbons).]
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The primary structures of representative bacterial ADP-Glc PPases and catalytic subunits from photosynthetic organisms were aligned. The region shown here encompasses residues located at and near the putative Glc-1-P-binding site according to our homology model. The consensus sequence is also shown at the bottom. The residues studied in this work are highlighted in gray.

**FIGURE 3. Sequence alignment of E. coli ADP-Glc PPase and its homologs.** The primary structures of representative bacterial ADP-Glc PPases and catalytic subunits from photosynthetic organisms were aligned. The region shown here encompasses residues located at and near the putative Glc-1-P-binding site according to our homology model. The consensus sequence is also shown at the bottom. The residues studied in this work are highlighted in gray.

**TABLE 1**
Comparison of the specific activities and apparent affinity for the substrate Glc-1-P of the E. coli wild-type and mutant enzymes

| Enzyme | kcat | Fold decrease | S0.5 (µM) | -Fold increase | kcat/Km |
|--------|------|---------------|-----------|---------------|---------|
| Wild-type | 370.0 ± 14.4 | 1.0 | 17 ± 2 | 1 | 21,765 |
| E194A | 15.43 ± 0.7 | 25.0 | 2812 ± 127 | 165 | 6 |
| E194D | 92.7 ± 11.7 | 4.0 | 6587 ± 1160 | 388 | 14 |
| E194Q | 80.7 ± 2.0 | 4.6 | 1441 ± 369 | 85 | 56 |
| S212A | 37.2 ± 2.1 | 1.1 | 241 ± 34 | 14 | 1440 |
| S212V | 22.4 ± 1.9 | 16.5 | 6416 ± 886 | 377 | 4 |
| S212T | 179.0 ± 0.7 | 2.1 | 4659 ± 274 | 274 | 38 |
| S212Y | 16.1 ± 0.1 | 231.2 | 90 ± 3 | 5 | 184 |
| Y216F | 29.0 ± 1.0 | 12.8 | 785 ± 39 | 46 | 37 |
| D239A | 32.7 ± 2.0 | 11.3 | 524 ± 126 | 31 | 7 |
| D239E | 347.7 ± 15.9 | 1.1 | 169 ± 27 | 10 | 2146 |
| D239N | 169.0 ± 3.7 | 2.2 | 264 ± 38 | 16 | 640 |
| F240A | 171.7 ± 5.7 | 2.2 | 204 ± 10 | 17 | 812 |
| F240N | 297.0 ± 8.0 | 2.8 | 376 ± 16 | 22 | 1046 |
| W274A | 384.0 ± 23.4 | 1.0 | 50 ± 3 | 3 | 5320 |
| W274F | 260.0 ± 8.0 | 1.4 | 416 ± 32 | 31 | 463 |
| W274L | 247.5 ± 0.6 | 1.5 | 1076 ± 127 | 100 | 267 |
| D276A | 0.36 ± 0.1 | 1027.8 | 1706 ± 380 | 982 | 12 |
| K195Q | 193.3 ± 50 | 1.9 | 1670 ± 380 | 982 | 12 |

Data are from Ref. 16.
kinetic parameter were observed with mutations at Glu\textsuperscript{194} and Ser\textsuperscript{212}. Glc-1-P saturation curves obtained for the Glu\textsuperscript{194} mutants are shown in Fig. 4 as an example to illustrate the shift in the $S_{0.5}$ between the wild type and Glu\textsuperscript{194} mutants. The E194A mutant showed a 165-fold increase (Table 1) compared with the wild type; therefore, we made substitutions to aspartic acid and glutamine to evaluate the importance of the charge and side chain size in such an effect. Mutation to glutamine increased the $S_{0.5}$ for Glc-1-P by 85-fold, pointing out the importance of the negative charge for substrate binding. However, mutation to aspartic acid, which also bears a negative charge, caused a larger negative effect on this kinetic parameter (Table 1), highlighting the significance of the side chain size. These two mutations, E194D and E194Q, caused 4- and 5-fold reduced $V_{\text{max}}$ values with respect to the wild type, whereas the E194A mutation decreased the $V_{\text{max}}$ by 24-fold. The apparent affinities for ATP, Mg\textsuperscript{2+}, and the activator Fru-1,6-P\textsubscript{2} were not significantly affected by any of these mutations of Glu\textsuperscript{194} (Table 2). Our results validate the hydrogen bonds observed in the structural model (Fig. 2) and strongly suggest that Glu\textsuperscript{194} plays a role in Glc-1-P binding.

Our structural model proposes that Ser\textsuperscript{212} binds O-3 and O-4 of the sugar moiety of the ligand through hydrogen bonds with the side chain and backbone, respectively (Figs. 2 and 6). Here, we probed the role of the side chain in Glc-1-P binding. All Ser\textsuperscript{212} mutations maintained apparent affinity properties for ATP, Mg\textsuperscript{2+}, and Fru-1,6-P\textsubscript{2} at wild-type levels (Table 2). S212A also showed similar $k_{\text{cat}}$ values compared with the wild type, but it displayed a 14-fold increased $S_{0.5}$ for Glc-1-P (Table 1). S212V and S212T caused dramatic effects on the apparent affinity for Glc-1-P, with 377- and 274-fold increased $S_{0.5}$ values, respectively (Table 1). Mutation to valine decreased the $k_{\text{cat}}$ by \sim 16-fold, whereas mutation to threonine did so by \sim 2-fold compared with the wild type. Surprisingly, replacement of Ser\textsuperscript{212} with tyrosine increased the apparent affinity for Glc-1-P by only 5-fold. The $k_{\text{cat}}$ for this mutant was, however, 231-fold lower compared with that for the wild type. These results strongly suggest that Ser\textsuperscript{212} is located in the Glc-1-P-binding pocket and that it contributes to the affinity of the enzyme for this substrate.

Asp\textsuperscript{276} was replaced with alanine, asparagine, and glutamic acid. The three mutations decreased the apparent affinity for Glc-1-P by 100-, 85-, and 24-fold, respectively (Table 1). Our results point out the importance of Asp\textsuperscript{276} for Glc-1-P binding and the significance of both the negative charge and size of its side chain on such effect. The analyses of these mutants suggest an additional role for Asp\textsuperscript{276} besides Glc-1-P interaction given that other kinetic parameters were also affected. D276A and D276N had \sim 1000-fold lower $V_{\text{max}}$ values compared with the

### Table 2

Kinetic parameters of *E. coli* wild-type and mutant ADP-Glc PPases

Reactions were performed in the synthesis direction (Assay B) as described under “Experimental Procedures.” Data represent the mean of two or three identical experiments \pm the mean difference of the duplicates or triplicates.

| Enzyme     | $S_{0.5}$  | ATP $k_{\text{cat}}$ |  | Mg\textsuperscript{2+} $k_{\text{cat}}$ | Fru-1,6-P\textsubscript{2} $k_{\text{cat}}$ |
|------------|------------|----------------------| | $k_{\text{cat}}$ | $k_{\text{cat}}$ |
|            | $V_{\text{max}}$ | $\text{Fold increase}$ | | $V_{\text{max}}$ | $\text{Fold increase}$ | |
| Wild-type  | 0.59 ± 0.03 | 1.0                  | | 4.7 ± 0.05 | 1.0                  | | 59.4 ± 4.7 | 1.0                  |
| E194A      | 1.20 ± 0.04 | 2.0                  | | 4.7 ± 0.05 | 2.7                  | | 320.0 ± 48.1 | 5.4                  |
| E194D      | 0.49 ± 0.01 | 0.8                  | | 5.3 ± 0.05 | 2.0                  | | 87.0 ± 17.0 | 1.5                  |
| E194Q      | 0.17 ± 0.08 | 0.3                  | | 4.5 ± 0.2  | 1.7                  | | 20.9 ± 2.8  | 0.4                  |
| S212A      | 0.68 ± 0.07 | 1.2                  | | 5.1 ± 0.2  | 2.0                  | | 85.3 ± 1.2  | 1.4                  |
| S212V      | 0.98 ± 0.07 | 0.7                  | | 3.1 ± 0.05 | 1.2                  | | 38.2 ± 4.4  | 0.6                  |
| S212T      | 0.41 ± 0.04 | 0.7                  | | 3.3 ± 0.2  | 1.3                  | | 37.3 ± 4.2  | 0.6                  |
| S212Y      | 0.43 ± 0.03 | 0.7                  | | 3.7 ± 0.05 | 1.4                  | | 121.9 ± 11.0 | 2.0                 |
| Y216F      | 0.35 ± 0.02 | 0.6                  | | 7.8 ± 0.6  | 3.0                  | | 126.0 ± 7.0  | 2.1                 |
| D239A      | 0.16 ± 0.03 | 0.3                  | | 3.7 ± 0.5  | 1.4                  | | 168.0 ± 9.0  | 2.8                  |
| D239E      | 0.96 ± 0.03 | 1.6                  | | 5.2 ± 0.1  | 2.0                  | | 118.2 ± 19.6 | 2.0                 |
| D239N      | 0.56 ± 0.05 | 0.9                  | | 5.6 ± 0.3  | 2.2                  | | 76.6 ± 2.9  | 1.3                  |
| F240A      | 1.14 ± 0.10 | 1.9                  | | 4.2 ± 0.2  | 1.6                  | | 109.0 ± 40.1 | 1.8                |
| F240M      | 1.98 ± 0.14 | 3.4                  | | 5.8 ± 0.3  | 2.2                  | | 72.0 ± 2.1  | 1.2                  |
| W274A      | 1.04 ± 0.04 | 1.8                  | | 6.0 ± 0.2  | 2.3                  | | 304.0 ± 11.0 | 5.1                |
| W274F      | 0.28 ± 0.01 | 0.5                  | | 2.4 ± 0.1  | 0.9                  | | 59.5 ± 9.2  | 1.0                  |
| W274L      | 0.48 ± 0.03 | 0.8                  | | 3.21 ± 0.01 | 1.2                | | 226.7 ± 7.0  | 3.8                  |
| D276A      | 2.03 ± 0.02 | 3.4                  | | 11.2 ± 0.4 | 4.3                  | | 403.0 ± 36.4 | 7.0                |
| D276N      | 2.3 ± 0.1   | 3.9                  | | 13.5 ± 2.6 | 5.2                  | | 760.4 ± 49.0 | 12.8              |
| D276E      | 4.77 ± 0.04 | 8.1                  | | 15.2 ± 0.2 | 5.8                  | | 281.8 ± 46.8 | 5.0                |
| K195Q\*    | 0.19 ± 0.01 | 0.3                  | | 3.4 ± 0.1  | 1.3                  | | 21 ± 2      | 0.4                  |

\* Data are from Ref. 16. 

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**FIGURE 4.** Steady-state kinetic measurement for Glc-1-P dependence for the wild-type and E194A, E194D, and E194Q mutant enzymes. Initial velocities were determined in the ADP-Glc synthesis direction using Assay B. For wild type ( ), E194A (○), E194D (▲), and E194Q (■), $V_{\text{max}}$ values were 111.0, 46.7, 27.8, and 24.2 units/mg, respectively. Reactions for each enzyme were performed in the presence of saturating concentrations of ATP, MgCl\textsubscript{2}, and Fru-1,6-P\textsubscript{2}. 

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**TABLE 2**

Kinetic parameters of *E. coli* wild-type and mutant ADP-Glc PPases
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FIGURE 5. Superposition of the amino acids in the Glc-1-P site from three NDP-Glc PPases. A, superposition of residues from the E. coli ADP-Glc PPase model with carbons in white (this work) and from the crystal structures of P. aeruginosa RmlA (Protein Data Bank code 1G23) with carbons in magenta (18) and S. typhi CDP-Glc PPase (Protein Data Bank code 1TZF) with carbons in cyan (19). The ADP-Glc PPase model has root mean square deviations of 1.9 Å with RmlA and 2.2 Å with CDP-Glc PPase. We show the CDP-Glc PPase active site as it is in the active enzyme, with Asp131, Trp232, and Asp236 belonging to one subunit and Glu178 and Lys179 from the neighboring monomer. The ADP-Glc molecule was modeled in ADP-Glc PPase, and Mg^{2+} is present in the CDP-Glc PPase crystal structure. B, Ala140, Ser212, and Tyr216 in E. coli ADP-Glc PPase (white carbons) overlaid with the homologous residues Tyr219, Asp188, and Phe192 in CDP-Glc PPase (cyan carbons) and Leu108, Val172, and Tyr176 in RmlA (magenta carbons). C, Asp239 and Phe240 in E. coli ADP-Glc PPase (white carbons) overlaid with the homologous residues Thr236 and Phe240 in CDP-Glc PPase (cyan carbons) and Glu198 and Ile199 in RmlA (magenta carbons).

wild type (Table 1) and 3.4- and ~4-fold higher S_{0.5} values for ATP, respectively (Table 2). D276E displayed a 3-fold decreased V_{max} with respect to the wild type (Table 1) but a bigger change in the apparent affinity for ATP, characterized by an 8-fold increased S_{0.5} for this substrate (Table 2). On the other hand, all three mutations decreased the apparent affinity for Mg^{2+} by ~4–6-fold (Table 2). These results correlate with the role of the Mg^{2+} ion chelator proposed for the homologous residue (Asp^{276}) in the S. tuberosum enzyme (14).

Furthermore, the three Asp^{276} mutants had 5–15-fold higher A_{0.5} values for Fru-1,6-P_2 compared with the wild type (Table 2). To investigate whether this residue is involved in the activator site, we studied another mutant. Previous reports showed that deletions of 11 and 15 residues from the N terminus of E. coli ADP-Glc PPase render activated enzymes even in the absence of Fru-1,6-P_2, with all other kinetic parameters similar to those of the wild type (39, 49). On the basis of these results, we combined both the N-terminal deletion and the single mutation D276N to create EcNA15-D276N. The activity of the partially purified double mutant was 0.027 ± 0.003 units/mg, similar to that of the partially purified D276N single mutant (data not shown), whereas the A_{0.5} for Fru-1,6-P_2 was 51 μM, similar to the that of the wild type (Table 2). This strongly suggests that Asp^{276} is not directly involved in activator binding but is a pivotal residue for the correct interaction of the substrates with the enzyme, influencing the resulting conformational changes upon their binding.

The role of the size and aromaticity of Trp^{274} was studied by substituting it with alanine, leucine, and phenylalanine. Mutation to alanine was characterized by a 22-fold decrease in the apparent affinity for Glc-1-P and did not have significant effect on the V_{max} of the enzyme (Table 1) or on the apparent affinities for ATP, Mg^{2+}, and Fru-1,6-P_2 (Table 2). We obtained similar results when leucine was placed in this position. In contrast, all parameters remained almost unchanged compared with wild-type levels when Trp^{274} was replaced with phenylalanine, indicating that aromaticity is required at this position for proper interaction of Glc-1-P with the enzyme.

Tyr^{216} is conserved not only among ADP-Glc PPases (Fig. 3) but also in RmlA (Tyr^{216}) (Fig. 5, A and B) (18). Mutation to phenylalanine allowed us to study the role, if any, of the side chain hydroxyl group in this position. The Y216F mutant displayed a 46-fold lower apparent affinity for Glc-1-P (Table 1) and showed small variations in the apparent affinities for ATP, Mg^{2+}, and Fru-1,6-P_2, with 1–3-fold increases in the respective kinetic constants (Table 2). However, this substitution, in which OH was removed, caused a 10-fold decrease in the V_{max} of the mutant enzyme. Our structural model does not show any direct interaction between this residue and bound ADP-Glc (Fig. 2). Our biochemical data suggest, however, that the side chain OH group plays a role in Glc-1-P interaction, possibly by driving the correct positioning of the substrate in the pocket, which also affects the concomitant catalytic reaction.

Asp^{239} and Phe^{240} are also conserved residues that are located in close proximity to the ligand and that do not show any evident interaction with it in the three-dimensional model. However, the D239A mutation decreased the apparent affinity for Glc-1-P by 31-fold and the V_{max} by 11-fold without significant changes in the other kinetic constants. Likewise, D239N and D239E increased the S_{0.5} for Glc-1-P by 16- and 10-fold, respectively, compared with the wild type. In contrast, the V_{max}
of the D239N mutant was 2-fold lower compared with that of the wild type and was not affected by the D239E substitution (Tables 1 and 2). Replacement of Phe240 with alanine and methionine affected the apparent affinity for Glc-1-P, with $S_0.5$ values 12- and 7-fold higher, respectively, compared with the wild type. No significant changes in the $V_{max}$ and all other kinetic parameters analyzed here were observed with F240A and F240M (Tables 1 and 2). Together, these results suggest that Asp239 and Phe240 are important residues for Glc-1-P binding.

**DISCUSSION**

In this work, we have reported the first detailed characterization of the sugar phosphate site and the three-dimensional structure of *E. coli* ADP-Glc PPase, the Glc-1-P site. We selected a set of residues implicated in shaping this substrate pocket by examination of the primary sequences of several ADP-Glc PPases and the three-dimensional structural model of the *E. coli* enzyme complexed with ADP-Glc obtained by homology modeling. The role of the selected residues in binding Glc-1-P was probed by site-directed mutagenesis and steady-state kinetics. The kinetic characterization of the individual mutants revealed the importance of the replaced amino acids.

Knowledge of the three-dimensional structure of *E. coli* ADP-Glc PPase is essential to understand the complex network of interactions established between the protein and the substrate for proper binding. The first published ADP-Glc PPase crystal structure is that of the homotetrameric potato tuber small subunit solved by Jin *et al.* (14), which we used as a template to build a model of the *E. coli* enzyme. The sequence identity between these two proteins is 33%, which is close to the lowest range of accepted homology for performing modeling (45). However, the functional similarity between our query and template proteins and a careful inspection of the sequence alignment, which included information on predicted secondary structures and functional conserved residues, increased the probabilities of obtaining a reliable model.

*E. coli* ADP-Glc PPase has been the subject of numerous structure-function relationship studies, including those aimed to elucidate the functional role of individual amino acids. Previously, Lys195 was studied (16) and showed a very specific effect on Glc-1-P interaction. The reported effect of this residue increased by 100–10,000-fold the $S_0.5$ for Glc-1-P without affecting other kinetic constants. To illustrate this, data reported for mutant K195Q have been included here in Tables 1 and 2. These results are consistent with a very specific role of Lys195 in the binding of Glc-1-P, probably by ionic interaction between the positively charged side chain $\varepsilon$-amino group and the negative phosphate group of Glc-1-P (16, 17). It is possible that the rest of the amino acids in the substrate pocket, which are the subject of this work, interact with the sugar hydroxyls to increase the affinity of the binding and to provide the correct positioning of the ligand for catalysis.

The three-dimensional model of ADP-Glc PPase that we obtained here allowed us to visualize the spatial arrangement of a set of conserved residues potentially involved in the interaction between the enzyme and the substrate Glc-1-P. It has been reported that, although proteins can bind carbohydrates in many different ways, certain amino acids show high propensity to be in a sugar-binding site (50, 51). Some examples are aromatic rings that can pack against the hydrophobic face of a sugar (52) and carboxylates that can form bidentate hydrogen bonds with two adjacent hydroxyls of a saccharide (50). In our model, we identified Trp274, Tyr216, and Phe240, as well as asparagine, and glutamic acid, respectively. A negative charge at position 239 is necessary to guarantee the stability of the enzyme at temperatures higher than the optimum for activity.
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Glu$^{194}$, Asp$^{239}$, and Asp$^{276}$, some of which show direct contacts with the modeled ligand ADP-Glc.

We also performed a close examination of the reported three-dimensional structures of enzymes that catalyze reactions very similar to those catalyzed by ADP-Glc PPase. These enzymes are *P. aeruginosa* (Protein Data Bank code 1G23) (18) and *S. typhi* CDP-Glc PPase (Protein Data Bank code 1TZF) (19). We inspected closely their active sites and identified residues homologous to Glu$^{194}$, Lys$^{195}$, Asp$^{276}$, and Trp$^{274}$, as well as the catalytic Asp$^{142}$ (40), of ADP-Glc PPase in their active sites (Fig. 5A). Interestingly, CDP-Glc PPase is a trimeric enzyme with three active sites formed in the interface of adjacent monomers (19). Most of the residues contributing to the architecture of the Glc-1-P site belong to one of the subunits, except for Glu$^{178}$ and Lys$^{179}$, homologous to Glu$^{194}$ and Lys$^{195}$ in *E. coli* ADP-Glc PPase, which are provided by the neighboring subunit (19).

The ADP-Glc PPase structural model shows the hexose moiety of ADP-Glc largely engaged in hydrogen bonds to surrounding residues (side chains of Lys$^{195}$, Glu$^{194}$, Ser$^{212}$, and Asp$^{276}$) (Fig. 2) and the protein backbone (Ser$^{212}$ and Gly$^{179}$) (Fig. 6). Lys$^{195}$ interacts with the β-phosphate of the ADP-Glc molecule. This observation is validated by the biochemical characterization reported by Hill et al. (16) and discussed above.

Glu$^{194}$ is proposed to interact with O-2 and O-3 of the sugar ring by a bidentate hydrogen bond. The Glu$^{194}$ mutants displayed the greatest changes in Glc-1-P apparent affinity when substituted with other residues (Table 1). Removal of the negative charge, as observed with the glutamine mutant, caused a large decrease in this kinetic parameter (85-fold), suggesting its importance for substrate interaction. Still, the size of the side chain is also essential given that substitution with aspartic acid decreased the apparent affinity for Glc-1-P by >380-fold. Given that the distance between two atoms engaged in a hydrogen bond is crucial for the establishment of such an interaction, the effect observed with a shorter side chain at position 194 supports the existence of a hydrogen bond between the ligand and Glu$^{194}$. In addition, the enzyme activity seems to be affected by modifications at this position. It is possible that Glu$^{194}$ plays a key role in positioning the substrate in the correct orientation for catalysis, which also agrees with a critical contribution of size to the functionality of this residue. Our results support the central role of Glu$^{194}$ in Glc-1-P binding and explain the absolute conservation of this amino acid in the ADP-Glc PPase family (Fig. 3) and other NDP-Glc PPases such as RmlA and CDP-Glc PPase (Fig. 5A).

Ser$^{212}$ may bind Glc-1-P through hydrogen bonds between the side chain and backbone and O-3 and O-4 of the sugar ring, respectively (Figs. 2 and 6). We probed the role of the side chain OH group in this interaction by making conservative and non-conservative mutations. Although to various degrees, all Ser$^{212}$ mutants affected the apparent affinity for Glc-1-P. Homology modeling of the Ser$^{212}$ mutant active-site residues complexed with ADP-Glc showed that the interaction predicted in the wild-type enzyme model between the Lys$^{195}$ e-amino group and the phosphate of the ligand is disrupted when Ser$^{212}$ is replaced with other amino acid (supplemental Fig. 1). The 14-fold increase in the $K_{m}$ for Glc-1-P caused by the S212A mutant might be explained by the disruption of one hydrogen bond between the side chain and O-3 of the glucose moiety of the ligand. Surprisingly, the effect of the side chain OH group provided by threonine is counteracted by the presence of an additional methyl group in comparison with serine. A similar situation is observed with valine in position 212. This extra methyl group largely disrupts the proper conformation of the binding pocket. The model predicts that Ser$^{212}$ is spatially close to secondary structures containing Glu$^{194}$ and Lys$^{195}$ which, as indicated previously, are important in Glc-1-P interaction. Ser$^{212}$ is also largely engaged in a hydrogen bond network with these structures (Fig. 6). These observations might explain how some of the mutations of Ser$^{212}$ affected the apparent affinity for Glc-1-P, as mutations of Glu$^{194}$ and Lys$^{195}$ did. Surprisingly, substitution of Ser$^{212}$ with a bulky side chain amino acid, tyrosine, caused a slight change in the apparent affinity for this substrate specifically. It is possible that, as the homology model predicts, the preferred rotamer for a tyrosine in this position is one directing the phenyl group away from the Glc-1-P pocket, burying the side chain against other hydrophobic side chains and stabilizing this position by a hydrogen bond between the phenyl OH group and an adjacent backbone (data not shown). It is possible that the burying of the phenyl group causes structural arrangements, which probably extend to other parts of the active site, affecting specifically an important catalytic residue. This would be explained by the dramatic reduction in the $k_{cat}$.
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displayed by mutant S212Y. Therefore, the side chain of Ser212 might contribute to the overall affinity for Glc-1-P by making direct interactions with O-3 of the sugar ring and with adjacent backbones containing important residues for the positioning of this substrate. On the other hand, the model shows the Ser212 peptide carbonyl group binding O-4 of the hexose through a hydrogen bond (Figs. 2 and 6). This interaction can also be observed in the crystal structures of the NDP-Glc PPases RmlA (18) and CDP-Glc PPase (19). The peptide carbonyl groups of Val172 in RmlA and Asn188 in CDP-Glc PPase, homologous to Ser212 in ADP-Glc PPase (Fig. 6), also make hydrogen bonds with the substrate, implying that this interaction is important for the correct geometry of Glc-1-P in the binding pocket. Apart from the specific interactions, the size of the side chain is important for the proper architecture of the Glc-1-P-binding site.

Asp276 is important for the enzyme interaction with Glc-1-P, and it may bind O-6 of the hexose through a hydrogen bond (Fig. 2). Substitutions with other residues affected the apparent affinity for this substrate by ~25–100-fold, supporting this hypothesis. However, Asp276 might have a broader role rather than exclusively interacting with the Glc-1-P molecule because the Vmax and the apparent affinity for the other substrates were also affected by the mutations studied (Tables 1 and 2). Asp276 is spatially close to the catalytic Asp142 (40), and its homologous residue in potato tuber ADP-Glc PPase (Asp280) has been proposed as an Mg2+ chelator (14). These observations explain why the different substitutions of Asp276 also affected other kinetic parameters besides the Glc-1-P apparent affinity. In contrast to the mutations of other residues in the Glc-1-P site, the activation by Fru-1,6-P2 was also altered in the Asp276 mutants (Table 2). The results obtained with EcNA15-D276N strongly suggest that this amino acid does not directly participate in activator binding.

Asp276 may be located in a hinge-like region of the active site between the ATP and Glc-1-P subdomains. Apart from interacting with the sugar ring and Mg2+, it may also contact other residues from adjacent secondary structures, establishing a network of interactions driving the conformational changes experienced upon binding the substrates. Comparison of the potato tuber ADP-Glc PPase crystal structures complexed with ATP or ADP-Glc illustrates such subdomain movement (14). The observations of Haugen and Preiss (53) also contribute to explaining the negative effects on all the kinetic properties of the enzyme when Asp276 was mutated. They demonstrate that (a) ATP alone displays half-site occupancy in the homotetrameric enzyme; (b) Glc-1-P does not bind to the enzyme unless MgCl2 and ATP are present; and (c) ATP displays full-site occupancy in the presence of Fru-1,6-P2 and ATP was also reported. Thus, the cooperative properties and the heterotrophic interactions between substrates and effectors (53) also explain the broad effect on the kinetic properties of the enzyme when the physicochemical properties of this strategically located residue are modified.

Aromatic residues, typically Trp and Phe, are key components of several saccharide-binding sites (52). Usually, these aromatic rings have been found to be involved in stacking interactions against the face of a sugar (50). However, in our structural model, none of the three aromatic residues in close proximity to the glucosyl moiety of the ligand orients its side chain parallel to the sugar ring. The great conservation of Trp274 observed among ADP-Glc PPases (Fig. 3) and other pyrophosphorylases (Fig. 5A) might be explained by its structural role within the Glc-1-P site. Substitution with short aliphatic side chain amino acids such as alanine and leucine not only affected the apparent affinity for Glc-1-P (Table 1) but also greatly decreased the thermal stability of the enzyme (Table 3). These effects were less when Trp274 was mutated to phenylalanine, suggesting that aromaticity is important at this position. This amino acid might provide the necessary stacking interactions to shape the Glc-1-P site correctly while establishing the proper hydrophobic interactions that increase the thermal stability of the protein.

Tyr216 is also located close to the ligand, but no evident interaction is observed between the sugar ring and the side chain OH group. We evaluated the role of such a group in Glc-1-P binding with the Y216F mutation, which lower the Vmax by 10-fold and the apparent affinity for this substrate by 46-fold (Table 1). Tyr216 is conserved in all ADP-Glc PPases studied so far (Fig. 3) and is present in RmlA (Tyr176) (Fig. 5, A and B). In contrast CDP-Glc PPase bears a phenylalanine (Phe192) in the homologous position, but Tyr129, located in an adjacent β-strand, orients its side chain so that the OH group overlaps with those of Tyr216 in ADP-Glc PPase and of Tyr176 in RmlA (Fig. 5B). Given the conservation of the aromatic ring at this position, it is possible that Tyr216 plays a structural role in the Glc-1-P site architecture. On the other hand, the OH group could also make a hydrogen bond with a water molecule in direct contact with the substrate, as observed with Tyr176 in RmlA (18). This interaction might be crucial to drive the correct positioning of the Glc-1-P molecule for the enzymatic reaction because not only the apparent affinity for this substrate but also the catalytic activity was affected by the removal of the side chain OH group.

We also analyzed the possible roles of Asp239 and Phe240 as part of the Glc-1-P site. Our results with mutant D239E showed that the change in size significantly affected the apparent affinity for Glc-1-P and that a negative charge at position 239 is necessary to maintain significant enzyme activity and thermal stability (Tables 1 and 3). On the other hand, substitutions with asparagine and alanine caused the greatest alteration in apparent affinity for Glc-1-P, catalytic activity (Table 1), and thermal stability (Table 3). The structures of RmlA and CDP-Glc PPase show other hydrogen bond donors in the position homologous to Asp239. Glu198 and Thr208, respectively (Fig. 5C). Moreover, in the RmlA structure, Glu198 interacts with O-2 of the dTDP-Glc molecule through a bridging water molecule. Hydrogen bonds and ion pairs with ordered water molecules are considered important interactions that increase the thermal stability of the protein (54) and the binding affinity and specificity for the substrate (50). We cannot rule out the possibility that Asp239 also interacts with the solvent, which is critical for the correct positioning of Glc-1-P and may also affect the enzyme activity.

The data obtained with Phe240 mutants demonstrate that a hydrophobic bulky residue is needed to maintain the properties
of the enzyme at wild-type levels. The role of Phe$^{240}$ might be merely structural, and the effects on Glc-1-P apparent affinity may be a consequence of the close proximity to Asp$^{239}$. In the three-dimensional model, Phe$^{240}$ is surrounded by a hydrophobic environment, and it is probably necessary to anchor the loop containing Asp$^{239}$ in the correct position. Phe$^{240}$ is conserved in most of the ADP-Glc PPases (Fig. 3), except in those from the *Mycobacterium* sp. taxonomic group, which bear methionine in the homologous position. Similarly, CDP-Glc PPase has Trp$^{209}$, whereas RmlA has a smaller hydrophobic residue (Ile$^{199}$) (Fig. 5C). Together with our biochemical results, these observations support the role of Phe$^{240}$ as an important structural component of the Glc-1-P site.

In this work, we have presented data supporting that key amino acids in ADP-Glc PPase have a role in the affinity of the enzyme for Glc-1-P. Whether establishing direct hydrogen bonds with the hydroxyls in the sugar ring or solvent molecules or properly shaping the substrate pocket, they all have an important role in determining the architecture of the Glc-1-P site. This is the first thorough biochemical characterization performed on ADP-Glc PPases. We combined biochemical data with information from the three-dimensional model, which allowed us to hypothesize the structural basis of substrate binding. Comparison of our model with other NDP-Glc PPases reveals remarkable similarities, suggesting that the architecture of the Glc-1-P site is conserved. Biochemical data involving the amino acids examined have not been reported on other PPases to date. We believe that the results reported in this work can be extended to other members of the NDP-Glc PPase family, providing new insights toward the understanding of the evolution of these enzymes.

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