Crystal structure of peptidyl-tRNA hydrolase from a Gram-positive bacterium, *Streptococcus pyogenes* at 2.19 Å resolution shows the closed structure of the substrate-binding cleft

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**ABSTRACT**

Peptidyl-tRNA hydrolase (Pth) catalyses the release of tRNA and peptide components from peptidyl-tRNA molecules. Pth from a Gram-positive bacterium *Streptococcus pyogenes* (SpPth) was cloned, expressed, purified and crystallised. Three-dimensional structure of SpPth was determined by X-ray crystallography at 2.19 Å resolution. Structure determination showed that the asymmetric unit of the unit cell contained two crystallographically independent molecules, designated A and B. The superimposition of Cα traces of molecules A and B showed an r.m.s. shift of 0.4 Å, indicating that the structures of two crystallographically independent molecules were identical. The polypeptide chain of SpPth adopted an overall α/β conformation. The substrate-binding cleft in SpPth is formed with three loops: the gate loop, Ile91–Leu102; the base loop, Gly108–Gly115; and the lid loop, Gly136–Gly150. Unlike in the structures of Pth from Gram-negative bacteria, the entry to the cleft in the structure of SpPth appeared to be virtually closed. However, the conformations of the active site residues were found to be similar.

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1. Introduction

The translation of genetic information by mRNA to ribosome often gets terminated due to a variety of reasons [1–4]. The premature termination results in the release of peptidyl-tRNA molecules which are toxic to the cell [5–8]. The loss of tRNA molecules to such a by-product reduces the availability of free tRNAs. This affects the protein synthesis adversely. However, such a condition can be corrected by an enzyme known as peptidyl-tRNA hydrolase (Pth). It has been shown by mutation studies that this enzyme is essential for the survival of bacteria [9]. Pth is an esterase which can be corrected by an enzyme known as peptidyl-tRNA hydrolase (Pth). It has been shown by mutation studies that this enzyme is essential for the survival of bacteria [9]. Pth1 is an esterase which can be corrected by an enzyme known as peptidyl-tRNA hydrolase (Pth). It has been shown by mutation studies that this enzyme is essential for the survival of bacteria [9], and human Pth2 enzyme has 116 amino acid residues [23]. Structurally also, Pth1 is significantly different from archaeal Pth2 [21,22] and human Pth2 enzymes [24,25]. Pth1 is found in the cells of bacteria and eukaryotes [15–17] while Pth2 is found in archaea and eukaryotes [18–20]. In eukaryotes, both Pth1 and Pth2 as well as some additional types of Pth enzymes are present [14,18,19]. While Pth1 is essential in bacteria for restoring the translational function of ribosome [9,17], it is not absolutely essential in eukaryotes [17–19]. Pth1 consists of approximately 190 amino acid residues while archaeal Pth2 has approximately 120 amino acid residues [21,22] and human Pth2 enzyme has 116 amino acid residues [23]. Structurally also, Pth1 is significantly different from archaeal Pth2 [21,22] and human Pth2 enzymes [23]. Since the present paper deals with the investigations on bacterial Pth1 enzyme only, it will be referred hereafter as Pth.

So far, crystal structures of Pth enzymes are available from a few Gram-negative bacteria including *E. coli* (EcPth) (PDB: 2PTH) (PDB: 2PTH, 3VJR) [24], *Acinetobacter baumannii* (AbPth) (PDB: 3V9H, 4JKW, 4JX9) [25], *Pseudomonas aeruginosa* (PaPth) (PDB: 4J4C, 4FYJ) [26,27], *Francisella tularensis* (FrPth) (PDB: 3NEA) [28] and *Burkholderia thailandensis* (BtPth) (PDB: 3V21) [29]. In addition to these structures, crystal structures of Pth enzymes from *Mycobacterium tuberculosis* (MtPth) (PDB: 2Z2I) [30] and *Mycobacterium smegmatis* (MsPth) (PDB: 3KZJ) [31] are also known. However, so far, no structure of Pth from any Gram-positive bacteria is known. We report here, the first crystal structure of Pth from a Gram-positive...
bacterium, *S. pyogenes* (SpPth). SpPth shows sequence identities ranging from 30% to 32% (Fig. 1) with Pth enzymes of other bacteria whose structures are known [24–31]. The three-dimensional structure of SpPth has been determined at 2.19 Å resolution.

2. Results

2.1. Overall structure

The structure determination of SpPth revealed the presence of two crystallographically independent molecules, A and B in the asymmetric unit. Each molecule consists of 189 amino acid residues. Molecules A and B were arranged side-by-side with a buried surface area of 535 Å². They formed three intermolecular hydrogen bonds and several van der Waals contacts. However, the gel-filtration profile of SpPth (Fig. S1A), when compared with a calibration curve obtained using proteins of known molecular weights (Fig. S1B), indicated a molecular weight of about 24 kDa. It suggested that SpPth did not form a stable dimer in solution. Thus, the association of two crystallographically independent molecules in the crystals may not be considered as a dimer.

The r.m.s. shift of 0.4 Å was obtained when Cα traces of the two crystallographically independent molecules, A and B were superimposed on each other. It showed that the structures of molecules, A and B were identical. Therefore, hereafter only one molecule will be used in the subsequent discussion. The structure of SpPth consisted of six α-helices and seven β-stands (Fig. 2A). Four parallel β-strands, β4, β1, β5 and β7 formed a twisted β-sheet in the centre of the molecule. The antiparallel β-strands, β2 and β3 cross the sheet at the centre. The β-strand β6 is antiparallel to β-strand, β7 and parallel to the longest α-helix, α5 in the structure. The β-structure is surrounded from three sides by α-helices among...
which two \( \alpha \)-helices, \( \alpha 2 \) and \( \alpha 3 \) provided a cover from one side while \( \alpha 5 \) and \( \alpha 6 \) are located on the opposite side of \( \alpha 2 \) and \( \alpha 3 \). The \( \alpha \)-helix, \( \alpha 1 \) covers the \( \beta \)-sheet structure from the third side. The fourth side is open and may be required to allow the hinge motion involving the \( \beta \)-sheet. The residues of N-terminus are predominantly hydrophobic in nature while those at the C-terminus are hydrophilic. It may be mentioned here that the structures of N- and C-terminal segments were clearly defined in the electron densities.

The structure of AbPth (PDB: 3WH4) was used for comparison as it represented a native unbound state of Pth from Gram-negative bacteria and also has the maximum sequence identity of 32% with SpPth. The superimposition of \( C^\alpha \) traces of SpPth on those of AbPth gave an r.m.s. shift of 1.6 Å indicating an appreciable deviation between the paths of polypeptide chains of two Pth enzymes representing two different classes of bacteria (Fig. 2B). In contrast, the corresponding r.m.s. shifts among \( C^\alpha \) traces of Pth enzymes belonging to Gram-negative bacteria [24–29] were found to be in the range of 0.7–1.4 Å. It showed that the path of the polypeptide chain of a Pth enzyme from a Gram-positive bacterium differed slightly from those of Gram-negative bacteria.

The values of length and width of the molecular structure of SpPth as estimated in terms of distance between His55 \( C^\alpha \) on one end and Gly141 \( C^\alpha \) on the opposite end and the distance between Lys42 \( C^\alpha \) belonging to one side and Gln185 \( C^\alpha \) belonging to the opposite side were 49.6 Å and 36.2 Å, respectively. The corresponding values in the structure of AbPth were found to be 44.7 Å and 35.8 Å, respectively. However, it may be mentioned here that both regions to which His55 and Lys42 belonged have higher values of B factors as compared to the rest of the protein. This may be due to conformational flexibility.

### Substrate binding cleft

The substrate binding cleft in Pth enzymes is formed with three segments, a base loop, Gly108–Gly115, a gate loop, Ile91–Leu102 and a lid loop, Gly136–Gly150 (Fig. 3A1). The base loop forms one side of the cleft while lid loop and gate loop form the opposite side of the cleft. The minimum distance from any atom of the base loop, Gly108–Gly115 to any atom of lid loop, Gly136–Gly150 in the native unbound state of SpPth was found to be 3.47 Å. This showed that the structure of SpPth in the native unbound state had the narrowest opening between the base and the lid loops among all the known structures of Pth enzyme reported so far (Table 2A). Similarly, the shortest distance between any atom of base loop, Gly108–Gly115 and any atom of gate loop, Ile91–Leu102 was found to be 3.88 Å. The width of the gate in SpPth was found to be smaller than those observed in Pth enzymes of Gram-negative bacteria (Table 2B). Only in the case of MtPth, the corresponding distance of 3.33 Å was smaller than that of SpPth (Table 2B).

Furthermore, in the structure of SpPth, the residues from opposite sides of the cleft formed several van der Waals contacts (Fig. 3A2). These distances indicated that the substrate binding cleft was closed at both lid and gate loops as well as the region between the closed lid loop and the closed gate loop (Fig. 3B1). In MtPth, it is fully closed at the gate loop but it is wide open at the lid loop (Fig. 3B2). On the other hand, the substrate binding channel in EcPth is wide open as both lid and gate loops are away from the base loop (Fig. 3B3). It shows that the catalytic site in the structure of SpPth appeared to be inaccessible. For the catalytic action to occur, the lid loop, Gly136–Gly150 and the gate loop Ile91–Leu102 will have to move away from their respective positions in the native state for the substrate to reach the active site residues. As seen from the superimposition of substrate binding clefts of SpPth (green) and EcPth (grey) (Fig. 4) together with the bound C-terminal tripeptide from the neighbouring molecule in EcPth, there were steric constraints for the binding of such a peptide in the native state of SpPth. This further indicated that the binding of ligands in the substrate binding cleft in SpPth may occur only if the lid loop, Gly136–Gly150 and the gate loop, Ile91–Leu102 moved away considerably.

### Catalytic site

The structural and point mutation studies have suggested that the amino acid residues, His20, Asp93, Asn10, Asn68 and Asn114 may play a role in the catalytic action of Pth enzymes [24,32–34]. The structural studies of the complexes of AbPth and PaPth with ligands have also shown that residues, Asn10, His20, Asn68 and Asn114 are involved in the binding with ligands in the

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**Fig. 2.** (A) Overall structure of SpPth in ribbon representation. Secondary structure elements, \( \alpha \)-helices, 1–6 and \( \beta \)-strands, 1–7 are labelled. (B) Superimpositions of \( C^\alpha \) traces of SpPth (green) and AbPth (grey) showing variations in the paths of their polypeptide chains. The r.m.s. shift in the positions of \( C^\alpha \) atoms of the two structures was found to be 1.6 Å. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
substrate binding cleft [25,27]. However, Out of these residues, the conformation of the side chain of Asn114 was found to be very sensitive when Pth interacted with other molecules through the substrate binding cleft. The torsion angle, $\chi_1$ of $63^\circ$ for the side chain of Asn114 as observed in the structure of SpPth defined the conformation of Pth in the native unbound state. The value of the corresponding torsion angle in the native unbound state of AbPth [PDB: 3WH4] was $69^\circ$. Also, in the structure of the complex of EcPth with tRNA CCA-acceptor-TΨC domain [34] in which there are no interactions with the active site region, the value of corresponding torsion angle, $\chi_1$ was $60^\circ$. Similarly, the values of $63^\circ$ and $97^\circ$ were observed for the corresponding torsion angles in the native structures of MtPth [30] and MsPth [31], respectively. These values clearly indicated that the torsion angle, $\chi_1$ for the side chain of Asn114 in the native unbound state of Pth was centred at $60^\circ$ [24,25,27]. Based on these observations, it may be concluded that the conformation of the side chain of Asn114 provided a clear indication about the binding state at the substrate binding site in Pth.

3. Discussion

The structure determination of SpPth revealed that the gate loop, Ile91–Leu102 and the lid loop, Gly136–Gly150 formed one side of the cleft while the base loop, Gly108–Gly115 represented the opposite wall. It may be mentioned here that all the three loops in the structure of SpPth had well defined structures with values of 16 Å$^2$ for the average B factors. The lid loop, Gly136–Gly150 in SpPth is internally well stabilized with a number of intra-loop interactions (Fig. 5A). The segment, Val144–Met149 of this loop was found to adopt a stable $\alpha$-helical structure which was stabilized by hydrogen bonds, Val144 O$^\cdot\cdot\cdot$N Val148 = 2.95 Å and Ile145 O$^\cdot\cdot\cdot$N Met149 = 2.85 Å. A tetra peptide of this loop, Leu139–Lys140–Gly141–Met142 formed a tight type II $\beta$-turn structure with a hydrogen bond between Leu139 O$^\cdot\cdot\cdot$N Met142 = 3.04 Å. Additionally, the side chain of Pro138 was stacked with the aromatic ring of His147. Similarly, the gate loop, Ile91–Leu102 (Fig. 5B) was also well stabilized in the structure of SpPth. In this loop, a tetra peptide, Asp94–Leu95–Asp96–Met97 adopted a type I $\beta$-turn conformation which was stabilized by a hydrogen bond between Asp94 O$^\cdot\cdot\cdot$O and Met97 N. An intra-residue hydrogen bond was also observed between Asp96 O$^\cdot\cdot\cdot$O and Asp96
N. Similar tight β-turn was not observed in the structures of other Pth enzymes [24–31] due to specific amino acid sequence differences in the corresponding loops. Furthermore, due to these tight β-turns in the loops, Ile91–Leu102 and Gly136–Gly150, the paths of the polypeptide chain after the β-turns in SpPth differed significantly from that reported in the native unbound structure of AbPth (Fig. 6A). Also such a conformation was found to influence the positions of the side chains of Leu95 and Val144 which shifted towards the base loop, Gly108–Gly115. It reduced the free space in the substrate binding cleft. The comparison of distances between Val144 of lid loop and Gly112 of base loop in various Pth enzymes are given in Table 2A. These distances indicated that the positions of the lid loops in various Pth enzymes varied considerably. The observed van der Waals contacts involving residues, Val144 and Leu95 with the atoms of the base loop, Gly108–Gly115 from the opposite side of the cleft (Fig. 3A2) and showed that the substrate binding cleft in SpPth was packed somewhat differently.

The substrate binding cleft in SpPth is expected to open on ligand binding where both the gate loop, Ile95–Leu102 and the lid loop, Gly136–Gly150 will be required to move away from their closed positions so that the substrate could align with the catalytic residues. Due to tight internal packings of these two loops in the structure of SpPth, the lid loop may have slightly different movement from those observed in other Pth enzymes where the corresponding loops are relatively more flexible [24–31].

Previously, in the structures of Pth enzymes of various bacteria, the distances between Gly113 Cα and Asp98 Oδ2 were described to determine the widths of the gate [25,26,30,31]. In the present structure, the side chain of corresponding residue Asp96 occupied a different position because Asp96 Oδ3 formed a hydrogen bond with Arg137 Nε (Fig. 6B). As a result of this, the distance estimated for the width of the cleft with respect to Asp96 Oδ in SpPth was not the shortest distance. In this case, the shortest distance was provided by Leu95 Cδ3. The corresponding interaction involving Asp96 is not present in Pth enzymes of Gram-negative bacteria because those enzyme consisted of His138 instead of Arg137 [24–29]. Although in MtPth and MsPth, the corresponding residue is Arg139, it still occupied a different position in the structure because it followed a slightly different chain path due to a unique sequence of Pro139–Pro140 [30,31]. The corresponding sequence in SpPth is Pro138–Leu139 while in Pth of Gram negative bacteria, it is Pro139–Gly140 [24–29]. Due to such variations in the nature of interactions involving Asp96, the side chain of Asp96 may occupy different positions. Thus, using Asp96, as used previously in the structures of Pth enzymes [25,26,30,31] as a reference residue for the determination of the width of the gate to the substrate binding site did not provide an accurate generalisation. The actual estimation of the width of the gate should be based on the criterion of the shortest distance between any atom of the base loop, Gly108–Gly115 and any atom of the gate loop, Ile91–Leu102. The comparison of these distances in Pth structures are given in Table 2B. It is interesting to note that the shortest distance in SpPth is between Gly112 and Leu95 while in rest of the structures, the shortest distances are estimated between Gly111/Gly113 and Asp96/Asp98.

It may also be mentioned here that the structure of SpPth is the first structure of Pth enzyme from a Gram-positive bacterium. It
The forward Fw genomic DNA was isolated and the Pth gene was amplified using bial Type Culture Collection (MTCC) Number 1924. From this, the Institute of Microbial Technology, Chandigarh (India) with Micro-(Luria–Bertani) containing 100
into pGEMT-easy cloning vector and sub-cloned into pET-28a restriction sites, respectively. The amplified gene was cloned Xho1
loop. The overall variations in the structures of the substrate bind-
shown. Asp96 in SpPth forms an ionic interaction with Arg137 while Asp98 of AbPth is not involved in such an interaction. As a result, Asp96 in SpPth, has moved away from Gly112 of the base loop Gly108–Gly115. The residue corresponding to Arg137 of SpPth is His138 in AbPth and other enzymes from Gram-negative bacteria. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

revealed notable differences in the structural features of the lid and gate loops when compared with those of other Pth enzymes. The region between the lid and gate loops was found to be tightly packed. Also the structure of SpPth provided a new rationalisation for the measurements of the width of the gate. The structure determination also revealed that the substrate binding cleft is obstructed at two positions, one by the lid loop and by the gate loop. The overall variations in the structures of the substrate bind-
ing clefts of Pth enzymes from various bacteria may be helpful in the designing of specific inhibitors against the enzymes of different bacteria.

4. Materials and methods

4.1. Cloning, expression and purification of SpPth

The freeze dried culture of S. pyogenes was obtained from CSIR-Institute of Microbial Technology, Chandigarh (India) with Micro-
bial Type Culture Collection (MTCC) Number 1924. From this, the genomic DNA was isolated and the Pth gene was amplified using the forward Fw 5′-CGGGATCATGATGAAAAATGATTGTTGGTC-3′ and reverse Rw 5′-CCGCTGGATATCCATTAAATTTCTGCATTG-3′ primers. The forward and reverse primers contained BamH1 and XhoI restriction sites, respectively. The amplified gene was cloned into pGEMT-easy cloning vector and sub-cloned into pET-28a expression vector. The sequence of the constructed plasmid was verified by DNA sequencing. The verified plasmid was introduced into E. coli BL21 (DE3) expression system.

A single freshly transformed colony was inoculated in 10 ml LB (Luria–Bertani) containing 100 μg/ml kanamycin and was kept in water bath overnight at 37 °C in shaking condition. 1% culture from primary inoculation was added in 1000 ml LB medium containing 100 μg/ml kanamycin and kept in orbital shaker at 37 °C in shaking condition until the optical density (OD) of the culture at 600 nm reached the level of 0.4–0.6. 5 ml of this secondary culture were removed and kept at 4 °C as an uninduced culture. The remaining secondary culture of recombinant cells containing the insert was induced with 0.4 mM isopropyl-1-thio-β-D-galactopyranoside (IPTG) at 20 °C for 16 h. The cells were harvested by centrifugation at 9803g for 15 min.

The cell pellet was dissolved in 50 mM Tris–HCl buffer containing 300 mM NaCl and 10 mM imidazole, pH 8.0, 1 mM protease inhibitor cocktail (Roche, Basel, Switzerland). The cells were disrupted using sonicator (Model UP50H, Hielscher, Brandenburg, Germany). The cleared lysate was applied to an Ni-NTA Super-flow column (Qiagen, Hilden, Germany) which was pre-equilibrated in lysis buffer containing 50 mM Tris–HCl buffer, 300 mM NaCl and 10 mM imidazole, pH 8.0 and purified using stepwise washing with 30 mM imidazole followed by 300 mM imidazole in lysis bu-
fer. The eluted fractions were examined using 10% sodium dodecyl sulphate–polyacrylamide gel electrophoresis (SDS–PAGE). A single band of protein corresponding to molecular weight of approximately 24 kDa was observed. The final purification step was car-
ried out using gel filtration chromatography with Sephadex G-50 column. The purity of the protein was established using SDS–PAGE which showed a single band at approximate molecular weight of 24 kDa.

4.2. Crystallisation of SpPth

The freshly prepared sample of purified protein was used for crystallization with hanging drop vapour-diffusion method at 298 K using 24 well limbro crystallization plates. The initial crystal-
ization screening experiments were carried out using Hampton Research crystallization kit (HR2110–HR2112). The protein concentration was set to 16 mg/ml in 50 mM HEPES buffer, pH 6.5. The crystals of SpPth were obtained by equilibrating 4 μl protein drops (containing 2 μl protein solution and 2 μl reservoir solution) against a reservoir solution containing 25% (v/v) PEG 10,000 and 50 mM HEPES buffer, pH 6.5. The crystals grew to approximate dimensions of 0.4 × 0.4 × 0.4 mm3 in about 2 weeks.

4.3. X-ray intensity, data collection and processing

Crystals of SpPth were stabilized by adding 25% glycerol to the mother liquor for data collection at low temperature. A single crys-
tal was mounted in a nylon loop and flash-frozen in liquid nitrogen. The data were collected at 100 K on MAR 225 CCD detector (MAR RESEARCH, Norderstedt, Germany) using beamline, BM14 at ESRF, Grenoble, France. A complete data set was collected using

Fig. 6. (A) The superimpositions of the regions consisting of loops, Gly108–Gly115, Ile91–Leu102 and Gly136–Gly150 of SpPth (green) on AbPth (grey). The comparison shows that the residues in SpPth occupy significantly different positions in the cleft and form van der Waals contacts involving inter-loop residues. (B) The interactions of Asp96 in SpPth (green) and of the corresponding Asp98 in AbPth (grey) are shown. Asp96 in SpPth forms an ionic interaction with Arg137 while Asp98 of AbPth is not involved in such an interaction. As a result, Asp96 in SpPth, has moved away from Gly112 of the base loop Gly108–Gly115. The residue corresponding to Arg137 of SpPth is His138 in AbPth and other enzymes from Gram-negative bacteria. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Table 1
Crystallographic data for Peptidyl-tRNA hydrolase from *Streptococcus pyogenes* at 2.19 Å resolution. Values in parentheses are for the highest resolution shell.

| Data collection statistics | P1 |
|---------------------------|----|
| Space group               | P1 |
| Unit-cell dimensions      |    |
| a (Å)                     | 36 |
| b (Å)                     | 43 |
| c (Å)                     | 65.1 |
| α (°)                     | 90.3 |
| β (°)                     | 105.7 |
| γ (°)                     | 112.5 |
| Number of molecules in unit cell | 2 |
| Vm (Å³/Da)                | 2 |
| Resolution range (Å)      | 35.40–2.19 |
| No. of measured reflections| 64,832 |
| No. of unique reflections  | 17,325 |
| Overall completeness (%)  | 98.3 (97.3) |
| R cryst (%)               | 16.8 (19.7) |
| Rfree (5% data) (%)       | 19.8 (21.6) |
| Number of protein atoms   | 2980 |
| Number of water oxygen atoms | 281 |

| r.m.s. deviations         |    |
| Bond length (Å)            | 0.014 |
| Bond angles (°)            | 1.6 |
| Dihedral angles (°)        | 17.6 |

| Mean B factor (Å²) for     |    |
| Main chain atoms           | 15.5 |
| Side chain and water oxygen atoms | 19.8 |
| Overall                    | 17.8 |

| Ramachandran plot statistics |    |
| Residues in the most favoured regions (%) | 93.2 |
| Residues in the additionally allowed regions (%) | 6.2 |
| Residues in the disallowed regions (%) | 0.6 (Phe68 A, Phe68 B) |

An oscillation range of 1° with an exposure time of 3 s per image using a wavelength of 0.98 Å. A total of 400 images were collected. The crystals diffracted to 2.19 Å resolution. The indexing of data indicated that the crystals belonged to space group P1 with unit cell dimensions, a = 36.0 Å, b = 43.0 Å, c = 65.1 Å, α = 90.3°, β = 105.7°, γ = 112.5°. The Matthews constant, Vm [35] of 2.0 Å³/Da was calculated for two molecules in the asymmetric unit which corresponded to a solvent content of 41.3%. The data were processed with AUTOMAR and SCALEPACK from HKL package [36]. The summary of data collection and processing details is presented in Table 1.

### 4.4. Structure determination and refinement of SpPth

The structure of SpPth was determined with program Phaser [37] in CCP4i suite [38] using coordinates of AbPth (PDB: 4WH4) as the search model. The amino acid sequence identity between SpPth and AbPth was 32% (Fig. 1). The Phaser program gave a solution for two molecules in the asymmetric unit. The coordinates obtained from the output of calculations using Phaser program were subjected to several cycles of maximum likelihood refinement with REFMACS [39]. The whole chain was rebuilt, segment by segment, using omit maps which were calculated by removing segments of a protein chain followed by 10 cycles of refinement with REFMAC5. The refined model was adjusted manually using programs, O [40] and COOT [41]. The structure gradually improved on fitting the protein molecule in electron density maps calculated with (Fo–Fc) and (2Fo–Fc) coefficients. This was followed by 10 rounds of refinement cycles. The model was improved by carrying out further manual model building. The additional cycles of refinement were also carried out to locate water oxygen atoms. The water oxygen atoms were placed at the electron density peaks in the (Fo–Fc) map where the peaks were above 3σ and formed hydrogen bonds with protein atoms or with other water molecules. The B-factors were refined isotropically. Finally, the refinement converged to values of 0.168 and 0.198 for R cryst and Rfree factors, respectively. The structural validation was carried out using the program PROCHECK [42]. The final refined model consists of 2980 protein atoms and 281 water oxygen atoms. The data quality and final refinement statistics are reported in Table 1. The coordinates and structure factors have been deposited in the Protein Data Bank with accession code PDB: 4QT4.

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### Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.fob.2014.10.010.

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