Structural and functional characterization of the Helicobacter pylori cytidine 5′-monophosphate-pseudaminic acid synthase PseF: molecular insight into substrate recognition and catalysis mechanism

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Abstract: The bacterium Helicobacter pylori is a human gastric pathogen that can cause a wide range of diseases, including chronic gastritis, peptic ulcer and gastric carcinoma. It is classified as a definitive (class I) human carcinogen by the International Agency for Research on Cancer. Flagella-mediated motility is essential for H. pylori to initiate colonization and for the development of infection in human beings. Glycosylation of the H. pylori flagellum with pseudaminic acid (Pse; 5,7-diacetamido-3,5,7,9-tetradecoxy-L-glycero-L-manno-nonulosonic acid) is essential for flagella assembly and function. The sixth step in the Pse biosynthesis pathway, activation of Pse by addition of a cytidine 5′-monophosphate (CMP) to generate CMP-Pse, is catalyzed by a metal-dependent enzyme pseudaminic acid biosynthesis protein F (PseF) using cytidine 5′-triphosphate (CTP) as a cofactor. No crystal-structural information for PseF is available. This study describes the first three-dimensional model of H. pylori PseF obtained using biocomputational tools. PseF harbors an α/β-type hydrolase fold with a β-hairpin (HP) dimerization domain. Comparison of PseF with other structural homologs allowed identification of crucial residues for substrate recognition and the catalytic mechanism. This structural information would pave the way to design novel therapeutics to combat bacterial infection.

Keywords: H. pylori, motility, glycosylation, homology modeling, pseudaminic acid

Introduction
Motility is crucial for the bacterium Helicobacter pylori to colonize in the human stomach and to develop an infection. More than half of the global population is infected with H. pylori. Helicobacter pylori can cause acute gastritis, duodenal ulcers, mucosa-associated lymphoid tissue lymphoma, gastric carcinoma and other extra-gastrointestinal diseases, including iron-deficiency anemia, idiopathic thrombocytopenic purpura, cardiovascular diseases, liver illness, pancreatic cancer, respiratory illness, skin diseases and diabetes. This is the first bacterium to be classified as a definitive human carcinogen by the International Agency for Research on Cancer. The current treatment for H. pylori eradication is the simultaneous use of a proton pump inhibitor with two or more broad-spectrum antibiotics. However, the emergence of multiple antibiotic resistance has narrowed down the choice of treatment. Therefore, identification and development of novel therapeutics to treat H. pylori infection is an urgent need.
*H. pylori* uses flagella-mediated motility to facilitate colonization in mucosa in the human stomach. *H. pylori* flagella need to be O-glycosylated with an unusual nine-carbon sugar pseudaminic acid (Pse) for proper assembly and function. Pse is synthesized in *H. pylori* via five enzymatic steps and then activated by addition of a cytidine 5'-monophosphate (CMP) before being transferred onto the flagellin. The activation reaction is catalyzed by a metal-dependent enzyme CMP-Pse synthase (also known as pseudaminic acid biosynthesis protein F [PseF], EC 2.7.7.38) using cytidine 5'-triphosphate (CTP) as a cofactor. This enzyme belongs to the nucleotide–diphosphate–sugar transferases superfamily. Insertional inactivation of the *PseF* gene in *H. pylori* resulted in the loss of formation of functional flagella and rendered the bacterium nonmotile. This suggests the importance of PseF for biosynthesis of flagella and bacterial virulence. Therefore, the Pse biosynthesis pathway is a potential target for the development of novel drugs targeting bacterial motility.

A functional homolog from *Neisseria meningitidis* CMP-5-N-acetylenuraminic acid synthetase (NmCNS; 24% sequence identity with *H. pylori* PseF [HpPseF]) catalyzes activation of N-acetylenuraminic acid (NeuAc) to form a CMP-neuraminic acid (CMP-Neu5Ac) in the presence of Mg²⁺ using CTP as a cofactor. The CMP-Neu5Ac is then transferred onto the sugar moiety that forms part of lipopolysaccharide and capsule, which are essential for bacterial virulence. These glycoproteins serve an important role in cellular development, immunity and other functions.

Structural analysis of NmCNS structure revealed that it exists as a domain-swapped homodimer architecture consisting of a catalytic hydrolase fold with a short α-hairpin (HP) domain.

The CMP-5-N-acetylenuraminic acid synthetase (CNS) enzymes follow ordered–kinetic mechanism where the CTP binds first, which is followed by the sialic acid to catalyze the reaction. Horsfall et al. reported that CNS enzymes require two Mg²⁺ ions for proper orientation of the substrates and activating the α-phosphate of CTP. An ordered solvent molecule has been proposed to serve as the general base for the reaction.

Currently, to the best of knowledge, no crystallographic three-dimensional (3D) structural information on HpPseF is available. In the current study, the physicochemical, structural and functional properties of HpPseF along with its protein–protein interaction (PPI) network information were obtained via different in silico approaches. In addition, the modeled 3D structure is compared with other structurally characterized hydrolyzing enzymes to study the structure–activity relationship and identify crucial residues for substrate recognition and catalytic mechanism.

### Materials and methods

#### Sequence retrieval, physicochemical properties and secondary structure analysis

The amino acid sequence of PseF from *H. pylori* P12 (UniProtKB id: B6JKQ2, 229 amino acids) was retrieved from the protein database of National Center for Biotechnology Information (NCBI). The ExPASy ProtParam tool was used to predict the physicochemical characteristics of HpPseF, and the results are listed in Table 1. The computed physicochemical parameters included molecular mass, isoelectric point, extinction coefficient, instability index, aliphatic index and grand average of hydropathicity (GRAVY). The CYS_REC tool from SoftBerry and the CONCORD server were used to predict the existence of disulfide bridges and secondary structural components of HpPseF, respectively.

#### TM helix prediction and subcellular localization

The subcellular localization of HpPseF was predicted by the consensus web server PSORTb. Membrane topology and transmembrane (TM) helix of HpPseF were determined by the TOPCONS server.

#### Model building, refinement and validation

The HpPseF model was predicted with the following servers: IntFOLD, I-TASSER, SWISS-MODEL, RaptorX, Phyre2 and M4T. The quality of the predicted models was

### Table 1 Physicochemical properties of the HpPseF

| Properties                      | Value               |
|---------------------------------|---------------------|
| Sequence length                 | 229                 |
| Molecular weight (kDa)          | 26.1                |
| Theoretical pl                  | 6.96                |
| Extinction coefficients (M⁻¹cm⁻¹) | 15150 v             |
| Instability index               | 40.9                |
| Aliphatic index                 | 88.7                |
| GRAVY index                     | −0.241              |
| Estimated half-life             | 30 h (mammalian reticulocytes, in vitro), >20 h (yeast in vivo), >10 h (E. coli in vivo) |
| Total number of positively charged residues (Arg + Lys) | 28                  |
| Total number of negatively charged residues (Asp + Glu) | 28                  |

**Abbreviations**: HpPseF, Helicobacter pylori pseudaminic acid biosynthesis protein F; pl, isolectric point; GRAVY, grand average of hydropathicity; E. coli, Escherichia coli.
determined by the protein structure and model assessment tools in the SWISS-MODEL workspace. The best model was selected according to the QMEAN4 Z-score, which is a linear combination of four geometrical descriptors of protein. Model refinement was performed by the 3Drefine web server. The final model was validated using the ProQ and Verify3D programs. The accuracy and stereochemical features of the final 3D model were checked by the RAM-PAGE and ERRAT servers. PPI networks analysis was carried out by the STRING database. The HpPseF model was achieved in the Protein Model Database under the access code of PM0080987. Structure figures were prepared using PYMOL (http://www.pymol.org/).

Ligand docking
The HpPseF model was used for detailed structural analysis and prediction of the ligand-binding site using the 3DLigand-Site and I-TASSER web servers. The CTP was modeled in the ligand-binding pocket. The program LigPlot* was used to depict ligand-binding residues of HpPseF.

Results and discussion
Physicochemical and functional characterization
Analysis of the physicochemical characteristics of HpPseF revealed that the protein is neutral (estimated isoelectric point [pI] = 7.0) with a molecular weight of 26115 Da (Table 1). Analysis of the amino acid composition of HpPseF revealed that leucine is the most predominant amino acid (12.2%). The instability index of the protein is >40, which suggests that the protein would be less stable in solution. The protein is expected to be cytoplasmic and harbors no TM helix. The computed negative GRAVY index (−0.24) suggests that HpPseF is hydrophobic and most likely to be insoluble in water. Therefore, expression of the protein for X-ray crystal-structure would be difficult. No disulfide bonding pattern among cysteine (Cys) residues has been predicted in HpPseF. Estimation of the secondary structure element revealed that HpPseF is an α/β protein and consists of mainly α-helices (45%), followed by the random coil (20%), extended β-strands (27%) and β-turn (8%).

Model building and refinement
The HpPseF models were predicted using different web servers (Table 2). The best model was selected based on the global QMEAN4 Z-score. The global QMEAN4 Z-score is an estimate of the absolute quality of a model and calculated based on the structure comparison between the modeled structure and high-resolution X-ray structures obtained from the protein database. Structure with QMEAN4 Z-score equivalent to 1 is similar to X-ray-diffracted crystal structure. The detailed analysis of the predicted models is presented in Table 2. RaptorX generated a model with a QMEAN4 Z-score of −3.10 (Table 2). The RaptorX model 1 was further refined and optimized by the 3Drefine web tool. The estimated QMEAN4 Z-score of the refined HpPseF model was −2.65 (Figure 1).

Model validation
The stereochemical properties of the HpPseF model was analyzed by using two web servers, ProQ and Verify3D.

Table 2 Comparison of the predicted models of the HpPseF

| Program     | Model no | QMEAN4 Z-score |
|-------------|----------|----------------|
| RaptorX     | 1        | −3.10          |
| SWISS-MODEL | 1        | −3.90          |
| I-TASSER    | 1        | −5.92          |
| Phyre2      | 1        | −4.68          |
| M4T         | 1        | −4.03          |

Notes: (A) The QMEAN quality analysis of the HpPseF 3D model (presented by a red star) is compared to the X-ray crystallographically determined structures of similar size proteins. (B) The QMEAN4 Z-score of the final HpPseF model.

Figure 1 Quality analysis of the refined 3D model of PseF from H. pylori. Most used by the STRING database. The HpPseF model was achieved in the Protein Model Database under the access code of PM0080987. Structure figures were prepared using PYMOL (http://www.pymol.org/).
The HpPseF model showed a ProQ-LG score of 4.48, which suggests that the quality of the refined model is very good. Analysis with Verify3D revealed that >79% of the residues in the model had scored ≥0.2 in the 3D/one-dimensional (1D) profile, indicating that residues are in the favorable region (Figure 2).

The HpPseF model was further evaluated for overall quality, backbone torsion angles and side chain interaction energies. The overall quality of the model was assessed by the ProSA-web server. A total of 95% of the residues (n = 215) were in the favored region, and the other 5% of the residues (n = 12) were present in the allowed region. Errors in non-bonded interactions were calculated by the ERRAT server, and the obtained score of 94.11 means that 94% of the residues in the model were below the rejection limit (experimental structures had scores ~90%; Figure 4). Altogether, this suggests that the range of structures of proteins of similar size determined by X-ray crystallography. The distribution of backbone torsion angles was checked by the RAMPAGE (Figure 3).
3D model of H. pylori PseF

**Figure 4** ERRAT result shows the overall quality factor of the HpPseF model (error-axis represents the error values to reject regions that exceed error value).

Abbreviation: HpPseF, Helicobacter pylori pseudaminic acid biosynthesis protein F.

**Figure 6** Comparison of HpPseF with other hydrolyzing homologous enzymes.

Notes: (A) HpPseF. (B) NmCNS. (C) MmCNS. α-helices, loops and β-strands are colored red, gray and green, respectively.

Abbreviations: HpPseF, Helicobacter pylori pseudaminic acid biosynthesis protein F; NmCNS, Neisseria meningitidis CMP-5-N-acetylneuraminic acid synthetase; MmCNS, Mus musculus CMP-5-N-acetylneuraminic acid synthetase; CMP, cytidine 5'-monophosphate.

The final HpPseF model is of a good quality and could be used for further studies.

**Overall structure of HpPseF and comparison to other homologs**

The HpPseF has an α/β-hydrolase domain with an ~35-residue inserted β-HP domain. The core domain is a αβα three-layer sandwich architecture consisting of seven β-strands with the topological order (↑β3-↑β2-↑β1-↑β4-↓β8-↓β5-↓β9; Figure 5). The central β-sheet is flanked by four α-helices (α1, α2, α5 and α9) on one side and three α-helices (α3, α4 and α8) on the other side. The loop between strand β1 and helix α1 is known as P-loop (Figure 5), which interacts with the pyrophosphate arm of the cofactor CTP in other structural homologs.20,46 The HP domain protrudes out from the central β-sheet and contains two antiparallel β-strands (β6 and β7) and two α-helices (α6 and α7). In other structurally characterized CNS enzymes, the HP domain plays an important role in the formation of a homodimer.20,46

Comparison of PseF against the published structures deposited in the Protein Data Bank that have been described in the literature, using the Dali server,47 revealed that mammal (Mus musculus) CMP-5-N-acetylneuraminic acid synthetase (MmCNS; root-mean-square deviation [RMSD] = 1.3, sequence = 23%, Z-score = 32.5, PDB code: 1QWJ[46]) showed the highest similarity followed by NmCNS (RMSD = 2.2, sequence = 22%, Z-score = 28.1, PDB code: 1EYR[20] Figures 6 and 7). Detailed analysis of the superimpositions revealed that structural similarity extends over the entire fold and includes all the secondary structural elements and the HP domain.
Prediction of the ligand-binding pocket and ligand docking

The CASTp server\(^48\) predicted the putative ligand-binding site in the core of the hydrolase domain. The calculated surface area and volume of the pocket are \(~438.2\, \text{Å}^2\) and \(~1711.6\, \text{Å}^3\), respectively. Prediction of the ligand-binding residues in HpPseF was performed using the 3DLigandSite\(^43\) and I-TASSER servers.\(^30\) The ligand CTP was modeled at the putative pocket. Figure 8 shows residues that form interactions with the CTP-modeled structure. The pyrophosphate moiety interacts with the residues of the P-loop between \(\beta1\) and \(\alpha1\), while the sugar moiety is accommodated by the residue from the \(\beta3–\alpha3\) region. The CTP forms hydrogen bonds and van der Waals interactions with six (Lys14, Lys19, Arg70, Ala75, Asp76 and Tyr104) and five residues (Leu8, Arg10, Ser13, Asn20 and Asp209), respectively. Prediction of the metal-binding site has revealed that Tyr104, Asp207 and Asp209 would likely interact with the metal cofactor. Sequence alignment of HpPseF with other homologs shows that residues involved in CTP binding are conserved among other homologous enzymes (Figure 7).

Analysis of the acceptor substrate-binding site and catalysis mechanism

The closest bacterial functional homolog NmCNS was solved in complex with CDP,\(^20\) while the murine homolog (MmCNS) was solved in complex with the reaction product CMP-NeuAc.\(^46\) Both CNS proteins form a homodimer, where the HP domain from one monomer interacts with the catalytic domain of the opposite monomer and contributes to the formation of the active site. Detailed structural

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**Figure 7** A sequence alignment of HpPseF, NmCNS and MmCNS.

**Notes:** The elements of the secondary structure and sequence numbering for HpPseF are shown above the alignment. Conserved residues are highlighted in red. Metal binding and catalytic residue are presented as blue stars. Yellow stars represent residues that interact with pyrophosphate arm of cofactor CTP. Residues that form a hydrophobic pocket to determine substrate specificity and contribute in proper positioning of the substrate NeuAc in NmCNS are represented as black and green stars.

**Abbreviations:** HpPseF, Helicobacter pylori pseudaminic acid biosynthesis protein F; NmCNS, Neisseria meningitidis CMP-5-N-acetylneuraminic acid synthetase; MmCNS, Mus musculus CMP-5-N-acetylneuraminic acid synthetase; CMP, cytidine 5′-monophosphate; CTP, cytidine 5′-triphosphate; NeuAc, N-acetylneuraminic acid.

**Figure 8** Ligplot\(^+\) results showing the residues of PseF from *H. pylori* interacting with the modeled CTP molecule.

**Note:** Residues making van der Waals contacts with CTP are indicated by red arcs with spokes radiating toward the ligand moieties they contact.

**Abbreviations:** PseF, pseudaminic acid biosynthesis protein F; *H. pylori*, Helicobacter pylori; CTP, cytidine 5′-triphosphate.
analysis of the NmCNS active site revealed that the enzyme has a hydrophobic pocket formed by the residues (Tyr179, Phe192 and Phe193) to aid binding of the methyl group of the N-acetyl moiety of acceptor substrate Neu5Ac and thus determines substrate specificity. Alanine substitution of these residues resulted in the greatest loss of enzyme activity and thus confirmed their crucial role in substrate recognition. The HpPseF model also adopts a hydrophobic pocket consisting of Y177, I190 and F191, which would likely play a crucial role in substrate recognition (Figures 7 and 9).

The CNS enzymes require two metal ions (preferably, manganese Mg$^{2+}$) to catalyze the condensation reaction. In NmCNS and MmCNS, two Asp residues (D211 and D209) are involved in binding the catalytic Mg$^{2+}$ ion, while another Gln residue (Q104 in NmCNS and Q141 in MmCNS) is believed to play a crucial role in binding the second metal ion at the intermediate stage of the reaction. A mutagenesis study has confirmed their role in catalysis. Structural and sequence alignment analyses have shown that HpPseF harbors two Asp residues (D207 and D209) and one Tyr residue (Y104) in the corresponding positions (Figures 7, 10 and 11).

The CNS enzymes follow an ordered bi–bi sequential kinetic mechanism where binding of a CTP with the enzyme is followed by sialic acid to generate CMP-NeuAc. The role of both Mg$^{2+}$ ions is believed to be in proper orientation of the substrates and activation of the α-phosphate moiety of CTP. The sugar hydroxyl group of NeuAc is activated by the catalytic Mg$^{2+}$ ion. The structural similarities among HpPseF and other well-characterized homologs (NmCNS and MmCNS) suggest that HpPseF would likely follow the same sequential catalytic mechanism (Michaelis–Menten equation) and catalyze the reaction via formation of a tetrahedral intermediate.

**Protein network analysis**

Analysis of PPI networks is necessary for better understanding of intricate molecular mechanisms and identification of novel therapeutics to combat multidrug-resistant bacteria. Figure 12 shows PPI partner proteins of HpPseF derived from the STRING database. The functional partners are mainly flagella glycosylation pathway proteins (PseB, PseC, PseH, HpPseG, PseI) and flagellar basal body L-ring protein (FlgH). In addition, HpPseF interacts with other proteins including polysialic acid capsule expression protein (KpsF), 50S ribosomal protein L34 (RpmH) and three hypothetical proteins (HP0465; HP0114, HP1570).

**Conclusion**

The present study presents physicochemical properties, structural and functional analysis and crucial residues of substrate recognition and the catalytic mechanism of HpPseF derived by using a broad range of biocomputational tools. The deduced structural information would serve as a fundamental basis to
design in vivo biochemical and X-ray crystallographic assays for a better understanding of the catalytic mechanism and molecular biology of Pse and sialic acid synthase enzymes. The HpPseF 3D model would pave the way for the design and production of inhibitors and mimetics of Pse and sialic acid biosynthesis pathway enzymes targeting motility to combat drug-resistant bacterial infection in the near future.

**Disclosure**

The author reports no conflicts of interest in this work.

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