A modeled structure for amidase-03 from *Bacillus anthracis*

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**Abstract**

Homology models of amidase-03 from *Bacillus anthracis* were constructed using Modeller (9v2). Modeller constructs protein models using an automated approach for comparative protein structure modeling by the satisfaction of spatial restraints. A template structure of *Listeria monocytogenes* bacteriophage PSA endolysin PlyPSA (PDB ID: 1XOV) was selected from protein databank (PDB) using BLASTp with BLOSUM62 sequence alignment scoring matrix. We generated five models using the Modeller default routine in which initial coordinates are randomized and evaluated by pseudo-energy parameters. The protein models were validated using PROCHECK and energy minimized using the steepest descent method in GROMACS 3.2 (flexible SPC water model in cubic box of size 1 Å instead of rigid SPC model). We used G43a1 force field in GROMACS for energy calculations and the generated structure was subsequently analyzed using the VMD software for stereo-chemistry, atomic clash and misfolding. A detailed analysis of the amidase-03 model structure from *Bacillus anthracis* will provide insight to the molecular design of suitable inhibitors as drug candidates.

**Keywords:** Homology modeling; modeller; amidase-03; hydrolase enzyme; *Bacillus anthracis*,

**Background:**

*Bacillus anthracis* secretes a number of enzymes capable of hydrolyzing the cell wall peptidoglycan. The hydrolysis of the cell wall imparts an important role in maintaining its structure. The structure of the cell wall therefore continuously undergoes modifications such as hydrolysis, insertion and release of old molecules [1, 2, 3]. Some enzymes, called autolysins cause cell lysis where processes like cell separation, cell wall turnover and competence occur. Autolysins also play an important role in the amplification of inflammatory responses [4]. These hydrolyzing enzymes are divided into N-acetylmuramidases, N-acetylglucosaminidases, and N-acetylmuramyl-L-alanine amidases based on substrate specificity [3]. The etiologic agent of anthrax, *Bacillus anthracis* contains two virulent plasmids pXO1 and pXO2 causing cutaneous, gastrointestinal and inhalation anthrax. *B. anthracis* produces protective antigen (83 kDa), edema factor (84 kDa) and lethal-factor (83 kDa) causing inflammatory response with the release of toxic compounds and mediators [5]. The genome sequencing and annotation shows the presence of several amidases like proteins in the *B. anthracis* genome. The most studied amidase gene from *B. anthracis* is amiA, belonging to class amidase-03 of autolysins [4]. Many autolysins usually increase the inflammatory response. However, amidase-03 decreases the inflammatory response in host by hydrolyzing the bond between glycan and peptide thereby reducing the amplification to response [4].

**Figure 1: Sequence alignment of amidase-03 (PDB ID: 2IR0) with template 1XOV**
Methodology:

Template Selection

Template selection was done using BLASTp for the query sequence against PDB (Protein Data Bank) available at NCBI [10]. BLOSUM62 was applied as matrix. The target subsequently selected was the x-ray crystal structure with PDB code 1XOV with resolution 1.8 Å and R-value 0.122. Sequence alignment for template 1XOV and query (Figure 1) was done by using python script align2d.py in Modeller [11].

Generation of the initial structures and model evaluation

Sequence alignment was used to generate several 3D models of amidase-03 using auto-model class. Five models were built by Modeller default routine in which initial coordinates are randomized and evaluated by pseudo-energy parameter. If several models are calculated for the same target, the best model can be selected in several ways. We evaluated the model with the lowest value of the Modeller objective function and PROCHECK statistics, which is also recognized by ADIT tool at PDB Server [12]. The stereo-chemical properties were assessed using PROCHECK [13].

Energy minimization of the structure: Molecular Dynamics

Energy minimization of the selected model was done by using steepest descent for removing the bad vander-waal contacts from the model. The calculations were done by using GROMACS 3.2 [14] software with G43a1 force field with flexible SPC water model in cubic box of size 1 Angstrom instead of rigid SPC model.

Visualization and structural analysis

Structure analysis and visualization were carried out with Visual Molecular Dynamics (VMD) [15] software. Structural alignment and RMSD calculation with template and several models were also done with VMD.
Discussion:

BLAST results had shown higher similarity with the crystal structure of 1XOV and 1JWQ, while the template was selected on the basis of greater identity. 1XOV template showed 35% sequence identity, 47 conserved residues and 55% of sequence similarity with amidase-03, while the 1JWQ showed 26% sequence identity and 55% sequence similarity. A putative domain was also found against pfam database entry pfam01520, a family of hydrolase enzyme consisting of catalytic domain which cleaves peptidoglycan of the bacterial cell wall by MurNAc-L-Ala linkage and has 51 conserved regions in BLASTp results (Figure 2). So, the 245 amino acids protein from B. anthracis genome encoded by GenBank accession NP_844822 consists of amidase-03 domain known to have catalytic functions to hydrolyze the cell wall. This protein in this study is referred as amidase-03 hereon.

The template structure 1XOV consists of two domains, a C-terminus and N-terminus while we used only N-terminus of the protein to build the model. The Figure 1 shows the alignment of 171 residues of amidase-03 with 316 residues of 1XOV. After 175 initial residues, Modeller put a long gap with C-terminal of the template as we used only N-terminus of the protein to build models of amidase-03. Due to the high target and template similarity, there are only two gaps inserted at 96th and 97th position in the N-terminus of the sequence alignment. Using crystal structure for Homology modeling provides high quality structure alignment for structure prediction [16]. Phi and Phi torsion angles of predicted models were checked by PROCHECK. A comparative table can be found with supplementary data (Table 1). In Ramachandran plot analysis, one residue of each model was found in disallowed region. For a conformational asset, a PROCHECK analysis was done with template 1XOV, which also confirmed one residue in disallowed region of template. So, the probable error can be negotiated here. The total number of residues in selected model was found as: 94.3% residues in most favored region, 3.8% in additional allowed region and 1.3% in generously allowed region of the Ramachandran plot.

Structural alignment was done using VMD to calculate the structure deviation between template (1XOV) and selected model of amidase-03 of B. anthracis (Figure 3). The RMSD between template and selected model showed a value of 0.0976 nm, which is an indicative of a good quality predicted model. However, all predicted models had less than 0.15 nm RMSD values. The predicted model of amidase-03 consists of six α-helices and six β-sheets. All β-sheets are surrounded with helices forming inner core of the protein and providing the amidase moiety. The overall confirmation of the model was found similar to N-terminal of 1XOV (Figure 3), which is an enzymatically active autolysin as reported in Listeria monocytogenes bacteriophage PSA endolysin PlyPSA [17]. These autolysins have unusual substrate specificity and high activity towards degradation of the murein layers and lysis of susceptible cells [17]. Based on their antimicrobial properties and substrate specificity, these enzymes show great potential to be effective therapeutic agents [18]. The increasing resistance to the known antibiotics has driven attention towards finding newer ways to combat pathogenic organisms and thus the investigation of the structures for the development of new drugs/ drug targets from the molecules like amidases have become reasonably essential especially for pathogenic organisms like B. anthracis.

Conclusion

We described the model structure for 245 amino acids amidase-03 from Bacillus anthracis using x-ray crystal structure of Listeria monocytogenes bacteriophage PSA endolysin PlyPSA (PDB ID: 1XOV) as template. The predicted structure was submitted to Protein Data Bank (PDB-id 2IR0). Molecular dynamics simulation of the model is described elsewhere [9].

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