Crystal Structure of Helicobacter pylori Pseudaminic Acid Biosynthesis N-Acetyltransferase PseH: Implications for Substrate Specificity and Catalysis

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Abstract

Helicobacter pylori infection is the common cause of gastroduodenal diseases linked to a higher risk of the development of gastric cancer. Persistent infection requires functional flagella that are heavily glycosylated with 5,7-diacetamido-3,5,7,9-tetraeodeoxy-L-glycero-L-manno-nonulosonic acid (pseudaminic acid). Pseudaminic acid biosynthesis protein H (PseH) catalyzes the third step in its biosynthetic pathway, producing UDP-2,4-diacetamido-2,4,6-trideoxy-β-L-altropyranose. It belongs to the GCN5-related N-acetyltransferase (GNAT) superfamily. The crystal structure of the PseH complex with cofactor acetyl-CoA has been determined at 2.3 Å resolution. This is the first crystal structure of the GNAT superfamily member with specificity to UDP-4-amino-4,6-dideoxy-β-L-AltNAc. PseH is a homodimer in the crystal, each subunit of which has a central twisted β-sheet flanked by five α-helices and is structurally homologous to those of other GNAT superfamily enzymes. Interestingly, PseH is more similar to the GNAT enzymes that utilize amino acid sulfamoyl adenosine or protein as a substrate than a different GNAT-superfamily bacterial nucleotide-sugar N-acetyltransferase of the known structure, WecD. Analysis of the complex of PseH with acetyl-CoA revealed the location of the cofactor-binding site between the splayed strands β4 and β5. The structure of PseH, together with the conservation of the active-site general acid among GNAT superfamily transferases, are consistent with a common catalytic mechanism for this enzyme that involves direct acetyl transfer from AccoA without an acetylated enzyme intermediate. Based on structural homology with microcin C7 acetyltransferase MccE and WecD, the Michaelis complex can be modeled. The model suggests that the nucleotide- and 4-amino-4,6-dideoxy-β-L-AltNAc-binding pockets form extensive interactions with the substrate and are thus the most significant determinants of substrate specificity. A hydrophobic pocket accommodating the 6'-methyl group of the altrose dictates preference to the methyl over the hydroxyl group and thus to contributes to substrate specificity of PseH.
**Introduction**

*Helicobacter pylori* is a Gram-negative, microaerophilic bacterium that colonizes the stomachs of more than half of world’s population [1]. *H. pylori* infections are associated with a number of gastroduodenal disorders ranging from gastritis, gastric and duodenal ulcers to gastric adenocarcinoma and mucosa-associated lymphoid tissue lymphoma [2, 3]. It was the first bacterium to be classified as a group I (definite) carcinogen for human gastric cancer by the International Agency for Research on Cancer [4]. *H. pylori* has a unipolar bundle of two to six sheathed flagella that enable the bacteria to drill into the highly viscous mucus lining of the stomach and reach the gastric epithelium [5]. Flagella-mediated motility is required not only for initial colonization but also for attaining robust infection and persistence of *H. pylori* in the high-flow and rapid-turnover environment of the stomach [6, 7]. *H. pylori* flagellins are O-glycosylated on serines and threonines with an unusual nine-carbon sugar pseudaminic acid (Pse) that has only been found in bacteria. Flagellin glycosylation is essential for assembly of flagellar filaments and motility, and hence for virulence [8, 9]. Therefore, the Pse biosynthesis pathway can be a potential target for novel therapeutics.

The first two enzymes in this pathway in *H. pylori*, UDP-N-acetylglucosamine dehydratase PseB and a pyridoxal-5-phosphate-dependent aminotransferase PseC, convert UDP-N-acetylglucosamine to UDP-4-amino-4,6-dideoxy-β-L-AltNAc (Fig. 1) [10]. The latter acts as a substrate for the 21-kDa Pse biosynthesis protein H (PseH), also known as flagellin modification protein H (FlmH) or flagellar protein G (FlaG1) [10, 11]. PseH is N-acetyltransferase (EC 2.3.1.202) that catalyzes transfer of an acetyl group from acetyl-CoA (AcCoA) to the C4 amino group of the nucleotide-linked sugar to produce UDP-2,4-diacetamido-2,4,6-trideoxy-β-L-Alt. Mutation in the *pseH* gene of the closely related species *Campylobacter jejuni* resulted in a non-motile phenotype that lacked flagella filaments and hook structures, indicating that PseH plays an essential role in flagella assembly [12].

Analysis of the PseH primary structure revealed low-level similarity to the GCN5-related N-acetyltransferase (GNAT) superfamily that covers more than 10,000 different enzymes from all kingdoms of life [13–15]. Members of the GNAT superfamily catalyze transfer of an acetyl group from acetyl-CoA (AcCoA) to a wide variety of substrates, including aminoglycosides, histones, aryalkylamines, glucosamine-6-phosphate, spermine, spermidine and serotonin [15]. Previous structural studies revealed that although different enzymes of this superfamily show only moderate pairwise sequence homology, they share a common core fold comprising a central highly curved mixed β-sheet flanked on both sides by α-helices, with the topology β0-β1-α1-α2-β2-β3-β4-α3-β5-α4-β6 [15–21].

The proposed reaction mechanism of most of the GNAT superfamily enzymes involves direct acetyl transfer from AcCoA without an acetylated enzyme intermediate [15]. In the first reaction step, a (non-conserved) general base abstracts a proton from the primary amine of the substrate to produce a lone pair of electrons, which then perform a nucleophilic attack on the thioester acetate. This leads to the formation of a transient bisubstrate intermediate that decomposes through proton transfer from a general acid (conserved tyrosine or serine) [15].

Limited structural information is available on enzymes that are functionally homologous to PseH. Acetyl transfer from AcCoA to the 4-amino moiety of the nucleotide-linked sugar substrate in a different biosynthetic pathway leading to legionaminic acid in *C. jejuni* is catalyzed by PgL1 which has a left-handed β-helix (LbH) fold and shows no detectable sequence similarity to PseH [22]. A different example of a bacterial nucleotide-sugar N-acetyltransferase, the *Escherichia coli* dTDP-fucosamine acetyltransferase WecD, belongs to the GNAT superfamily but shares only 15% sequence identity with PseH [17].
Here, we report the crystal structure of the \textit{H. pylori} PseH complex with AcCoA solved at 2.3 Å resolution, which allowed us to address the molecular details of substrate binding and catalysis of this enzyme. This is the first crystal structure of the GNAT superfamily member with specificity to UDP-4-amino-4,6-dideoxy-β-L-AltNAc.
Materials and Methods

Purification, determination of the oligomeric state, crystallization, preparation of derivatives and data collection

Recombinant PseH from *H. pylori* was purified as previously described [14]. The oligomeric state of PseH in solution was determined by passing it through a Superdex 200 HiLoad 26/60 gel-filtration column (GE Healthcare) equilibrated with 50 mM Tris/HCl pH 8.0, 200 mM NaCl and calculating the molecular weight (MW) using a calibration plot of log MW versus the retention volume \[ V_{\text{retention}} \text{(ml)} = 549.3 - 73.9 \times \log \text{MW} \] available at the EMBL Protein Expression and Purification Core Facility website http://www.embl.de/pepcore/pepcore_services/protein_purification/chromatography/hiload26-60_superdex200/index.html. The PseH-AcCoA crystal complex was obtained by co-crystallization with 5 mM AcCoA as described [14]. The crystals belong to space group I212121 with unit-cell dimensions \( a = 107.8 \) Å, \( b = 145.6 \) Å, \( c = 166.2 \) Å and three protein subunits in the asymmetric unit. Two different mercury derivatives were obtained by soaking crystals overnight in either mercury chloride (1 mM) or mercury potassium iodide (1 mM). To perform data collection at cryogenic temperatures, the crystals were briefly soaked in a cryo-stabilizing solution containing 1.0 M di-ammonium tartrate, 0.1 M sodium acetate trihydrate pH 3.8, 20% (v/v) glycerol and 5.0 mM AcCoA, and flash-frozen by plunging them into liquid-nitrogen. X-ray diffraction data for the native crystal were collected to 2.3 Å resolution using the MX2 beamline of the Australian Synchrotron. Diffraction data for the mercury chloride-derivitized crystal were collected to 2.4 Å resolution using the Australian Synchrotron MX1 beamline. Diffraction data for the mercury potassium iodide-derivitized crystal were collected to 2.8 Å resolution using the in-house Rigaku MicroMax-007 microfocus rotating-anode generator. All diffraction data were processed and scaled using iMOSFLM [23] and AIMLESS [24] from the CCP4 software suite [25]. Data collection statistics are summarized in Table 1.

|                         | Data set | Native | HgCl₂ | K₂Hgl₄ |
|-------------------------|----------|--------|-------|--------|
| Wavelength (Å)          | 0.95     | 1.01   | 1.01  |
| Resolution (Å)          | 2.3 (2.4–2.3) | 2.4 (2.5–2.4) | 2.8 (3.0–2.8) |
| Total No. of observed reflections | 168,694 (12,801) | 385,414 (56,182) | 117,999 (16,375) |
| No. of unique reflections | 54,332 (4,535) | 51,674 (7,446) | 32,747 (4,705) |
|Completeness (%)         | 94 (96)  | 99 (99) | 99 (99) |
|Redundancy               | 3.0 (2.8) | 7.5 (7.5) | 3.6 (3.5) |
|Mean \( I/\sigma(I) \)   | 10.3 (2.3) | 15.9 (5.4) | 11.9 (3.5) |
|\( R_{merge} \)          | 0.06 (0.30) | 0.08 (0.35) | 0.08 (0.31) |
|Overall \( B \) factor from Wilson plot (Å²) | 34       | 27     | 25     |

Values in parentheses are for the highest resolution shell.

\[
R_{merge} = \left( \frac{\sum_{h} \sum_{i} \left| I_{hi} - \langle I_{h} \rangle \right|}{\sum_{h} \sum_{i} \left| I_{hi} \right|} \right), \quad \text{where } I_{hi} \text{ is the intensity of the } h\text{th observation of reflection } h.
\]

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Structure determination

The structure of PseH was determined using the method of multiple isomorphous replacement coupled with anomalous scattering (MIRAS). The locations of the four Hg sites for the mercury...
chloride derivative and seven sites for the mercury potassium iodide derivative were found using Autosol [26] from the PHENIX software suit [27]. The overall figure of merit of the resulting phase set was 0.24 for data between 30 and 2.4 Å. An initial partial model generated using AutoBuild within PHENIX was manually completed using COOT [28] and then refined against the 2.3 Å resolution native data set using PHENIX. The electron density indicated that one acetate ion was bound to each PseH subunit. A complete model including water molecules, AcCoA and acetate ions was built through iterative cycles of re-building with COOT and refinement with PHENIX. Analysis of the stereochemical quality of the model was accomplished using MOLPROBITY [29]. The final refined model of the PseH-AcCoA complex contains 532 of the expected 555 amino acid residues, three acetate ions, three AcCoA molecules and 228 water molecules (R-factor 0.178, R_free 0.218). All the non-glycine residues lie in permitted regions of the Ramachandran plot with 97% of these in the most favoured regions. Refinement statistics are given in Table 2. Structure figures were prepared using PYMOL [30]. Accessible surface area was calculated using AREAIMOL from the CCP4 software suite [25] with a probe radius of 1.4 Å.

**Protein Data Bank accession number**

The coordinates of the PseH complex with AcCoA have been deposited in the Protein Data Bank (RCSB) under accession code 4RI1.

| Table 2. Refinement statistics. |
|--------------------------------|
| Resolution range (Å) | 30–2.3 |
| Reflections | 54302 |
| Atoms | 4902 |
| a R-factor | 0.178 |
| b R_free | 0.218 |
| Bond-length deviation from ideality (Å) | 0.008 |
| Bond-angle deviation from ideality (°) | 1.1 |

| Molprobity scores |
|-------------------|
| Ramachandran regions (%) |
| Favored | 97 |
| Allowed | 3 |
| Outliers | 0 |
| Clashscore | 4.0 |
| Average B (protein atoms) (Å²) | 40 |
| Average B (water molecules) (Å²) | 43 |
| Average B (acetate ions) (Å²) | 42 |
| Average B (AcCoA) (Å²) | 67 |

\[ a \ R = \frac{\sum \left( |F_{\text{calc}}| - |F_{\text{calc}}| \right)}{\sum \left( |F_{\text{calc}}| \right)} \]

\[ b \text{ The free R-factor was calculated on 5% of the data omitted at random.} \]
Results and Discussion

Overall structure of PseH and comparison to other members of the GNAT superfamily

Although unliganded PseH did not crystallize, co-crystallization with AcCoA readily yielded crystals. The structure of recombinant H. pylori PseH (residues 1–180 plus an additional GIDPFT fragment (cloning artifact)) was determined to 2.3 Å resolution by using the multiple isomorphous replacement coupled with anomalous scattering (MIRAS) method with two mercury derivatives. The asymmetric unit contains three molecules. To determine the correct oligomeric assembly, we performed size-exclusion chromatography and analysis of the packing of individual subunits in the crystal. When subjected to gel filtration, the protein eluted as a single peak with an apparent molecular weight of approximately 36 kDa, indicating that PseH behaves as a dimer in solution. In line with this, analysis of probable assemblies in the crystal using the PDBe PISA server (http://www.ebi.ac.uk/msd-srv/prot_int/cgi-bin/piserver) also suggested that PseH likely exists as a stable dimer in solution; two of the three molecules in the asymmetric unit form a non-crystallographic dimer, and the third molecule forms a similar dimer with a symmetry-related neighbor. The dimer is stabilized by an interface with a surface area per monomer (~1,000 Å²) that is approximately 10% of the total surface area of a single monomer (~10,100 Å²).

The PseH structure has a central twisted seven-stranded β-sheet flanked by five α-helices (Fig. 2A). The β-strands and α-helices are arranged in the topological order ββααβββαβαβαβ. The β-strands form a β-sheet in the order 01234576 (Fig. 2B). Strands β4 and β5 are splayed apart, creating a channel through the molecule (Fig. 2C) which is a signature of the GNAT fold. Helices α1 and α2 pack against one face of the β-sheet, helices α3 and α4 against the other, whereas helix α5 forms a C-terminal extension of strand β7.

In a comparison of PseH against the structures in the RCSB Protein Data Bank [31] that have been described in the literature, using the protein structure comparison service Fold at European Bioinformatics Institute (http://www.ebi.ac.uk/msd-srv/ssm) [32], significant similarities were found with other members of the GNAT superfamily. PseH has the closest structural similarity to E. coli microcin C7 self immunity acetyltransferase MccE (PDB ID code 3R9G [16]) and Salmonella typhimurium ribosomal protein L12 Nα-acetyltransferase RimL (PDB ID code 1S7N [19]) (rms deviation of 2.4 Å and 2.2 Å for the superimposition of 160 Cα atoms (Fig. 3A), showing 18% and 14% sequence identity over equivalenced positions). MccE acylates the product of unwanted processing of the antibiotic microcin C7 in E. coli, thus inactivating it [16]. RimL possesses the same activity as MccE [33] and, in addition, converts the ribosomal protein L12 to L7 by acetylating its N-terminal amino group [19]. PseH, RimL and the acetyltransferase domain of MccE adopt a very similar fold, despite the limited sequence homology (less than 8% global sequence identity and less than 15% identity for pairwise alignments of PseH with MccE and RimM; see Fig. 3B). Structural similarity extends over the entire fold and includes all the secondary elements, except an additional C-terminal helix α5 in PseH (Fig. 3A). Furthermore, the mode of dimerization of PseH in the crystal is very similar to that of RimL [19] (Fig. 3C), although the second closest homologue (the acetyltransferase domain of MccE) is monomeric [16].

Further structural comparisons show that the PseH fold is very similar to the other members of the GNAT superfamily. Structural conservation of the GNAT fold has been related to its function as a scaffold for residues essential for AcCoA binding and catalysis [15]. In this respect, it is interesting to note that the structure of PseH is more similar to the GNAT enzymes that utilize amino acid sulfamoyl adenosine (MccE, RimL) or protein (RimL) as a substrate than a different GNAT-superfamily bacterial nucleotide-sugar N-acetyltransferase of the
known structure, the *E. coli* dTDP-fucosamine acetyltransferase WecD (Fig. 3D) [17]. Like PseH, WecD transfers an acetyl group from AcCoA to the 4-amino moiety of the nucleotide-linked sugar substrate. Structural comparison shows that WecD contains an extra 70-amino-acid domain at the N-terminus (Fig. 3D) and a different number and order of strands in the β-sheet of the GNAT-domain, 2345617 (as compared to 01234576 in PseH). Alignment of the structures of PseH and the GNAT-domain in WecD resulted in a match of only 124 Cα atoms with rms deviation of 2.9 Å and 10% identity over equivalence positions.

**Fig 2. The overall fold of *H. pylori* PseH.** (A) Stereo diagram of the structure of the PseH monomer. β-strands and α-helices are represented as arrows and coils and each element of the secondary structure is labeled and numbered as in text. The bound AcCoA molecule is shown in black. (B) The topology of secondary structure elements PseH. The α-helices are represented by rods and β-strands by arrows. Residue numbers are indicated at the start and end of each secondary structure element. (C) The molecular surface representation of PseH showing the AcCoA-binding tunnel between strands β4 and β5, which is a signature of the GNAT fold.

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Fig 3. Comparisons of PseH with other GNAT superfamily enzymes. (A) Stereo ribbon diagram of the superimposed structures of PseH from *H. pylori* (black), RimL from *S. typhimurium* (red) and the acetyltransferase domain of MccE from *E. coli* (green). The side chains of the conserved tyrosine in PseH...
A common mechanism of the acetyl transfer in GNAT enzymes involves protonation of the leaving thiolate anion of CoA by a general acid [15]. Previous mutagenesis studies were consistent with the role of Ser553 in MccE (structurally equivalent to Ser141 in RimL) as the general acid in catalysis [34]. In the superimposed structures of PseH, the MccE acetyltransferase domain and RimL, the side chain of Tyr138 of PseH is positioned close to that of Ser553 in MccE and Ser141 in RimL (Fig. 3A). Further structural superimpositions show that Tyr138 is structurally conserved in many GNAT superfamily transferases, including PA4794 from Pseudomonas aeruginosa (PDB ID 4KOW [18]), GNA1 from Saccharomyces cerevisiae (PDB ID 1I1D [20, 35]), sheep serotonin N-acetyltransferase (PDB ID 1L0C [36]) and human spermidine/spermine N1-acetyltransferase (PDB ID 2JEV [21]), where its role as a general acid in catalysis has been confirmed by mutagenesis. This suggests that Tyr138 acts as a general acid in the PseH-catalysed reaction.

Binding of AcCoA and localization of the putative active site

Analysis of the difference Fourier map revealed an AcCoA binding site between the splayed strands β4 and β5, which is the common cofactor site of GNAT superfamily enzymes (Figs. 2A and 4) [15–21]. The density for the entire molecule was readily interpretable, although somewhat less defined for the 3'-AMP moiety (Fig. 4). The position and extended conformation of AcCoA was found to be very similar to that described for other GNAT enzymes. The acetyl group of AcCoA is located at the bottom of the active site pocket on the face of the molecule opposite the AcCoA binding site. The pocket is lined with polar and aromatic residues. The carbonyl group of the thioester forms a bifurcated hydrogen bond with the main-chain amide of Ile93 (strand β4) and the hydroxyl of Tyr138, the putative general acid catalyst in the reaction. The acetyl moiety of AcCoA is further stabilized by van der Waals contacts with Leu91, Leu125 and Glu126. The β-alanine and β-mercaptoethylamine moieties are hydrogen bonded to the main-chain carbonyl of Ile93 (strand β4) and the side-chain of Asn131, and also interact through van der Waals contacts with Asn34, Trp38, Met39, Tyr94 and Ala134. The carbonyl oxygen of the pantoic acid moiety forms a hydrogen bond with the main-chain amide of Lys95 (strand β4), while the pyrophosphate group is stabilized by hydrogen bonds to the main chain of Gly103 and the side-chain of Lys133. The pattern of hydrogen bonds between the pantetheine moiety of AcCoA and strand β4 resembles bonding interactions in an antiparallel β-sheet, which is a common feature of GNAT enzymes.

Model for UDP-4-amino-4,6-dideoxy-β-L-AltNAc binding and implications for catalysis

The observed remarkable similarity between the overall folds of PseH, RimL and the acetyltransferase domain of MccE is consistent with their common ability to bind nucleotide-linked substrates. Indeed, analysis of the superimposition of the structures of PseH and the MccE acetyltransferase domain in complex with AcCoA and AMP revealed that the structural similarity extends to the architecture of the pocket that is occupied by the nucleotide moiety of the substrate in MccE (PDB ID 3R96 [16]) (Fig. 4A,B). In the crystal structure of the latter, the
The adenosine ring is sandwiched between Trp453 and Phe466, which are part of a largely hydrophobic pocket lined with residues change numbering here Leu436, Met451, Val493 and Trp511. Our analysis of the PseH structure revealed that many of the residues that form the corresponding pocket on the surface of PseH are structurally conserved between PseH and MccE. As Fig. 5 illustrates, the location and orientation of Val26, Met39, Phe52, Val76 and Tyr94 in PseH are similar to those of Leu436, Met451, Phe466, Val493 and Trp511 in MccE, respectively. The observed structural conservation of the nucleotide-binding pocket in PseH and MccE allowed us to model the nucleotide moiety of the UDP-4-amino-4,6-dideoxy-β-L-AltNAc substrate bound to PseH in a mode similar to that seen in MccE, with the uracil ring sandwiched between the side chains of Arg30 and Phe52 and forming face-to-face π-π stacking interaction with the aromatic ring of the latter (Fig. 5B).

Our structural analysis suggests that there are no residues in the vicinity of the AcCoA acetyl group that could serve as an acetyl acceptor and, thus, it is unlikely that the reaction proceeds through an enzyme-acetyl intermediate. The 4-amino-4,6-dideoxy-β-L-AltNAc moiety of the substrate has therefore been modeled next to the acetyl group of AcCoA, with the C4-N4 bond positioned optimally for the direct nucleophilic attack on the thioester acetate (Fig. 6) and in an orientation similar to that described for the functional homologue of PseH, WecD [17]. The model has been optimized to remove steric clashes and bring the bond length, bond angle and
torsion angle values close to ideal by using the structure idealization protocol implemented in Refmac [37].

Analysis of this model (Fig. 6) suggests that the pyrophosphate moiety makes minimal contacts with the protein. In contrast, the nucleotide- and 4-amino-4,6-dideoxy-β-L-AltNAc-binding pockets form extensive interactions with the substrate and are thus the most significant determinants of substrate specificity. Calculations of the surface area of the uracil and 4-amino sugar rings shielded from the solvent upon this interaction give the values of 55% and 48%, confirming good surface complementarity between the protein and the substrate in the model. Hydrogen bonds between the protein and the substrate involve the side-chains of Arg30, His49, Thr80, Lys81, Tyr94 and the main-chain carbonyl of Leu91 (Fig. 6). Van der Waals contacts with the protein involve Met39, Tyr40, Phe52, Tyr90 and Glu126. Notably, the 6'-methyl group of the altrose points into a hydrophobic pocket formed by the side-chains of Met39, Tyr40, Met129 and the apolar portion of the β-mercaptopethylamine moiety of AcCoA, which dictates preference to the methyl over the hydroxyl group and thus to contribute to substrate specificity of PseH.

The proposed catalytic mechanism of PseH proceeds by nucleophilic attack of the 4-amino group of the altrose moiety of the substrate at the carbonyl carbon of the AcCoA thioester.
group. In our model of the Michaelis complex, the C4-N4 bond lies directly over the acetyl group with the angle formed between the C4 of the amino-altrose, N4 of amino-altrose and the thioester carbonyl carbon being approximately 120° (Fig. 6). The model is therefore consistent with the geometry of approach required for nucleophilic attack by the substrate. At physiological pH, the 4-amino group of the unbound substrate is positively charged [15]. How does PseH promote its deprotonation, converting it into a nucleophile? Our analysis of the crystal structure of the PseH/AcCoA complex and the model of the Michaelis complex shows that there are no titratable side-chains in the vicinity of the thioester group or the 4-amino group of the modeled substrate that could be directly involved in deprotonation. However, we note that all three PseH subunits in the asymmetric unit contain a well-ordered water molecule that is hydrogen bonded to the side-chains of Ser78 and Thr80, and is located within a hydrogen-bond distance of the 3'-hydroxyl of the modeled 4'-amino-altrose, is represented as a grey-blue ball. Deprotonation of the substrate’s amine group may occur via the 3'-hydroxyl of the altrose and this intervening water molecule.
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Author Contributions
Conceived and designed the experiments: YCL AR. Performed the experiments: YCL AIU. Analyzed the data: AIU YCL AR. Wrote the paper: YCL AR.

References
1. World Gastroenterology Organisation (2011) World Gastroenterology Organisation global guideline: Helicobacter pylori in developing countries. J Dig Dis 12: 319–326. doi:10.1111/j.1751-2980.2011.00529.x PMID:21955424
2. Suerbaum S, Michetti P (2002) Helicobacter pylori infection. N. Engl. J. Med. 347, 1175–1186. PMID:12374879
3. Bauer B, Meyer TF (2011) The human gastric pathogen Helicobacter pylori and its association with gastric cancer and ulcer disease. Ulcers Article ID 340157, doi:10.1155/2011/340157
4. International Agency for Research on Cancer (1994) IARC Monogr Eval Carcinog Risks Hum 61: 177–240. PMID:7715070
5. Yoshiyama H, Nakazawa T (2000) Unique mechanism of Helicobacter pylori for colonizing the gastric mucus. Microbes Infect 2: 55–60. PMID:10717541
6. Ottemann KM, Lowenthal AC (2002) Helicobacter pylori uses motility for initial colonization and to attain robust infection. Infect Immun 70: 1984–1990. PMID:11895962
7. Baldwin DN, Shepherd B, Kraemer P, Hall MK, Sycuro LK, et al. (2007) Identification of Helicobacter pylori genes that contribute to stomach colonization. Infect Immun 75: 1005–1016. PMID:17101654
8. Josenhans C, Vossebein L, Friedrich S, Suerbaum S (2002) The neuA/flmD gene cluster of Helicobacter pylori is involved in flagellar biosynthesis and flagellin glycosylation. FEMS Microbiol Lett 210; 165–172. PMID:12044670
9. Schirm M, Soo EC, Aubry AJ, Austin J, Thibault P, et al. (2003) Structural, genetic and functional characterization of the flagellin glycosylation process in Helicobacter pylori. Mol Microbiol 48: 1579–1592. PMID:12791140
10. Schoenhofen IC, McNally DJ, Brisson J-R, Logan SM (2006) Elucidation of the CMP-pseudaminic acid pathway in Helicobacter pylori: synthesis from UDP-N-acetylgalactosamine by a single enzymatic reaction. Glycobiology 16: 8C–14C. PMID:16751642
11. Ishiyama N, Creuzenet C, Miller WL, Demendi M, Anderson EM, et al. (2006) Structural studies of FiaA from Helicobacter pylori reveal the mechanism for inverting 4,6-dehydratase activity. J Biol Chem 281: 24489–24495. PMID:16651261
12. Guerry P, Ewing CP, Schirm M, Lorenzo M, Kelly J, et al. (2006) Changes in flagellin glycosylation affect Campylobacter autoagglutination and virulence. Mol Microbiol 60: 299–311. PMID:16573682
13. Gough J, Karplus K, Hughey R, Chothia C (2001) Assignment of homology to genome sequences using a library of hidden Markov models that represent all proteins of known structure. J Mol Biol 313: 903–919. PMID:11697912
14. Liu YC, Ud-Din AI, Roujeinikova A (2014) Cloning, purification and preliminary crystallographic analysis of the Helicobacter pylori pseudaminic acid biosynthesis N-acetyltransferase PseH. Acta Cryst F 70: 1276–1279.
15. Vetting MW, De Carvalho LP, Yu M, Hegde SS, Magnet S, et al. (2005) Structure and functions of the GNAT superfamily of acetyltransferases. Arch Biochem Biophys 433: 212–226. PMID:15581578
16. Agarwal V, Meltitskaya A, Severinov K, Nair SK (2011) Structural basis for microcin C7 inactivation by the MccE acetyltransferase. J Biol Chem 286: 21295–21303. doi:10.1074/jbc.M111.226282 PMID:21507941
17. Hung M-N, Rangarajan E, Munger C, Nadeau G, Sulea T, et al. (2006) Crystal structure of TDP-fucosamine acetyltransferase (WecD) from Escherichia coli, an enzyme required for enterobacterial common antigen synthesis. J Bacteriol 188: 5606–5617. PMID:16855251
18. Majorek KA, Kuhn ML, Chruszcz M, Anderson WF, Minor W (2013) Structural, functional, and inhibition studies of a GCN5-related N-acetyltransferase (GNAT) superfamily protein PA4794: A new C-terminal lysine protein acetyltransferase from Pseudomonas aeruginosa. J Biol Chem 288: 30223–30235. doi: 10.1074/jbc.M113.501353 PMID: 24003232

19. Vetting MW, de Carvalho LP, Roderick SL, Blanchard JS (2005) A novel dimeric structure of the RimL Nα-acetyltransferase from Salmonella typhimurium. J Biol Chem 280: 22108–22114. PMID: 15817456

20. Peneff C, Mengin-Lecreulx D, Bourne Y (2001) The crystal structures of Apo and complexed Saccharomyces cerevisiae GNA1 shed light on the catalytic mechanism of an amino-sugar N-acetyltransferase. J Biol Chem 276: 16328–16334. PMID: 11278591

21. Hegde SS, Chandler J, Vetting MW, Yu M, Blanchard JS (2007) Mechanistic and structural analysis of human spermidine/spermine N1-acetyltransferase. Biochemistry 46: 7187–7195. PMID: 17516632

22. Olivier NB, Imperiali B (2008) Crystal structure and catalytic mechanism of PglD from Campylobacter jejuni. J Biol Chem 283: 27937–27946. doi: 10.1074/jbc.M801207200 PMID: 18667421

23. Battye TGG, Kontogiannis L, Johnson O, Powell HR, Leslie AGW (2011) iMOSFLM: a new graphical interface for diffraction-image processing with MOSFLM. Acta Cryst D 67: 271–281.

24. Evans PR, Murshudov GN (2013) How good are my data and what is the resolution? Acta Cryst D 69: 1204–1214.

25. Winn MD, Ballard CC, Cowtan KD, Dodson EJ, Emsley P, et al. (2011) Overview of the CCP4 suite and current developments. Acta Cryst D 67: 235–242.

26. Terwilliger TC, Adams PD, Read RJ, McCoy AJ, Moriarty NW, et al. (2009) Decision-making in structure solution using Bayesian estimates of map quality: the PHENIX AutoSol wizard. Acta Cryst D 65: 582–601.

27. Adams PD, Afonine PV, Bunkóczi G, Chen VB, Davis IW, et al. (2010) PHENIX: a comprehensive Python-based system for macromolecular structure solution. Acta Cryst D 66: 213–221.

28. Emsley P, Cowtan K (2004) Coot: model-building tools for molecular graphics. Acta Cryst D 60: 2126–2132.

29. Chen VB, Arendall WB, Headd JJ, Keedy DA, Immormino RM, et al. (2009) MolProbity: all-atom structure validation for macromolecular crystallography. Acta Cryst D 66: 12–21.

30. The PyMOL Molecular Graphics System, Version 1.2r3pre, Schrödinger, LLC.

31. Berman HM, Westbrook J, Feng Z, Gilliland G, Bhat TN, et al. (2000) The Protein Data Bank. Nucleic Acids Res 28: 235–242. PMID: 10592235

32. Krissinel E, Henrick K (2004) Secondary-structure matching (SSM), a new tool for fast protein structure alignment in three dimensions. Acta Cryst D 60: 2256–2268.

33. Kazakov T, Kuznedelov K, Semenova E, Mukhamedyarov D, Datsenko KA, et al. (2014) The RimL transacetylase provides resistance to translation inhibitors microcin C. J Bacteriol 196: 3377–3385. doi: 10.1128/JB.01584-14 PMID: 25002546

34. Novikova M, Kazakov T, Vondenhoff GH, Semenova E, Rozenski J, et al. (2010) MccE provides resistance to protein synthesis inhibitor microcin C by acetylating the processed form of the antibiotic. J Biol Chem 285: 12662–12669. doi: 10.1074/jbc.M109.080192 PMID: 20159968

35. Mio T, Yamada-Okabe T, Arisawa M, Yamada-Okabe H (1999) Saccharomyces cerevisiae GNA1, an essential gene encoding a novel acetyltransferase involved in UDP-N-acetylglucosamine synthesis. J Biol Chem 274: 424–429. PMID: 9867860

36. Scheinber NA, De Angelis J, Burley SK, Cole PA (2002) Investigation of the roles of catalytic residues in serotonin N-acetyltransferase. J Biol Chem 277: 18118–18126. PMID: 11884405

37. Murshudov GN, Vagin AA, Dodson EJ (1997) Refinement of macromolecular structures by the maximum-likelihood method. Acta Cryst D 53: 240–255.