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Structural and functional characterization of *M. tuberculosis* sedoheptulose-7-phosphate isomerase, a critical enzyme involved in lipopolysaccharide biosynthetic pathway

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M. tuberculosis GmhA enzyme catalyzes the isomerization of D-sedoheptulose 7-phosphate into D-glycero-D- α -manno-heptose-7-phosphate in GDP-D-glycero- α -D-manno-heptose biosynthetic pathway. The D-glycero- α -D-manno-heptose is a major constituent of lipopolysaccharide and contributes to virulence and antibiotic resistance to mycobacteria. In current study, we have performed the structural and biochemical analysis of *M. tuberculosis* GmhA, the first enzyme involved in D-sedoheptulose 7-phosphate isomerization in GDP-D- α -D-heptose biosynthetic pathway. The *Mtb*GmhA enzyme exists as tetramer and small angle X-ray scattering analysis also yielded tetrameric envelope in solution. The *Mtb*GmhA enzyme binds to D-sedoheptulose 7-phosphate with $K_m \sim 0.31 \pm 0.06 \text{ mM}^{-1}$ and converts it to D-glycero-D- α -manno-heptose-7-phosphate with catalytic efficiency (k_{cat}/K_m) $\sim 1.45 \text{ mM}^{-1} \text{ s}^{-1}$. The residues involved in D-sedoheptulose 7-phosphate and Zn^{2+} binding were identified using modeled *Mtb*GmhA + D-sedoheptulose 7-phosphate + Zn^{2+} structure. To understand the role in catalysis, six site directed mutants of *Mtb*GmhA were generated, which showed significant decrease in catalytic activity. The circular dichroism analysis showed $\sim 46\%$ α -helix, $\sim 19\%$ β -sheet and $\sim 35\%$ random coil structures of *Mtb*GmhA enzyme and melting temperature $\sim 53.5 \text{ }^\circ\text{C}$. Small angle X-ray scattering analysis showed the tetrameric envelope, which fitted well with modeled *Mtb*GmhA tetramer in closed conformation. The *Mtb*GmhA dynamics involved in D-sedoheptulose 7-phosphate and Zn^{2+} binding was identified using dynamics simulation and showed enhanced stability in presence of these ligands. Our biochemical data and structural knowledge have provided insight into mechanism of action of *Mtb*GmhA enzyme, which can be targeted for novel antibiotics development against *M. tuberculosis*.

The GDP-D-glycero- α -D-manno-heptose is a key building block of lipopolysaccharide in mycobacteria and blocking of its biosynthetic pathway leads to high antibiotic susceptibility and reduced virulence of mycobacteria. The enzymes involved in GDP-D- α -D-heptose biosynthetic pathway offer an attractive target for novel antibiotics development. The GDP-D- α -D-heptose in mycobacteria is synthesized in four steps, (1) isomerization of D-sedoheptulose 7-phosphate into D-glycero-D- α -manno-heptose-7-phosphate by GmhA enzyme (2) Phosphorylation of D-glycero-D- α -manno-heptose-7-phosphate at C1 position by HddA enzyme, which forms D-glycero-D- α -manno-heptose-1,7-bisphosphate (3) Removal of phosphate at C7 position in D-glycero-D- α -manno-heptose-1,7-bisphosphate by GmhB enzyme, which leads to D-glycero-D- α -manno-heptose-1-phosphate and (4) modification of the phosphate at C1 position to form a phosphodiester linkage with GDP by HddC enzyme and leads to GDP-D-glycero- α -D-manno-heptose (Fig. 1A). GDP-D- α -D-heptose is incorporated in the S-layer glycoproteins of mycobacterial membrane by specific precursor.

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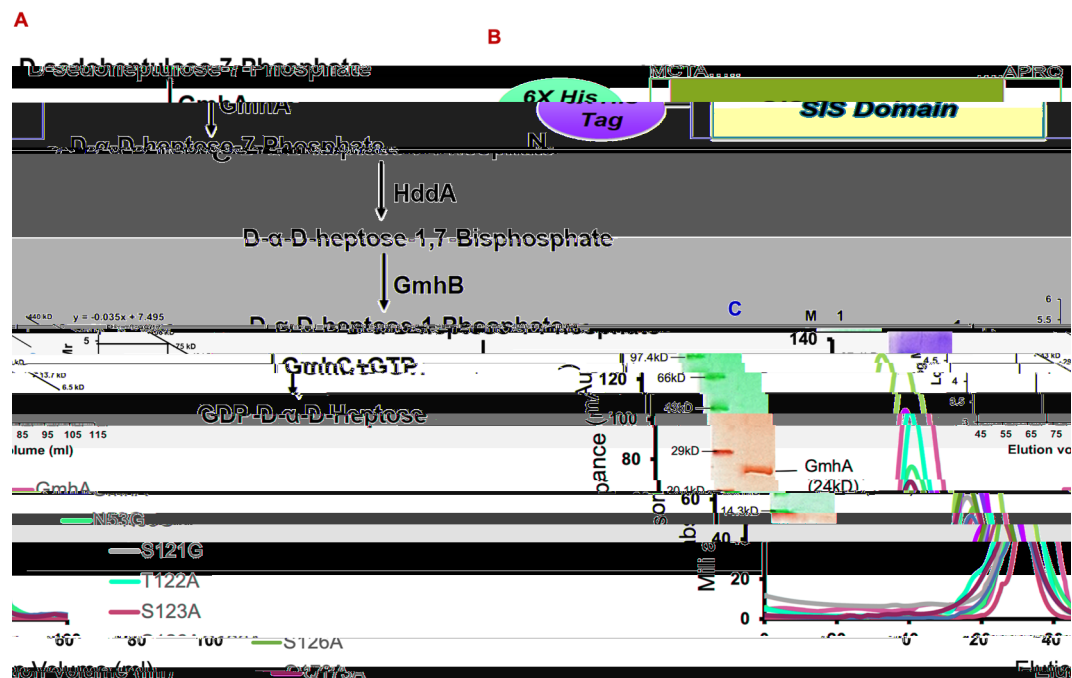


Figure 1. (A) Schematic diagram showing various enzymes involved in *M. tuberculosis* GDP-D- α -D-heptose biosynthetic pathway. (B) The gene construct used for *MtbGmhA* enzyme expression. SIS domain represents the sugar isomerase domain. (C) The elution profile of wild type and six mutants of *MtbGmhA* enzymes obtained from Superdex 200 column. Inset shows the SDS-PAGE of wild type enzyme; M = Mw marker, 1 = *MtbGmhA*. The molecular mass of *MtbGmhA* was calculated from standard curve generated using elution profiles of known molecular mass proteins.

The heptose sugar maintains the structural integrity of the outer membrane of bacteria and interacts with membrane proteins and divalent cations. The heptose sugar is observed in inner core of lipopolysaccharide of *E. coli* and *P. aeruginosa*¹. The heptose moiety is involved in ionic interactions and provides barrier to the passage of detergent, dyes and antibiotics through bacterial cell wall² and is essential for survival of *P. aeruginosa*³.

The Gram-positive bacterial species contain crystalline two dimensional protein arrays, known as S- layers⁴. In S-layers, protein clusters are covalently attached to glycan chains, made up of 20–50 identical repeating units of α -L-rhamnose and D-glycero- α -D-manno-heptose units. Glycosylation of bacterial surface proteins with heptose and rhamnose are significant and involved in many functions e. g. adherence⁵, evasion of host immune response and enhanced resistance to proteolytic attack.

The GDP-D- α -D-heptose biosynthetic pathway in *M. tuberculosis* was first discovered by Eidels and Osborn⁶. The *M. tuberculosis* is a Gram-positive organism, however exhibits both features of Gram-positive and Gram-negative bacteria⁷. The GmhA is the key enzyme and highly conserved in Gram-positive and Gram-negative organisms e. g. *A. thermoaerophilus* and *C. acetobutylicum*. The *MtbGmhA* showed 26.2% sequence identity with C-terminal region of L-glutamine-D-fructose-6-phosphateamido transferase, a ketose/aldose isomerase consists of sugar isomerase domain⁸. The D-glycero- α -D-manno-heptose is also observed as cellular component of *A. thermoaerophilus* DSM 10155, a member of the bacillus/clostridium group of Gram-positive organism⁹.

Crystal structures of GmhA enzymes from eight Gram-negative species have been determined^{10–13}. The GmhA enzymes from *P. aeruginosa* and *V. cholerae* were crystallized as dimer, while other GmhA orthologues were crystallized as tetramer in the asymmetric unit. The GmhA tetramer is found in two distinct conformations, “closed” and “open”. The “open” conformation is characterized by an extended α 3– β 2 loop, less ordered α 3' region, and loosely packed dimer-dimer interface. In “closed” conformation, unstructured α 3' region adopts a helical structure and α 3– β 2 loop is positioned inward, which allowed more extensive dimer-dimer interaction.

In current study, we have performed biochemical and structural analysis of *MtbGmhA* enzyme. The *MtbGmhA* enzyme was purified and its activity was analyzed by coupled assay involving *MtbGmhA*, *MtbHddA* and *MtbGmhB* enzymes. The binding analysis between all three enzymes was performed using surface plasmon resonance technique. The *MtbGmhA* residues involved in D-sedoheptulose 7-phosphate and Zn^{2+} binding were identified using modeled complex and their roles in catalysis were determined using site directed mutagenesis. The circular dichroism, small angle X-ray scattering, and dynamics simulation techniques were used to analyze the *MtbGmhA* structure and dynamics involved in D-sedoheptulose 7-phosphate and Zn^{2+} binding. Our structural and biochemical analysis on *MtbGmhA* explained the mechanism of action, which will contribute in the development of novel antibiotics against *M. tuberculosis*.

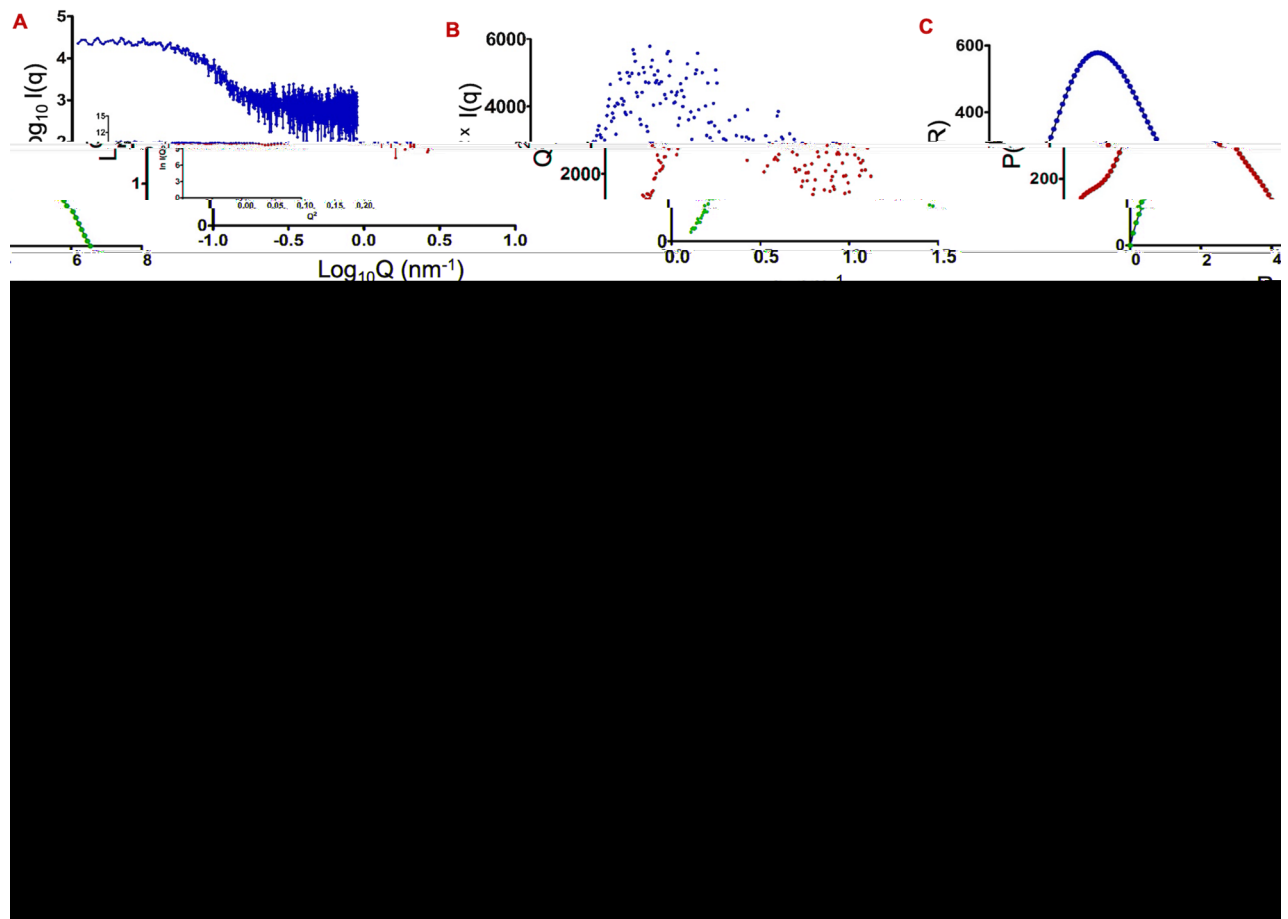


Figure 2. Small angle X-ray scattering analysis on *MtbGmhA* enzyme. (A) SAXS intensity profile of *MtbGmhA*. The inset shows linear fit to Guinier region of measured data. (B) Kratky plot of *MtbGmhA*. (C) The $P(R)$ curves computed for *MtbGmhA* indicating the frequency distribution of interatomic vectors in predominant scattering species. (D–F) Fitting of *MtbGmhA* tetramer (closed state) in shape restored from dummy atom modeling using small angle X-ray scattering data and shown after every 90° rotation along Y-axis. Fitting of *MtbGmhA* tetramer (open state) in shape restored from dummy atom modeling using small angle X-ray scattering data and shown after every 90° rotation.

Results and discussion

Purified *MtbGmhA* forms tetramer in solution. The Fig. 1A showed the various steps involved in GDP-D- α -D-heptose biosynthetic pathway in *M. tuberculosis*. The *MtbGmhA* enzyme (196 residues, Mw ~ 24 kDa) is the first enzyme involved in D-sedoheptulose 7-phosphate isomerization and converts it to D-glycero-D- α -manno-heptose-7-phosphate. The *MtbGmhA* gene was cloned in *pET28a* expression vector (Fig. 1B) and overexpressed in soluble fraction of *Escherichia coli*. The *MtbGmhA* enzyme was purified and eluted as tetramer (~ 100 kDa) from Superdex 200 (16/60) column, identified based on molecular mass standard (Fig. 1C, inset). The purified *MtbGmhA* showed more than 98% purity on SDS-PAGE (Fig. 1C, inset). The SDS-PAGE profile of 9 eluted fractions of *MtbGmhA* (E1–E9) are shown in Fig. S1. We have generated the six-point mutants e. g. N53G, S121G, T122G, S123G, S126G, Q173G of *MtbGmhA* enzyme and all mutant proteins were purified like wild type enzyme and eluted as tetramer (Fig. 1C). The primers used for gene amplification of six *MtbGmhA* mutants are shown in Table S1. PDBePISA server analysis also indicated that biological assembly of *MtbGmhA* enzyme is a tetramer¹².

The GmhA enzymes from *P. aeruginosa* and *V. cholera* form tetramer in solution^{10,14}. However, their crystallographic asymmetric unit contains dimer and functional tetramers were obtained using two-fold crystallographic symmetry. The *H. pylori* GmhA is the only homologue, which was observed as homodimer in solution^{15,16}.

Small angle X-ray scattering analysis revealed the tetrameric envelope of *MtbGmhA* enzyme.

The small angle X-ray scattering data was collected on *MtbGmhA* enzyme to analyze its low-solution shape in solution. The SAXS intensity profile $[I(Q)]$ as a function of momentum transfer vector Q , Guinier plot using globular approximation, Kratky plot and pair-distribution function $[P(R)]$ were calculated (Fig. 2A–C). The Fig. 2A showed the double logarithmic plot of small angle X-ray scattering data ($\log_{10} I(Q) \sim \log_{10} Q$). These data showed no inter particulate effect or aggregation of protein, as lack of upwards or downwards points were observed, when data approached to 0 nm^{-1} . The Fig. 2B showed the bell-shaped peak in Kratky plot, which

<i>MtbGmhA</i>	
Data collection parameters	
Instrument	SAX Space (Anton Paar)
Beam geometry	10 mm slit
Wavelength (Å)	1.5418
Q range (nm ⁻¹)	0.10–3.00
Exposure time (min)	30
Concentration (mg/ml)	1.22
Temperature (K)	283
Structural parameters	
I(0) (cm ⁻¹) [from P(R)]	25,650.00
R _g (nm) [from P(R)]	2.73 ± 0.11
I(0) (cm ⁻¹ from Guinier)	26,737.10 ± 492.71
R _g (nm) (from Guinier)	2.48 ± 0.17
D _{max} (nm)	7.9
Porod volume estimate (nm ³)	130.29
Molecular-mass determination	
Molecular mass M _r (kDa) I(0)	93.2
Calculated monomeric M _r from sequence (KDa)	24.105
Software employed	
Primary data reduction	SAXSquant
Data processing	PRIMUSQT
Ab initio analysis	DAMMIF
Validation and averaging	DAMAVR
Computation of model intensities	CRYOSOL
3D graphics representations	PyMOL

Table 1. Small angle X-ray scattering data collection and experimentally derived parameters of *MtbGmhA*. R_g = radius of gyration. D_{max} = maximum particle dimension.

indicated the globular shape of *MtbGmhA* in solution. The Guinier plot analysis yielded the radius of gyration R_G ~ 2.38 ± 0.11 nm, quite similar to theoretical R_G of *MtbGmhA* tetramer. The Fig. 2C showed the probability distribution of various interatomic vectors P(R), using small angle X-ray scattering data as a reference. These data yielded the maximum linear dimension (D_{max}) ~ 6.35 nm and R_G ~ 2.45 nm for *MtbGmhA* tetramer (Fig. 2C). The single peak in P(R) profile indicated the globular shape of *MtbGmhA* enzyme. Using Lysozyme data (conc. ~ 1 mg/ml), the intensity at zero angle (I₀) ~ 464 a. u. was observed for 1.0 kDa in 1 h exposure. Using it as control, the relative intensity of *MtbGmhA* protein (conc. ~ 1.2 mg/ml) has yielded the molecular mass ~ 93.2 kDa, which corresponds to *MtbGmhA* tetramer (Table 1).

Uniform density modeling protocol was used to restore the shape, which yielded 10 independent *MtbGmhA* models. All models were averaged, refined and yielded the normalized spatial disposition (NSD) ~ 1.30 ± 0.03, a measure of similarity between individual solutions. The NSD showed the high similarity between dummy residue shape of solved models. The modeled *MtbGmhA* tetramer in closed state fitted well into SAXS envelope (χ^2 ~ 1.17) (Fig. 2D–F), compared to open state (χ^2 ~ 3.29) (Fig. 2G–I). The CRYOSOL program was used to compute the theoretical small angle X-ray scattering profile from *MtbGmhA* tetrameric model and compared with experimental small angle X-ray scattering data. These data indicated that *MtbGmhA* exists as monodisperse, homogeneous and closed state tetramer in solution.

Activity assay using wild type and mutant *MtbGmhA* enzymes showed the roles of various active site residues.

Activity assay using wild type enzyme. For activity analysis, the *MtbHddA* and *MtbGmhB* enzymes were purified (data not shown), performed the coupling assay using all three enzymes and monitored the release of phosphate. As control, a phosphate standard curve was calculated using 0–50 μM phosphate, dissolved in coupling reaction buffer having no *MtbGmhA* enzyme. Following kinetic parameters, K_m ~ 0.31 ± 0.06 mM⁻¹, k_{cat} ~ 0.45 ± 0.02 s⁻¹ and catalytic efficiency (k_{cat}/K_m) ~ 1.45 mM⁻¹ s¹ were obtained for *MtbGmhA* enzyme (Fig. 3A, Table 2).

The *MtbGmhA* kinetic parameters were compared with other GmhA orthologues, as shown in Table 2. The catalytic efficiency of *MtbGmhA* enzyme was quite similar to *B. pseudomallei* and *E. coli* GmhA enzymes. However, *H. pylori* GmhA enzyme showed ~ 50-fold higher catalytic efficiency (k_{cat}/K_m) than *MtbGmhA* enzyme and other GmhA orthologues. The *H. pylori* GmhA was observed as dimer in solution, while tetramers were observed for GmhA enzymes from *M. tuberculosis*, *E. coli* and *B. pseudomallei*. It is not clear how oligomeric state of GmhA enzyme affects its catalytic efficiency.

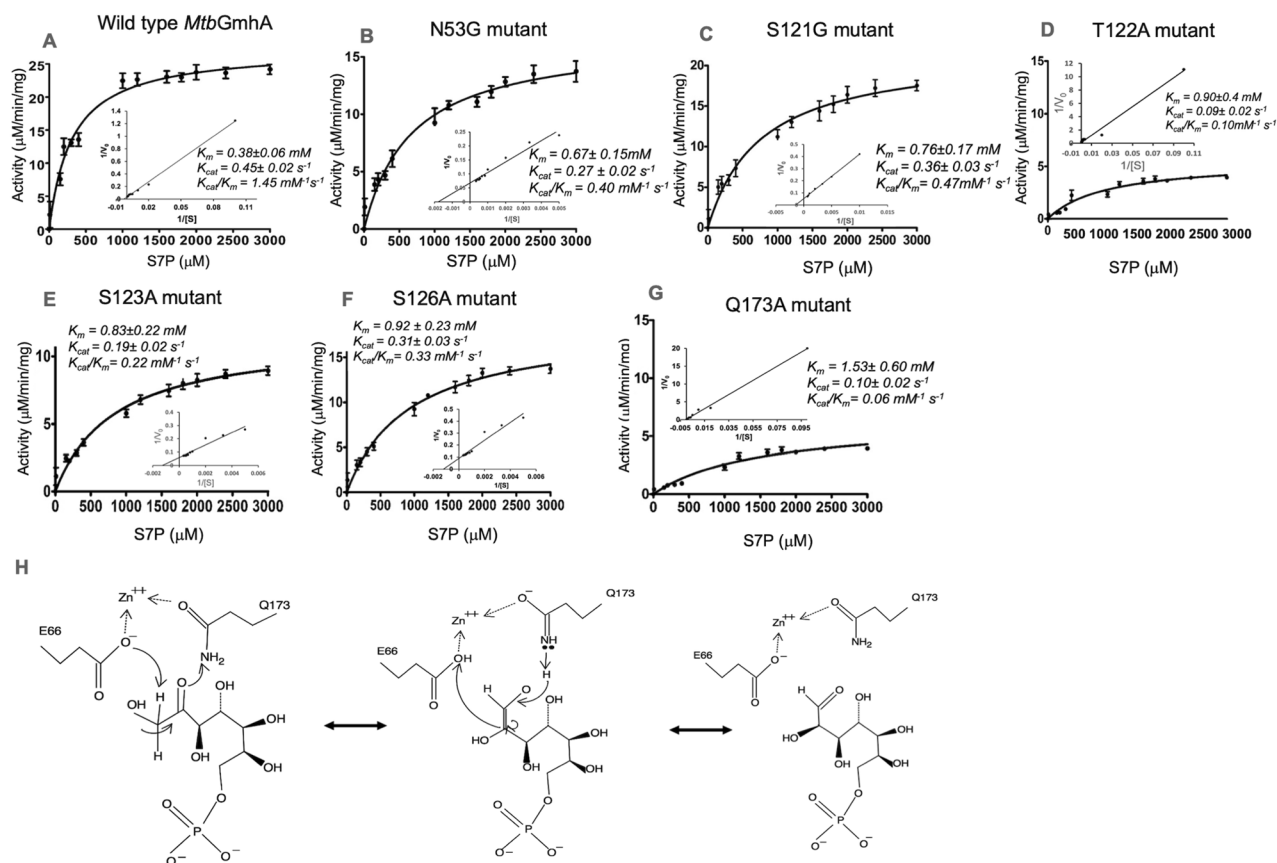


Figure 3. Activity assay on wild type and six mutants of *MtbGmhA* enzymes. Three replicates were performed for each measurement. The *MtbGmhA* activity was determined using Malachite green phosphate detection kit. A coupling reaction was performed using *MtbGmhA*, *MtbHddA* and *MtbGmhB* enzymes and release of inorganic phosphate was monitored at 600 nm. Following saturation curves were obtained, (A) wild type *MtbGmhA* (B) N53G mutant (C) S121G mutant (D) T122A mutant (E) S123A mutant (F) S126A mutant and (G) Q173A mutant. The 0 to 3 mM concentrations of D-sedoheptulose 7-phosphate were used in all assays and results are given in Table 2. (H) Proposed mechanism of *MtbGmhA* used in isomerization of D-sedoheptulose 7-phosphate to D-glycero-D- α -manno-heptose-7-phosphate through enediol intermediate, in which Gln173 acted as a catalytic acid and forms an imine stabilized by Zn^{2+} ion and Glu65 acted as catalytic base.

Protein	K_m (mM)	k_{cat} (s^{-1})	k_{cat}/K_m ($\text{mM}^{-1} \text{s}^{-1}$)
<i>MtbGmhA</i>	0.31 ± 0.06 (0.38)	0.45 ± 0.02 (0.52)	1.45 (1.36)
N53G mutant	0.67 ± 0.15 (0.62)	0.27 ± 0.02 (0.27)	0.40 (0.43)
S121G mutant	0.76 ± 0.17 (0.73)	0.36 ± 0.03 (0.33)	0.47 (0.45)
S123A mutant	0.92 ± 0.23 (0.93)	0.31 ± 0.03 (0.33)	0.33 (0.35)
S126A mutant	0.83 ± 0.22 (0.83)	0.19 ± 0.02 (0.18)	0.22 (0.21)
T122A mutant	0.90 ± 0.4 (0.89)	0.09 ± 0.02 (0.11)	0.10 (0.12)
Q173A mutant	1.53 ± 0.6 (1.54)	0.10 ± 0.02 (0.13)	0.06 (0.08)
<i>E. coli GmhA</i>	0.9 ± 0.3	0.4 ± 0.7	0.5
<i>B. pseudomallei GmhA</i>	0.4 ± 0.2	0.5 ± 0.1	1.2
<i>H. pylori GmhA</i>	0.5 ± 0.1	33.1 ± 2.1	71.9

Table 2. Kinetic parameters of wild type and six *MtbGmhA* mutants obtained using Michaelis Menten curve and Lineweaver–Burk plot (in red bracket). The kinetic parameters of *E. coli GmhA*, *B. pseudomallei GmhA* and *H. pylori GmhA* are shown for comparative analysis.

Activity assay using six *MtbGmhA* mutants. We build the *MtbGmhA* + S7P + Zn^{2+} complex and identified the six residues involved in D-sedoheptulose 7-phosphate binding. we have generated the six *MtbGmhA* mutants, performed coupling assay and release of free P_i was monitored (Fig. 3B–G, Table 2).

As seen in Table 2, Gln173 \rightarrow Ala mutation leads to ~ 24 fold decrease in catalytic efficiency and \sim fivefold decrease in binding affinity to D-sedoheptulose 7-phosphate substrate in *MtbGmhA*, when compared to wild

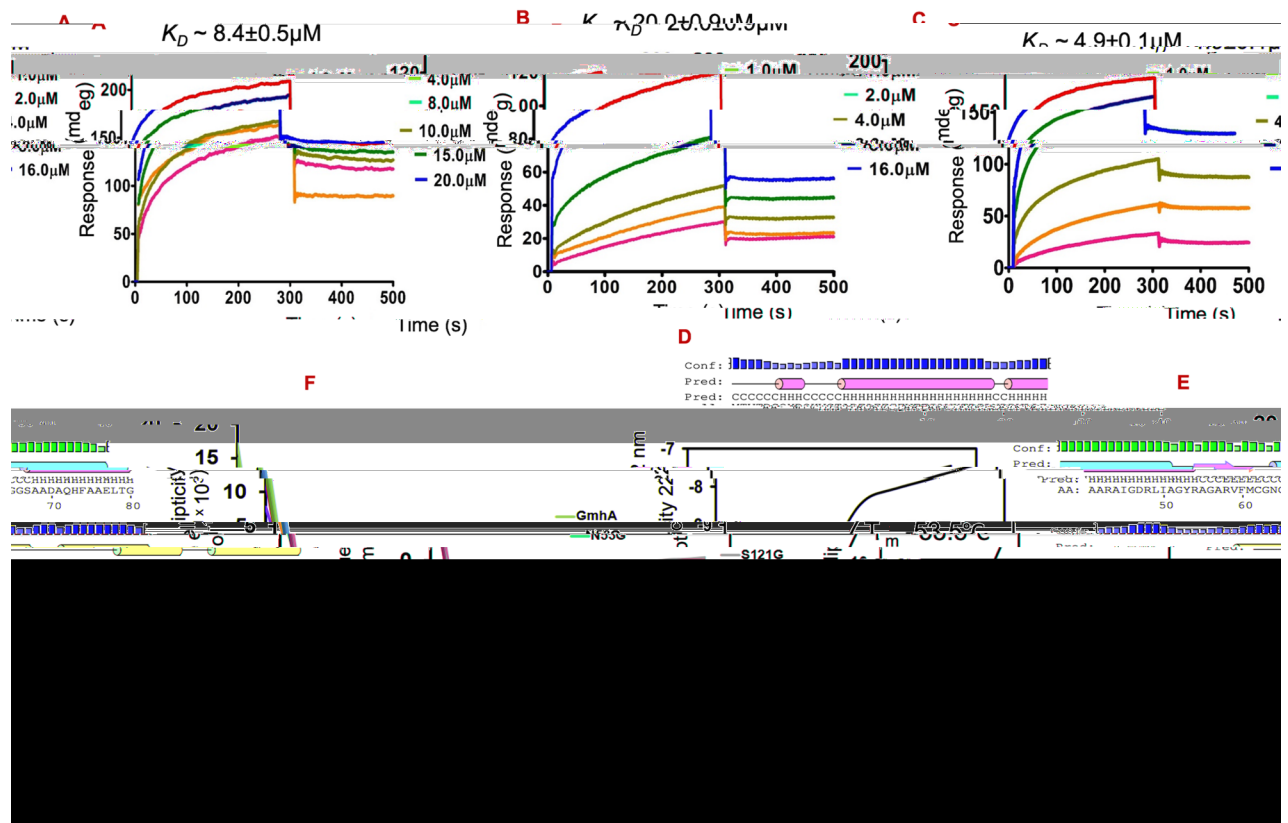


Figure 4. Interaction analysis between *MtbGmhA*, *MtbGmhB* and *MtbHddA* enzymes using surface plasmon resonance technique. (A) Sensogram showing the binding of *MtbHddA* on immobilized *MtbGmhA*. Five different concentrations of *MtbHddA* enzyme (1–16 μM) were used in this assay, which yielded the $K_D = 8.45 \pm 0.45 \mu\text{M}$. (B) Sensogram showing binding of *MtbHddA* on immobilized *MtbGmhB*. Five different concentrations of *MtbGmhB* (1–16 μM) were used in this assay, which yielded the $K_D = 20 \pm 0.95 \mu\text{M}$. (C) Sensogram showing the binding of *MtbGmhB* on immobilized *MtbGmhA*. Five different concentrations of *MtbGmhB* (4 to 20 μM) were used in current assay, which yielded $K_D = 4.9 \pm 0.11 \mu\text{M}$. (D) Psi-Pred program analysis on *MtbGmhA* sequence showing the secondary structural contents in *MtbGmhA* enzyme. (E) The circular dichroism data of wild type and six *MtbGmhA* mutants, shown in different colors. (F) The thermal denaturation profile on *MtbGmhA* enzyme, indicating the melting temperature $\sim 53.5^\circ\text{C}$ of the enzyme.

enzyme. Following data e. g. Asn53 \rightarrow Gly mutation (\sim twofold decrease in binding affinity and ~ 3.5 fold decrease in catalytic efficiency), Ser121 \rightarrow Gly mutation (\sim twofold decrease in binding affinity and ~ 3.5 fold decrease in catalytic efficiency), Thr122 \rightarrow Ala mutation (\sim threefold decrease in binding affinity and ~ 14.5 fold decrease in catalytic efficiency), Ser123 \rightarrow Ala mutation (\sim threefold decrease in binding affinity and ~ 4.5 fold decrease in catalytic efficiency), Ser126 \rightarrow Ala mutation (\sim threefold decrease in binding affinity and ~ 4.3 fold decrease in catalytic efficiency) were observed. The Asn53 forms bifurcated hydrogen bonds with O_4 and O_9 atoms of D-sedoheptulose 7-phosphate and Asn53 \rightarrow Gly mutation leads to ~ 3.5 fold decrease in catalytic efficiency. The Gln173 forms hydrogen bond with O_2 atom of D-sedoheptulose 7-phosphate and its mutation to Ala leads to ~ 24 fold decrease in catalytic efficiency. The bar diagram showing the catalytic efficiency of wild type and six *MtbGmhA* mutants are shown in Fig. S2.

In *Escherichia coli* and *P. aeruginosa* GmhA structures¹⁰, the Glu68 and His183 acted as acid and base in enzyme catalysis, which converted the D-sedoheptulose 7-phosphate to D-glycero-D- α -manno-heptose-7-phosphate. In *B. pseudomallei* GmhA structure¹², His183 was buried behind Zn^{2+} in the active site and Glu68 and Gln175 bind to Zn^{2+} in such a way, that their side chains acted as acid and base in enzyme catalysis.

In modeled *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn^{2+} tetramer, the Glu66 from one monomer and Gln173 from another monomer bind to D-sedoheptulose 7-phosphate and Zn^{2+} and may act as acid and base in enzyme catalysis (Fig. 3H). Mutation of Gln173 \rightarrow Ala has shown ~ 50 fold decrease in catalytic efficiency of *MtbGmhA* enzyme, indicating its involvement in enzyme catalysis. Similar mechanism has been observed in other GmhA orthologues, which contain Zn^{2+} in the active site.

***MtbGmhA*, *MtbHddA* and *MtbGmhB* enzymes interact each other in μM range.** Surface plasmon resonance technique was used to analyze the binding affinity between *MtbGmhA*, *MtbHddA* and *MtbGmhB* enzymes. Following K_D values were observed e. g. $\sim 8.4 \pm 0.5 \mu\text{M}$ between *MtbGmhA* and *MtbHddA* (Fig. 4A), $20.0 \pm 0.9 \mu\text{M}$ between *MtbHddA* and *MtbGmhB* (Fig. 4B) and $4.9 \pm 0.1 \mu\text{M}$ between *MtbGmhA* and *MtbGmhB*

Protein	α -helix (%)	β -sheet (%)	Random coil (%)
N53G mutant	46	21	33
S121G mutant	47	17	37
T122A mutant	38	16	46
S123A mutant	39	15	46
S126A mutant	40	17	43
Q173A mutant	39	16	45

Table 3. Secondary structural contents in six mutants of *MtbGmhA* obtained using circular dichroism spectroscopy.

(Fig. 4C). These data indicated that all three enzymes bind each other in μ M range and form a stable complex in GDP-D- α -D-heptose biosynthetic pathway.

The genetic organization in D- α -D-heptose biosynthetic pathway was first described in *A. thermoaerophilus* DSM 10155, in which all enzymes were located as cluster containing genes involved in synthesis and transfer of dTDP-rhamnose⁹. In *M. tuberculosis*, the genes for two different pathways were present at different locations and involved in synthesis of glycolipids and protein glycosylation. Interestingly, in CDC 1551 strain of *M. tuberculosis*, the GmhA and GmhB genes were fused as single gene, probably encoding a bifunctional enzyme having isomerase and phosphatase activities.

Circular dichroism analysis revealed the secondary structures and thermal stability of *MtbGmhA* enzyme. The PSIPRED program analysis on *MtbGmhA* sequence showed the secondary structures of enzyme (Fig. 4D). The Far-UV CD data on *MtbGmhA* and its six mutants were collected in 260–200 nm range (Fig. 4E). The K2D program has yielded the \sim 46% α -helix, \sim 19% β -sheet and \sim 35% random coil structures of wild type *MtbGmhA*, quite similar to structures observed in six *MtbGmhA* mutants (Fig. 4E, Table 3). Various secondary structure prediction programs have yielded quite similar secondary structures in *MtbGmhA* enzyme, as observed in CD analysis (Table S2).

For thermal stability analysis, the mean residue ellipticity (θ_{222}) data was collected on *MtbGmhA* enzyme in 25–90 °C range with 10 °C step (Fig. 4F). The helical structure of *MtbGmhA* enzyme was quite stable till 42 °C and disordered at 65 °C. A melting temperature, $T_m \sim$ 53.5 °C was obtained, which indicated the high thermostability of *MtbGmhA* enzyme.

The modeled *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn^{2+} tetramer showed the active site involved in D-sedoheptulose 7-phosphate and Zn^{2+} binding. The *MtbGmhA* model (1–196 residues) was obtained using I-TASSER (Iterative Threading ASSEMBLY Refinement) server, which used the PDB-2X3Y (Crystal structure of *B. pseudomallei* GmhA tetramer in closed state¹²) as the best input template (RMSD = 0.79, id1 = 0.39, id2 = 0.39, Conv = 0.98, Z-score = 3.2). The I-TASSER server also used the PDB-2I2W (Crystal structure of *Escherichia Coli* phosphoheptose isomerase in open state¹⁰) as another template for *MtbGmhA* modeling. However, current template did not yield suitable *MtbGmhA* model, as having following parameters (RMSD = 2.1, id1 = 0.35, TM-score = 0.82, Conv = 0.89). The D-sedoheptulose 7-phosphate and Zn^{2+} ions were docked into *MtbGmhA* tetramer using GLIDE module of Schrodinger program (Fig. 5A). A docking score of -5.44 and X-score of 8.1 were observed in docking analysis. The *P. pseudomallei* GmhA structures (PDB-2X3Y and PDB-2XBL) were used as reference to validate the results obtained in docking analysis of both ligands.

The *MtbGmhA* monomer consists of central five stranded parallel β -sheets, flanked by eight α -helices and forms a helix-beta-helix sandwich (Fig. 5A). Overall fold of *MtbGmhA* enzyme was quite similar to flavodoxin-type nucleotide-binding motif, as observed in other GmhA orthologues. The Electrostatic surface of *MtbGmhA* monomer (Fig. 5B) showed the positively charged surface at catalytic cleft, which accommodates the D-sedoheptulose 7-phosphate and Zn^{2+} ion. The LIGPLOT v.4.5.3 analysis of *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn^{2+} complex (Fig. 5C) showed that D-sedoheptulose 7-phosphate forms hydrogen bonds with Asn53 and Gly55 residues of β 1– α 3 helix region, Ser121, Thr122, Ser123 and Ser126 residues from β 3– α 6 helix region, Gln173 residue from α 8 helix in *MtbGmhA* enzyme. The Zn^{2+} ion forms hydrogen bond with Gln173 of *MtbGmhA*, which stabilize the D-sedoheptulose 7-phosphate in the active site. The *C. jejuni* and *P. aeruginosa* GmhA enzymes also adopt the closed structures¹⁴ and metal binding sites were observed in both enzymes. It appears that Zn^{2+} binding induces the closed state conformation of GmhA and more suitable for D-sedoheptulose 7-phosphate binding and catalysis.

Since *MtbGmhA* is observed as tetramer in solution, we build the *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn^{2+} tetramer using the same PDB-2X3Y as input template (Fig. 5D). The *MtbGmhA* tetramer forms a compact structure, in which four α -helices form hexagons, tilted at 120° and may help in structural stabilization. The α 4 and α 5 helices of each *MtbGmhA* monomer facing each other and forms a compact groove. 50% of α 1 helix of one monomer interacts with α 1 helix of the neighboring monomer. The Electrostatic surface of the *MtbGmhA* tetramer (Fig. 5E) showed the positively charged surface at catalytic pockets involved in D-sedoheptulose 7-phosphate and Zn^{2+} ion binding. Overall surface of *MtbGmhA* tetramer was negatively charged, except D-sedoheptulose 7-phosphate and Zn^{2+} binding pockets. The loop regions connecting β 1– α 3 and β 3– α 6 strands were involved in formation of D-sedoheptulose 7-phosphate binding pocket in *MtbGmhA* enzyme. The Fig. 5F

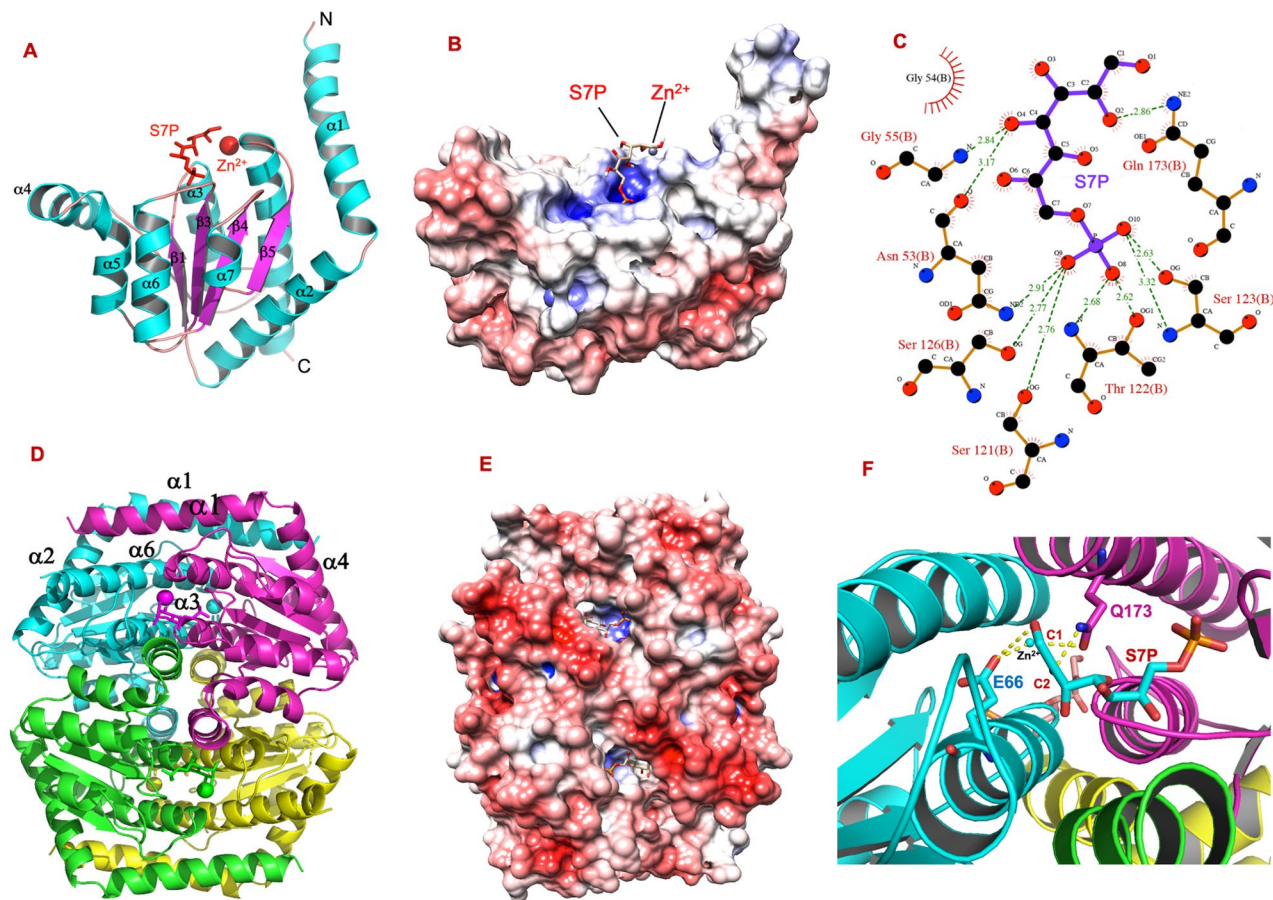


Figure 5. (A) The *MtbGmhA* + D-sedoheptulose 7-phosphate (S7P) + Zn^{2+} complex model showing the α -helices (in cyan), β -sheets (in magenta), loops (in orange). The D-sedoheptulose 7-phosphate and Zn^{2+} ion are shown in red colors. (B) Electrostatic surface diagram of *MtbGmhA* monomer showing the D-sedoheptulose 7-phosphate and Zn^{2+} ion in the active site (blue positive surface). (C) LigPlot analysis of *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn^{2+} complex showing the hydrogen bonds and van der Waals interactions between D-sedoheptulose 7-phosphate and Zn^{2+} with active site residues of *MtbGmhA* enzyme. The green dashed lines showed the hydrogen bonds and residues in “arc with spikes” are involved in van der Waals interactions with ligands. (D) *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn^{2+} tetramer showing each monomer in different color and D-sedoheptulose 7-phosphate and Zn^{2+} ligands in the active site of each monomer. (E) Electrostatic surface diagram of *MtbGmhA* tetramer showing four catalytic clefts (blue, positive charge), which accommodate the D-sedoheptulose 7-phosphate and Zn^{2+} in their binding pockets. (F) Interface between two *MtbGmhA* monomers, in which Glu66 residue (cyan) from one monomer and Zn^{2+} and Gln173 residue from another monomer binds to O1-C1 and O2-C2 of D-sedoheptulose 7-phosphate respectively and proposed as involved in isomerization of D-sedoheptulose 7-phosphate to D-glycero-D- α -manno-heptose-7-phosphate.

showed the Glu66 residue of one monomer (cyan) bind to O1-C1 of D-sedoheptulose 7-phosphate and Gln173 residue of another monomer (Pink) involved in binding to O2-C2 of D-sedoheptulose 7-phosphate and Zn^{2+} and involved in enzyme catalysis.

Differences between *MtbGmhA* and other known GmhA enzymes. *Sequence alignment and active site analysis.* The Fig. 6A showed the sequence alignment of *MtbGmhA* enzyme with 11 GmhA orthologues. The secondary structures of *MtbGmhA* model were placed on the top of sequence alignment. The Asn53, Gly55, Ser121, Thr122, Ser123, Ser126 residues (*) involved in D-sedoheptulose 7-phosphate and Zn^{2+} binding were found quite conserved. The active site residues e. g. Gln173 from one monomer and Glu66 from another monomer (shown as #), which bind to C1-O1 and C2-O2 of D-sedoheptulose 7-phosphate were also quite conserved. Other *MtbGmhA* residues involved in substrate binding e. g. Asn53 (fully conserved), Gly55 (G \rightarrow T in 2YVA, G \rightarrow S in 3TRJ), Ser121 (S \rightarrow T in 3TRJ), Thr122 (fully conserved), Ser123 (S \rightarrow R in 2YVA), Ser126 (fully conserved), Gln173 (fully conserved) and Glu66 (E \rightarrow S in 2YVA, E \rightarrow Q in 5BY2 and E \rightarrow K in 3TRJ) have shown degree of conservation with 11 other GmhA orthologues. In *Francisella tularensis*, Ser121 was replaced with Thr121, however not involved in catalytic cleft formation^{15,16}. Similarly, Ser123 \rightarrow Arg mutation in DiaA helps in timely initiation of chromosomal replication during cell cycle. In DiaA structure, important residues involved in enzyme catalysis were Ser52, Ala53, Arg71, Pro72, Asn83, Lys101, Leu190, and Phe191¹⁷.

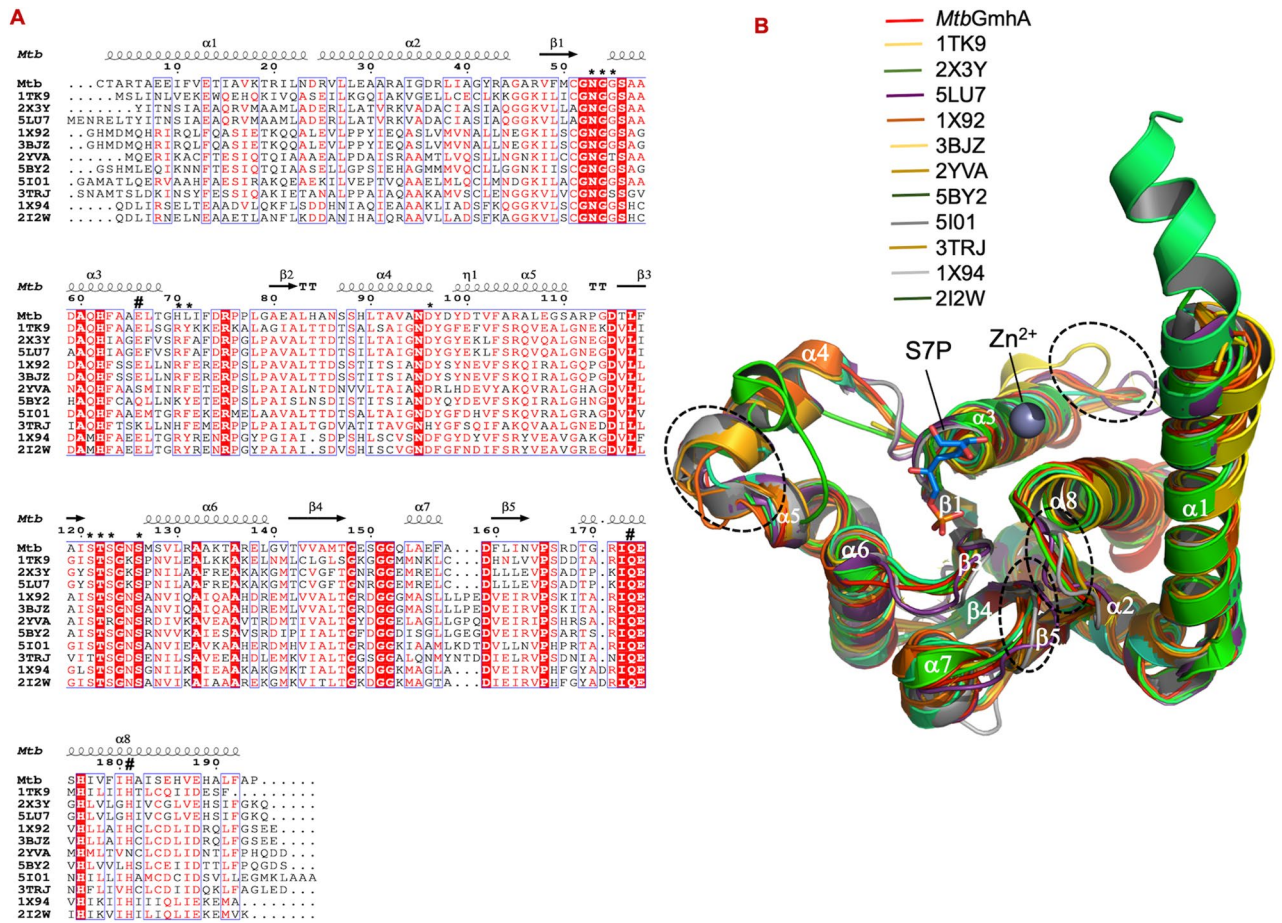


Figure 6. (A) Multiple sequence alignment of *MtbGmhA* with eleven GmhA orthologues, which showed highly conserved (red shade), semi-conserved (red letter) and dispersive residues (black letter). The residues involved in D-sedoheptulose 7-phosphate binding (*), in Zn²⁺ binding and enzyme catalysis (#) are shown above the sequence alignment. (B) Structural superposition of 11 GmhA orthologues on *MtbGmhA* model. These structures are 1TK9, 2X3Y, 5LU7, 1X92, 3BJZ, 5BY2, 5I0L, 3TRJ, 1X94, 112W. The 2YVA is not a *MtbGmhA* orthologue, but a homologue with different function. Major conformational changes are highlighted with dashed circles.

In addition, other *MtbGmhA* residues (Gly52, Gly54, Ala60, His62, Arg75, Asn95, Asp115, T116 (T → V in 11 enzymes), L117, G124, A132, A136, G148, Gly151, Gly152, Asp159, Pro165, Ile172, Glu174, His176) were fully conserved in sequence alignment with 11 GmhA orthologues.

Structural superposition. Structural superposition of 11 GmhA orthologues on *MtbGmhA* model have yielded following r.m.s. deviation e. g. 1TK9 (Crystal structures of two phosphate isomerases from *C. jejuni*)¹⁴ (0.65 Å, 146 Ca atoms), 2X3Y (Crystal structure of GmhA from *B. pseudomallei*)¹² (0.27 Å, 168 Ca atoms), 5LU7 (Crystal structure of heptose isomerase GmhA mutants)¹⁸ (0.15 Å, 168 Ca atoms), 1X92 (Crystal structure of *Pseudomonas Aeruginosa* phosphoheptose isomerase in complex with reaction product D-glycero-D-mannopyranose-7-Phosphate)¹⁰ (0.56 Å, 148 Ca atoms), 3BJZ¹⁰ (0.75 Å, 133 Ca atoms), 2YVA¹⁷ (Crystal structure of *E. coli* DiaA) (0.76 Å, 147 Ca atoms), 5BY2¹¹ (Sedoheptulose 7-phosphate isomerase from *Colwellia psychrerythraea* strain 34H) (0.63 Å, 147 Ca atoms), 5I0L (Crystal structure of Nucleoside Diphosphate kinase from *Schistosoma mansoni*)¹⁹ (0.53 Å, 147 Ca atoms), 3TRJ (Crystal structure of phosphoheptose isomerase from *Francisella tularensis*)^{15,16}, 1X94 (Crystal structure of two putative phosphoheptose isomerase from *Vibrio cholerae*)¹⁴ (0.64 Å, 116 Ca atoms), 112W (Crystal structure of *Bacillus licheniformis* BS3 class-A beta-lactamase and acyl-enzyme adduct formed with cefoxitin)²⁰ (0.69 Å, 128 Ca atoms) (Fig. 6B). Major changes were observed in the loop regions connecting α3–β2, α8–β5, β5–α6 strands and minor changes in other loops of *MtbGmhA* enzyme. The β1–α3 loop is involved in D-sedoheptulose 7-phosphate and Zn²⁺ binding and found quite conserved in 11 GmhA orthologues and showed least conformational change. Major conformational changes were observed in α4 and α7 helices of *E. coli* GmhA enzyme (2I2W, open state) to *MtbGmhA*, when compared to structures of 10 GmhA orthologues.

Dynamic simulation on *MtbGmhA* tetramer as (1) Apo (2) D-sedoheptulose 7-phosphate bound and (3) D-sedoheptulose 7-phosphate + Zn²⁺ bound state has revealed the dynamics

Models	Apo	D-S7P	D-S7P + Zn ²⁺
	(Tetramer)	(Tetramer)	(Tetramer)
Protein atoms	11,444	11,444	11,444
Water/ions	35,201(28)	23,372(36)	23,379(28)
Ligand atoms	–	88	88(S7P), 4 (Zn ²⁺)
Total atoms	117,075	81,684	81,701
Minimization (Steep)	50,000	50,000	50,000
NVT equilibration	100 ps	100 ps	100 ps
NPT equilibration	100 ps	100 ps	100 ps
Time steps (fs)	2	2	2
No. of steps	500,000	500,000	500,000
Simulation (ns)	1	1	1

Table 4. Dynamics simulation parameters of Apo, D-sedoheptulose 7-phosphate (D-S7P) bound and D-sedoheptulose 7-phosphate + Zn²⁺ (D-S7P + Zn²⁺) bound *MtbGmhA* tetramer.

involved in ligand recognition. 1 ns dynamics simulation was performed on *MtbGmhA* tetramer in (i) Apo (ii) D-sedoheptulose 7-phosphate bound and (iii) D-sedoheptulose 7-phosphate + Zn²⁺ bound state and analyzed the dynamics involved in D-sedoheptulose 7-phosphate and Zn²⁺ binding (Table 4). All three *MtbGmhA* tetramers obtained after dynamics simulation showed good stereochemistry and lie in allowed regions of Ramachandran plot (Figs. S3–S5).

Dynamics simulation on *MtbGmhA* tetramer in absence of ligand. To investigate the effect of ligand binding on dynamics fluctuation of *MtbGmhA* tetramer, we have performed the dynamics simulation on *MtbGmhA* tetramer in absence of ligand. Superposition of simulated *MtbGmhA* tetramer (Green) on starting structure (Grey) have yielded RMSD ~1.3 Å for 705 Ca atoms, indicating quite similar structure, except minor changes were observed in the loop regions of protein (Fig. 7A). High B-factors (~40–60 Å²) were observed for amino acid stretches, 20–30, 50–60, 70–80, 105–115, 120–130, 150–155 and 160–170 of *MtbGmhA* tetramer (Fig. 7B). The initial RMSD of *MtbGmhA* tetramer was ~0.0 and increased slightly (~0.2 nm) during 1 ns simulation (Fig. 7G). The radius of gyration (R_g) in *MtbGmhA* tetramer was quite stable during simulation period and found ~2.6 nm (Fig. 7H).

Dynamics simulation on *MtbGmhA* tetramer in complex with D-sedoheptulose 7-phosphate. 1 ns dynamics simulation was performed on *MtbGmhA* + D-sedoheptulose 7-phosphate tetramer to analyze the effect of substrate binding on *MtbGmhA* structure. Superposition of the simulated *MtbGmhA* + D-sedoheptulose 7-phosphate structure (red) on starting complex structure (grey) have yielded the RMSD ~1.1 Å for 740 Ca atoms (Fig. 7C). It showed that overall structure of *MtbGmhA* tetramer remains unperturbed after dynamics simulation, except minor changes in the loop regions of enzyme. The D-sedoheptulose 7-phosphate substrate was quite stable during simulation period. To examine, whether structural deviations occurred in local regions, or throughout the whole structure, we computed the B-factor plot for *MtbGmhA* tetramer during entire simulation period (Fig. 7D). The amino acid stretches 20–30, 70–80 and 150–155 showed the higher B-factor (~40–60 Å²) (Fig. 7D). The *MtbGmhA* residues *e.g.*, Asn53, Gly55 residues involved in C5-O5 binding and Ser121, Thr122, Ser123, Ser126 residues are involved in C7-PO₄ binding have shown significant decrease in B factor compared to wild-type *MtbGmhA* tetramer. The active site Glu66 and Gln173 residues of *MtbGmhA* showed low B-factor <20 Å² compared to 20–60 Å² observed in wild type enzyme. These residues bind to C1–O1 and C2–O2 of D-sedoheptulose 7-phosphate and involved in catalysis. Other residues, 20–30, 70–80 and 150–155 showed higher B-factors, though not involved directly in substrate binding.

Figure 7G showed the RMSD of *MtbGmhA* sampled through dynamics simulation from starting structure. Minor structural changes were observed in 1 ns simulation and reached to final ~7 Å. The radius of gyration (R_g) of complex was quite stable and found ~2.2 nm (Fig. 7H). These values were quite similar to R_g of *MtbGmhA* tetramer obtained in small angle X-ray scattering analysis (R_g ~2.38 nm). These data showed that D-sedoheptulose 7-phosphate binding to *MtbGmhA* tetramer enhanced the overall stability of enzyme.

Dynamics simulation on *MtbGmhA* tetramer in complex with D-sedoheptulose 7-phosphate and Zn²⁺. Dynamics simulation on *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn²⁺ tetramer was performed to examine the combined effect of Zn²⁺ and D-sedoheptulose 7-phosphate binding to *MtbGmhA*. Figure 7E showed the superposition of simulated *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn²⁺ (Blue) on starting complex structure (Grey) (RMSD ~0.5 Å for 752 Ca atoms). Overall structure of *MtbGmhA* tetramer was quite stable except minor difference in the loop region of the enzyme. The D-sedoheptulose 7-phosphate and Zn²⁺ ions were quite stable in four subunits of *MtbGmhA* tetramer, except minor movement of Zn²⁺ ions occurred in different subunits. It appears that Zn²⁺ binding further enhanced the stability of *MtbGmhA* + D-sedoheptulose 7-phosphate tetramer and essential for D-sedoheptulose 7-phosphate binding by *MtbGmhA* enzyme.

We have computed the B-factor plot for *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn²⁺ tetramer during the entire simulation period (Fig. 7F). The amino acid stretches, 20–30, 70–80, 120–130 and 150–160 showed the

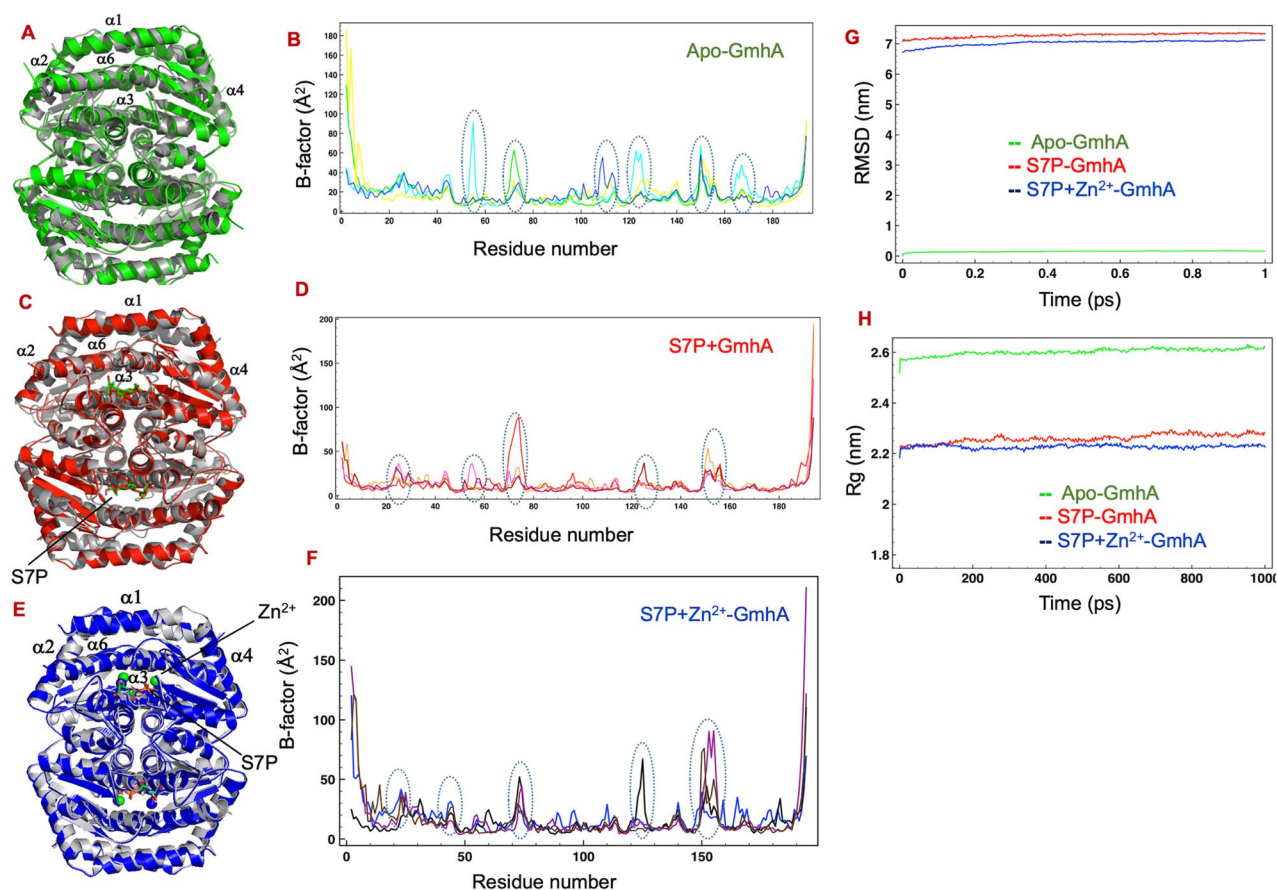


Figure 7. (A) Dynamic simulation on apo *MtbGmhA* tetramer. The simulated *MtbGmhA* tetramer (Green) is superposed on starting structure (Grey). (B) Plot showing the B-factor (Y-axis) and residue number (X-axis) of simulated *MtbGmhA* tetramer. (C) Dynamic simulation on *MtbGmhA* + D-sedoheptulose 7-phosphate tetramer. The simulated *MtbGmhA* + D-sedoheptulose 7-phosphate tetramer (red) is superposed on starting structure (grey). (D) The B-factor (Y-axis) and residue number (X-axis) of simulated *MtbGmhA* + D-sedoheptulose 7-phosphate tetramer. (E) Dynamic simulation on *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn^{2+} tetramer. The simulated complex (blue) is superposed on starting structure (grey). (F) The B-factor (Y-axis) and residue number (X-axis) of simulated *MtbGmhA* + D-sedoheptulose 7-phosphate + Zn^{2+} tetramer. (G) The plot showing the backbone RMSD ~ time for apo, D-sedoheptulose 7-phosphate bound and D-sedoheptulose 7-phosphate + Zn^{2+} bound *MtbGmhA* tetramer. (H) The plot showing the radius of gyration (R_g) ~ time, which showed the changes in degree of compactness of *MtbGmhA* tetramer after D-sedoheptulose 7-phosphate and Zn^{2+} binding.

higher B-factor ($\sim 40\text{--}60 \text{ \AA}^2$) (Fig. 7D). The RMSD increased slightly till 20 ps and remained stable throughout 1 ns simulation ($\sim 0.2 \text{ \AA}$) (Fig. 7G). The radius of gyration (R_g) was quite stable and found $\sim 2.2 \text{ nm}$ (Fig. 7H).

Conclusion

In current study, we have performed the structural and biochemical analysis of *MtbGmhA* enzyme involved in D- α -D-heptose biosynthetic pathway, critical for the development of novel antibiotics against *M. tuberculosis*. The *MtbGmhA* forms a tetramer in solution and small angle X-ray scattering analysis also yielded the tetrameric envelope of the enzyme. The *MtbGmhA* catalyzes the D-sedoheptulose 7-phosphate isomerization with $1.45 \text{ mM}^{-1} \text{ s}^{-1}$ rate and binds the *MtbHddA* and *MtbGmhB* enzymes in μM range. Site directed mutagenesis have identified the roles of various active site residues involved in D-sedoheptulose 7-phosphate binding and catalysis.

The circular dichroism analysis on wild type and six *MtbGmhA* mutants showed quite similar secondary structures and high thermostability of the enzyme. The small angle X-ray scattering analysis have revealed de novo SAXS envelope, which fitted well with modeled *MtbGmhA* tetramer in closed conformation. Dynamics simulation on Apo, D-sedoheptulose 7-phosphate bound and D-sedoheptulose 7-phosphate + Zn^{2+} bound *MtbGmhA* tetramer showed that ligand binding enhanced the overall stability of the enzyme. In *MtbGmhA* tetramer, asymmetric domain movement occurred, compared to isolated monomer. The D-sedoheptulose 7-phosphate binding restricts the domain movement and kept the enzyme in the active conformation. The small angle X-ray scattering analysis indicated that *MtbGmhA* adopts a compact globular conformation, usually observed in crystal structure. This compact conformation of *MtbGmhA* is catalytically relevant, not only for isomerization of D-sedoheptulose 7-phosphate, but also in binding to other enzymes involved in D- α -D-heptose biosynthesis pathway. Our structural and biochemical analysis on *MtbGmhA* have provided a new insight into mechanism,

which will be critical for novel antibiotics development against *M. tuberculosis*. As *MtbGmhA* is required to maintain the permeability in *M. tuberculosis*, current knowledge can be implied in other Gram-positive organisms.

Materials and methods

Expression and purification. The *MtbGmhA* gene (*Rv0113*) was amplified from *H37Rv* genomic DNA by polymerase chain reaction and cloned into *pET28a* expression vector (Novagen) using *NdeI* and *HindIII* restriction sites. The *pET28a-MtbGmhA* plasmid was confirmed by restriction-digestion and gene sequencing analysis. The *pET28a-MtbGmhA* plasmid was transformed in *E. coli* BL21(DE3) cells and cells were grown at 37 °C in luria bertani media (50 µg/ml kanamycin as antibiotic) till OD₆₀₀ ~ 0.6–0.7. The cell culture was induced with 0.5 mM isopropyl β-D-1-thiogalactopyranoside and grown for another 4 h. The *MtbGmhA* protein over-expressed in soluble fraction of the cell.

The cells were harvested by centrifugation at 10,000 × g for 10 min at 4 °C and suspended in lysis buffer containing (25 mM Tris/HCl pH 8.0, 500 mM NaCl, 10% (v/v) Glycerol, 3 mM β-mercaptoethanol, 1 mM Phenylmethylsulfonyl fluoride, 1 mM Benzamidine hydrochloride, 10 mM Imidazole and 0.2 mg/ml Lysozyme). The cells were homogenized, disrupted by sonication and lysate was centrifuged at 25,000 × g to collect the supernatant. The supernatant was loaded on Ni-NTA column (*GE Healthcare*), pre-equilibrated with buffer-A (25 mM Tris/HCl pH 8.0, 300 mM NaCl, 10% Glycerol, 3 mM β-mercaptoethanol, 1 mM Phenylmethyl sulfonyl fluoride and 1 mM Benzamidine hydrochloride). The column was washed with 0–25 mM gradient of imidazole and eluted the protein in buffer-A + 250 mM imidazole. The eluted protein was concentrated and loaded on Superdex 200 column (*GE healthcare Ltd*) pre-equilibrated with buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl, 10% Glycerol and 3 mM β-mercaptoethanol). The peak fractions were pooled and concentrated using 10 kDa cutoff ultracentrifugal device (*Millipore, USA*). The purified *MtbGmhA* was analyzed on 12% SDS-PAGE and mass spectrometry. Protein concentration was determined using absorbance at 280 nm and stored at –80 °C. The recombinant *MtbGmhA* contains 216 residues e.g. 6 residues from 6xHis tag and 14 residues from vector at N-terminal and 196 residues of *MtbGmhA* enzyme.

QuickChange II XL site directed mutagenesis protocol (*Stratagene*) was used to generate the six mutants of *MtbGmhA*. All mutants were purified with protocol used for wild type *MtbGmhA* purification²¹. All *MtbGmhA* mutants were confirmed by gene sequencing.

Small-angle X-ray scattering analysis. The small angle X-ray scattering data was collected by using SAXSpace instrument (Anton Paar, IMTECH, India) at 10 °C using line collimation optics. The *MtbGmhA* enzyme (conc. ~ 1.2 mg/ml in buffer ~ 25 mM Tris-HCl pH 8.0, 150 mM NaCl and 3 mM β-mercaptoethanol) was used for small angle X-ray scattering data collection. The small angle X-ray scattering data on Lysozyme (conc. ~ 5.0 and 3.4 mg/ml in buffer ~ 40 mM sodium acetate buffer pH 3.8, 150 mM NaCl) was collected for beam intensity estimation and comparison with *MtbGmhA* data. After data collection, the *MtbGmhA* enzyme was checked on SDS-PAGE analysis to see if any degradation occurred during data collection. SAXStreat software was used for beam position correction. 120 µl of *MtbGmhA* was loaded in quartz capillary flow cell and exposed to X-ray beam ($\lambda = 1.5418 \text{ \AA}$).

The small angle X-ray scattering data was collected in triplicate and averaged. SAXSquant 4.2.4 software was used to remove the buffer contribution and desmearing the line collimation. The small angle X-ray scattering data were processed using ATSAS 2.8.3 suite and ScÅtter²² and data containing scattering intensity (*I*) as a function of scattering vector *Q*, [where $Q = (4\pi/\lambda) \sin\theta$, θ is the scattering angle and λ is wavelength] were obtained. The Kratky plot [$I(Q)Q^2$ vs *Q*] was obtained from intensity profile. The radius of gyration (R_g) and radius of cross section (R_c) were estimated from scattering intensity at low *Q* region using Guinier analysis. Additionally, using R_g and R_c values, the length of an ellipsoidal structure *L* were estimated using the $L = [12(R_g^2 - R_c^2)]^{1/2}$ relationship. The GNOM45 program was used to estimate the shape and size of scattering entities, which consider both low and high *Q* data. Indirect Fourier transformation of scattering data over measured *Q* range was computed as a pairwise distribution function of interatomic vectors²³. This analysis also provided R_g and I_0 from the second moment and start of *P*(*R*) as well as D_{max} (maximum diameter).

Ten independent models of *MtbGmhA* were generated using DAMMIF program v. 1.1.2²⁴, averaged the aligned model and filtered at a given cutoff volume using DAMAVER v. 5.0 program²⁵. The *MtbGmhA* tetramer was superimposed on SAXS envelope using the SUPCOMB v. 2.3 program²⁶. The SAXS curve was calculated from the atomic model using the CRY SOL v. 2.8.3 program²⁷. The PyMOL v. 2.3 program²⁸ was used for structural visualization and to generate all superimposed structures. The curve fitting and data plotting were performed using Graphpad-Prism 5 program²⁹.

Enzyme assay. The activity of wild type and six mutants of *MtbGmhA* were determined by coupling reaction using *MtbHddA*, *MtbHddA* and *MtbGmhB* enzymes and release of P_i was monitored (as described in DeLeon *et. al.*³⁰). We prepared the 200 µl of reaction mixture containing (0.31 nM *MtbGmhA*, 0.11 nM *MtbHddA* and 0.4 nM of *MtbGmhB*) in 20 mM HEPES buffer pH 8.0, 10 mM MgCl₂, 10 mM KCl, 10 mM ATP and 0–3 mM D-sedoheptulose 7-phosphate were added in the reaction mixture and incubated at room temperature for 30 min. The P_i concentration was measured using 50 µl of P_i colorlock Gold and 0.5 µl of Accelerator (*Innova Biosciences Ltd*) in the reaction mixture. After 2 min, 20 µl of stabilization reagent was added in each sample and absorbance was taken at 630 nm. All reactions were performed in triplicate and 11 different concentrations of D-sedoheptulose 7-phosphate were used for each reaction. The kinetic parameters were determined by fitting the data to Michaelis–Menten equation using GraphPad Prism version 6.0.2 (GraphPad Software, La Jolla).

$$v = \frac{k_{cat} * E_t [S]}{(K_m + [S])}$$

Binding analysis. The D-sedoheptulose 7-phosphate binding to *MtbGmhA* was performed using Auto-lab Esprit Surface Plasmon Resonance equipment³¹. The gold surface [self-assembled monolayer of 11-mercaptoundecanoic acid (11-MUA)] was first activated by EDC (N-ethyl-N dimethyl amino propyl carbodiimide, conc. ~0.2 M)—NHS (N-hydroxysuccinimide, conc~0.05 M) coupling. 50 μ M of *MtbGmhA* was immobilized on gold surface using 20 mM sodium acetate buffer, pH 4.2. After immobilization, the surface was blocked with 100 mM ethanolamine pH 8.5. The binding experiment was performed at 25 °C in running buffer (20 mM HEPES pH 7.5, 150 mM NaCl, 5 mM MgCl₂, and 2% Glycerol). The D-sedoheptulose 7-phosphate substrate was injected at 30 μ l/min for 5 min, followed by dissociation for 5 min. After each experiment, the sensor surface was regenerated using 50 mM NaOH.

For interaction analysis between *MtbGmhA*, *MtbHddA* and *MtbGmhB*, 50 μ M of *MtbGmhB* was immobilized on gold surface activated by EDC-NHS coupling using 20 mM sodium acetate buffer, pH 4.2. The surface was blocked with 100 mM Ethanolamine pH 8.5. 50 μ M of *MtbGmhA* was injected at 30 μ l/min for 5 min followed by dissociation for 5 min. Similar protocol was used for interaction analysis between *MtbHddA* and *MtbGmhB* and *MtbGmhA* and *MtbHddA* enzymes. All experiments were performed at 25 °C. The sensograms were collected, processed, analyzed using BIAevaluation (GE Healthcare)³² and fitted with Langmuir 1:1 model using formula

$$R_{eq} = \frac{R_{max} * K_A * C_A}{K_A * C_A + 1}$$

where R_{eq} is binding response at steady state, R_{max} is maximal binding response, C_A is the concentration of analyte and K_A is equilibrium association constant.

Circular dichroism analysis. Circular dichroism analysis was performed to estimate the secondary structure of *MtbGmhA* enzyme³³. A far UV spectrum on *MtbGmhA* was collected using Chirascan spectropolarimeter (Applied Photophysics) with quartz cuvette of 1 mm path length. The *MtbGmhA* enzyme was transferred in Sodium phosphate buffer, pH 7.5. The circular dichroism data was collected in 200–260 nm range at 25 °C. The spectra from baseline was subtracted from protein spectra and obtained values were averaged for each dataset. The mean residue ellipticity (θ) was calculated from the observed spectra, as a function of wavelength. The K2D2 program was used to estimate the secondary structure of *MtbGmhA*³⁴. Similar protocol was used for CD data collection on all six *MtbGmhA* mutants. The theoretical secondary structure prediction on *MtbGmhA* was performed using various programs e.g., SOPMA³⁵, CFSP36, GOR³⁷, PHD³⁸, SIMPA96³⁹, DSC⁴⁰, HNN⁴¹, RaptorX⁴², JPred⁴³ and PsiPred⁴⁴.

For thermal stability analysis, the circular dichroism spectra of *MtbGmhA* was recorded at 222 nm from 25 to 90 °C range in 10 °C step. The melting temperature was calculated from scattering profile using polynomial fitting, taking temperature corresponding to half denaturation⁴⁵. A graph of ($\Delta A/\Delta T$) versus (T_{avg}) was plotted using Sigma Plot program. A bell-shaped melting curve was observed, and peak corresponds to melting temperature of the protein.

Multiple sequence alignment. The *MtbGmhA* sequence was retrieved from UniProt (P9WGG1) database and sequences of 11 GmhA orthologues were retrieved from RCSB database⁴⁶. The *MtbGmhA* sequence was aligned with 11 GmhA orthologues using MultAlin⁴⁷ and ESPript 3.0⁴⁸ programs.

Molecular modeling, docking and dynamics simulation. I-TASSER server was used to model the *MtbGmhA* tetramer and was ranked by the C-score and TM-score⁴⁹. The *MtbGmhA* sequence (1–196 residues) was given as input and server retrieved the template proteins of similar folds from Protein Data Bank by LOMETS (Local Meta-Threading-Server)⁵⁰. Additional steps were performed to remove the steric clashes and refined the global topology of the *MtbGmhA* models.

I-TASSER server identified the PDB-2X3Y (Crystal structure of GmhA tetramer in closed state from *B. pseudomallei*¹²) as the best template for *MtbGmhA* modeling (RMSD = 0.79, id1 = 0.39, id2 = 0.39, Conv = 0.98, Z-score = 3.2). The best *MtbGmhA* model was obtained having following parameters (C-score = 0.21 and TM-score = 0.88 \pm 0.07), which indicated a reliable model with correct global topology. The I-TASSER server also yielded PDB-2I2W (Crystal structure of *Escherichia Coli* Phosphoheptose Isomerase in open state¹⁰) as another template for *MtbGmhA* modeling, however following parameters (RMSD = 2.1, TM id1 = 0.35, TM-score = 0.82, Conv = 0.89) did not warrant a reliable *MtbGmhA* model in “open state”.

Docking of D-sedoheptulose 7-phosphate and Zn²⁺ into *MtbGmhA* tetramer was performed using GLIDE module of the Schrodinger-9.4 program⁵¹. The *MtbGmhA* + S7P + Zn²⁺ complex was generated using induced fit protocol of Schrödinger-9.4 program. Defaults parameters were used except Extra Precision (EP) scoring function was used in docking calculation.

The dynamics simulation of (i) Apo (ii) D-sedoheptulose 7-phosphate bound and (iii) D-sedoheptulose 7-phosphate + Zn²⁺ bound models of *MtbGmhA* tetramer were performed using GROMACS package 4.5 (Groningen MACHine for Chemical Simulations) using CHARMM all-atom force field parameters (Table 4)⁵². The *MtbGmhA* tetramer was solvated in the dodecahedron box at 1 nm distance from the boundary. Simple point charge water molecules were used for solvation together with Na⁺ and Cl⁻ ions to neutralize the total charge of the system. Energy minimization was performed using 50,000 steps of steepest descent minimization. To

maintain the 300 K constant temperature, protein and non-protein atoms were coupled to their temperature baths using V-scale thermostat using 100 ps NVT. The whole system was kept for 100 ps NPT equilibration and 1 bar pressure. For all simulations, the bond lengths and water molecules were restrained using the SETTLE and LINCS⁵³ algorithms respectively.

The 1 nm cutoff was used for the treatment of Van der Waals interactions and long-range electrostatic interactions and simulated using PME⁵⁴ method with 0.16 FF grid spacing and 4th order B-spline interpolation for the reciprocal sum space. Periodic boundary conditions were applied in all directions. Rigid body displacements and rotations were removed from all trajectories. The PRODRG server⁵⁵ was used to generate the topology of the D-sedoheptulose 7-phosphate. PyMol and Plot2⁵⁶ programs were used for generating the figures and all simulation trajectories. The *Mtb*GmhA tetramers obtained after dynamics simulation exhibited good stereochemistry. The structural validation on all models was performed using various servers e.g., ERRAT⁵⁷, Verify3D⁵⁸, and PROCHECK⁵⁹.

Received: 26 February 2019; Accepted: 15 October 2020

Published online: 30 November 2020

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Acknowledgements

Current project was supported by research grants from UGC-SAP, UGC-Networking and DST-PURSE from Jawaharlal Nehru University, New Delhi. Sumita Karan and Bhanu Pratap were supported by Senior Research Fellowship from UGC, India. The authors thank the staff members of Advanced Instrumentation Research Facility (AIRF) of Jawaharlal Nehru University for their help in CD experiments. The authors thank the staff at small angle X-ray diffraction facility at IMTECH, Chandigarh for their help in SAXS experiment.

Author contributions

S.K. designed the primers, cloned the genes, performed protein expression and purification of wild type and all mutants of *MtbGmhA* proteins. S.K. prepared the protein samples and performed functional and interaction analysis using surface plasmon resonance experiments. S.K. and B.P. prepared samples and analyzed CD and thermal denaturation data. S.K., S.P.Y. and FNU Ashish. collected the small angle X-ray scattering data, processed and fitted the structures. A.K.S. conceived, designed the strategies and techniques employed, supervised the research, and performed entire molecular modeling and dynamic simulation analysis. S.K. and A.K.S. wrote the entire paper.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at <https://doi.org/10.1038/s41598-020-77230-8>.

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