**In silico** Characterization of Biofilm-Associated Protein (Bap) Identified in a Multi-drug Resistant *Acinetobacter baumannii* Clinical Isolate

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**Introduction:** *Acinetobacter baumannii* (AB) is a Gram-negative bacteria associated with various hospital infections. The present study deals with in silico analysis of the biofilm-associated protein (Bap) in this pathogen. **Method:** Sixty-eight multi-drug resistant (MDR) AB were isolated from two hospitals in Kerman, Iran. Biofilm-formation was investigated using the microtiter method and PCR followed by sequencing to detect the bap gene in the strongest biofilm-forming isolate. The physicochemical parameters of Bap protein were determined by the ProtParam tool using the ExPasy program. The 3D models from the primary amino acid sequence were constructed using the I-TASSER modeling platform based on multiple-threading alignments by LOMETS. Nevertheless, to ensure the correct initial structure, the protein was minimized in energy through the 3DRefine software of the deep learning system. For the accuracy of predicted models, calculation of the orientation of dihedral angles, including the phi (φ) and psi (ψ) and backbone conformation using the PROCHECK module of the PDB Sum server was performed. The domains and key amino acids involved in protein structure were studied by the Pfam and Interpro softwares. **Results:** Analysis of the amino acid content of the Bap protein revealed the absence of Arg and Cys in the protein structure. Our Bap protein exhibited ~99.6% identity with other Bap sequences in the GenBank database. Stereochemical simulation identified 19 antiparallel β-sheets with two small α-helices. The N-terminal of Bap protein formed oligomers that mediate cellular adhesion. **Conclusion:** This study adds considerable information about Bap protein 3D structure, its conformation, domain analysis, and amino acids involved in cellular attachment.

**INTRODUCTION**

*Acinetobacter baumannii* is a Gram-negative pathogen associated with various hospital infections, particularly in intensive care units (ICU) of hospitals worldwide [1]. This species accounts for ~80% of reported *Acinetobacter* infections, including ventilator-associated pneumonia, bacteremia, meningitis, peritonitis, urinary tract infections, and wound infections [2]. With the emergence of multi-resistant, and pandrug-resistant strains of this bacterium, the treatment process has become challenging, resulting in increased death rates among immunocompromised patients [3]. One of the outstanding *A. baumannii* features is the biofilm formation on medical devices and living tissue, representing a crucial microbial virulence factor [4]. The biofilm-forming *A. baumannii* isolates are intrinsically resistant to many antibiotics, including carbapenems, fluoroquinolones, and third-generation cephalosporins [3]. One essential protein contributing to biofilm-formation in this bacterium is the biofilm-associated protein (Bap) [5].

The genus *Acinetobacter* hosts various large multidomain Baps proteins, all exhibiting similarity in structure, as they are internally repetitive and feature multiple (3-50 copies) immunoglobulin-like (Ig-like) domains. These domains have a particular three-dimensional structure known as Ig fold, comprising 70-100 amino acid (aa) residues in seven antiparallel β-strands organized in two β-sheets packed against each other in a sandwich structure [5]. Furthermore, size heterogeneity indicated changes in the number of amino acids due to recA-independent slipped-strand mispairing during DNA replication. Nevertheless, in the A.

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*baumannii* hospital isolates, the core unit of the Bap protein, is highly conserved [7]. Considering that Bap protein promotes adhesion to biotic and abiotic surfaces, it seems reasonable that the fragments containing the N-terminus of the protein become aggregation-prone and self-assemble into amyloid-like structures under acidic pH and low concentrations of calcium [8].

Our previous study detected a *bap* gene in a potent biofilm-forming *A. baumannii* isolate and analyzed its expression [9]. However, information on the structure of a large part of Bap protein and its domains activities is still poorly understood. The present study deals with the physicochemical characterization, stereochemical simulation, protein modeling, amino acid alignment, and phylogenetic tree analysis of a Bap protein obtained from the strongest biofilm-forming MDR *A. baumannii* clinical isolate identified herein.

**MATERIALS AND METHODS**

**Bacterial sources and antimicrobial susceptibility testing.** Bacterial source, gene identification, and antimicrobial susceptibility testing of 68 MDR clinical isolates of *A. baumannii* were determined as previously reported elsewhere [4].

**Biofilm formation.** Biofilm formation was performed by microtitre assay as described previously [12]. The quantification of cells in the biofilms was carried out by solubilization of dye with 200 µL of absolute ethanol at optical density (OD) of 480 nm. The OD cut-off value (ODc) was established as three standard deviations (SD) above the mean optical density (OD) of the negative control as shown in the formula: ODe=average OD of negative control +(3_SD of negative control). According to the optical density, the isolates were divided into four categories: (1) strong biofilm producer (4_ODc<OD), (2) medium biofilm producer (2_ODc<OD_4_ODc), (3) weak biofilm producer (ODc<OD_2_ODc), and (4) no biofilm formation (OD_ODe). Among the strong biofilm producers, one that showed the highest OD was selected for further studies. The assays were performed in duplicates, and *Pseudomonas aeruginosa* PAO1 was used as biofilm positive strain.

**Detection and sequencing of *bap* gene.** A clinical *A. baumannii* isolate that showed the strongest biofilm formation was selected and subjected to PCR using specific primer sequences to detect the *bap* gene [13]. The PCR product was sequenced by ABI Prism 373 DNA Sequencer (Applied Biosystems 373/3730XL, Bioneer, Korea) as described previously [4]. The generated sequence was compared with similar sequences deposited in the National Centre for Biotechnology Information (NCBI) database using BLAST comparison algorithm with default settings (https://www.ncbi.nlm.nih.gov/genbank/blast), and similarities were obtained.

**Nucleotide sequence accession number.** The *bap* gene and protein sequences were deposited in the GenBank database under the accession numbers KR080550 and AKL78797, respectively.

**Physicochemical characterization of the Bap protein.** Physicochemical properties of the Bap protein such as molecular weight, theoretical isoelectric point, atomic composition, extinction coefficient, instability index, aliphatic index, and grand average of hydropathicity were determined using the ExPasy ProtParam server (https://web.expasy.org/cgi-bin/protparam/protparam). Furthermore, the amino acid compositions based on percent quantity were determined by ExPasy ProtParam software.

**Classification, functional analysis, and phylogenetic tree of Bap protein.** The primary classification and functional analysis of Bap protein were determined using the InterPro software ver. 5.31-85 (www.Ebi.ac.uk/interpro); it provides an integrative classification of protein sequences into families and domains. The biological functions of the PmrA protein, including ligand-binding site, molecular and cellular functions in terms of Gene Ontology (GO), were inferred from the PDP database based on comparing sequence and structural similarity using the I-TASSER server. Nevertheless, the phylogenetic tree was constructed using the unweighted pair group method with arithmetic mean (UPGMA). The sequence was subjected to the amino acid homology analysis using the BLAST tool obtained from the UniProtKB database (https://www.uniprot.org/). The Bap amino acid sequences displaying the maximum identity to the query sequence were used for alignment and construction of phylogeny [15]. The dendrogram from the distance matrix was generated in MEGA 6 software using the maximum likelihood (ML) method based on the JTT matrix-based model. The bootstrap test of phylogeny was assessed with 1000 repetitions [15].

**Three-dimensional structure of Bap protein.** For constructing the correct 3D protein model from the primary Bap amino acid sequence, the iterative threading assembly refinement (I-TASSER) online software based on multiple-threading alignments LOMETS was used (https://zhanglab.dcmib.med.umich.edu/I-TASSER/). Five closest stereochemical structures were obtained, and the confidence of each model was quantitatively measured by confidence (C)-score. The C-score was calculated based on the significance of threading template alignments and the convergence parameters of the structure assembly simulations. The C-score is generally within the -5 to +2 range. Higher C-score signifies higher confidence in the model. Subsequently, the quality of the structure was estimated by the C, TM, and RMDS scores. In addition, the stability of the models was estimated using the structural validation algorithm ProSA [16]. For confirming the accuracy of the protein secondary structure, the PSSpred algorithm

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was used as described previously [17]. The orientation of dihedral angles (torsion) angles phi (Φ) and psi (ψ) were determined using a logistic representation Z-score of the Ramachandran plot and the SAVES program (https://saves.mbi.ucla.edu/). The predicted models were then subjected to the backbone conformation using the PROCHECK module of the PDBSum server (https://www.ebi.ac.uk/thornton-srv/software/PROCHECK/). Finally, a template with the highest quality was selected for the model building based on the confidence score of the Deep Learning Potential [18, 19].

Structural similarity search. For searching the structural similarity of the Bap protein, the MADOKA webserver engine (http://madoka.denglab.org/) was used. The server consisted of two-phase, searching similar protein structures by aligning the input with the whole PDB library.

Table 1. The physicochemical properties of Bap protein obtained in this study.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Molecular weight</td>
<td>39557.83</td>
</tr>
<tr>
<td>Number of amino acids</td>
<td>396</td>
</tr>
<tr>
<td>Theoretical pI</td>
<td>3.46</td>
</tr>
<tr>
<td>Total number of negatively charged residues (Asp + Glu)</td>
<td>56</td>
</tr>
<tr>
<td>Total number of positively charged residues (Arg + Lys)</td>
<td>7</td>
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Atomic composition

| Carbon C | 1718 |
| Hydrogen H | 2685 |
| Nitrogen N | 445  |
| Oxygen O  | 624  |
| Sulfur S  | 0    |

Formula 718H2685N445O624 5472 85.25 41940 1.060 1.060 15.48 -0.016

Extinction coefficients are in M⁻¹ cm⁻¹, at 280 nm measured in water. Abs 0.1% (g/L) 1.060. The aliphatic index of a protein was calculated according to the following formula: Aliphatic Index = X(Ala) + a X(Val) + b (X(Ile) + X(Leu)) where X(Ala), X(Val), X(Ile), and X(Leu) are mole percent (100 X mole fraction) of alanine, valine, isoleucine, and leucine.

Amino acids compositions of Bap protein. The primary analysis of the amino acid content of the Bap protein revealed an absence of Arg and Cys, whereas Glu with 9.2%, Lys with 7.8%, and Trp and Tyr with 6.1% were the most abundant amino acids residues in the protein structure (Fig. 1).

The classification, structural diversity, and phylogenetic tree. The InterPro tool classification of the generated Bap protein sequence included the superfamily Ca_tandamer, containing three regions, namely Big_3 and Big_16 and Ig-like sequences. The sequences inserted under each domain indicated order predictability (Fig. 2A). Interestingly in our case, the Ig-like sequence begins with Gly at position 111, while Big_3 and Big_16 sequences begin with Ala at position 217 and Asp at position 304, respectively. The Bap protein belonged to the Ig-like homologs superfamily with cell adhesion property and calcium-binding tandamer activity. Moreover, hierarchical information based on the
Structural diversity of this superfamily was compared to other superfamilies in the CATH database, which is a protein structures classification tool in the Protein Data Bank (Fig. 2B); protein structures are chopped up into domains and grouped into superfamilies, and the evolutionary relationships among them are determined through a combination of automated algorithms and expert manual curation. The sequence diversity was ~73%, indicating a relatively low diversity (Fig. 2B). This result was supported by phylogenetic tree analysis of Bap protein which showed 99.6% identity with other Bap sequences (Fig. 2C). The analysis showed 99.6% identity with *A. baumannii* MBO4071943.1 and 99.4% with *A. baumannii* WP_140983092.1, respectively.

**Sequence Alignments.** Multiple amino acid sequence alignments of the Bap protein (Fig. 3) indicated that the "IWAKFKDAENDADADSDADADADS" consensus sequence was responsible for extracellular adhesion, bringing together bacterial cells. Similarly, the surface binding domain sequence is demonstrated by yellow color. This region seems to be conserved in many Bap proteins. No amino acid substitution and mutation were detected in this study.

**Structure and quality of Bap protein.** The cartoon structure of Bap protein obtained by the I-TASSER webserver (Fig. 4A) contained 19 antiparallel β-sheets with two small α-helices. The stereochemical trajectory of the protein exhibited three main regions and one surface N-terminal oligomers that mediate biofilm development. This sequence was conserved in many Bap proteins. In contrast, the C-terminal region was membrane bond and variable in amino acid composition. The region-A consisted of 5 antiparallel β-sheets while region-B had six antiparallel β-sheets (Fig. 5A). Interestingly, region-D was situated in the C-terminal site of protein consisting of 6 β-sheets and one small α-helix.

**Quality estimation.** The results of the Ramachandran plot (Fig. 5A) indicated a clash score of 13.68%, Ramachandran favored 86.08%, and mole probability score = 2.49. The predicted contact and distance probability matrix of Bap protein (Fig. 5B) was displayed by darker color as a higher probability. Furthermore, the contact map of the bap protein defined by Cb-Cb distance showed ≤ 8 angstroms. Furthermore, ASP 273, ASN 149, and THR 246 were situated in the non-favored region. The predicted contact and distance probability matrix of Bap protein maps were also determined in the LOMETS platform (Fig. 5B). The darker color and increased yellow color intensity indicated more accuracy of model structure. Furthermore, to predict the exact secondary structure of Bap protein, five highly similar models were obtained. The highest C-score of 0.75 was selected for further study.

**Amino acids involved in adhesion.** The schematic representation of the amino acids involved in cellular adhesion of Bap protein (Fig. 6) showed that Pro144, Ile 143, Val 138, Tyr 161, Asp 146, Ile 136, Asn 142, and Lys 141 play an important role in biofilm formation.
Stereochemical analysis of Bap protein

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Fig. 2. A) Classification of the submitted sequence of the Bap protein in InterPro tool software. The Ig-like domains have three motifs, fitting the consensus TDnAGN, found in many bacterial surface proteins. B) Overview of the sequence and structural diversity of Bap superfamily by CATH system. C) The phylogenetic tree analysis of Bap protein. The analysis showed a close identity of Bap protein with other similar proteins reported in the GenBank database.
Fig. 3. Multiple amino acids alignments of Bap protein with closely related sequences using UniProtKB database (https://www.uniprot.org/help/sequence-alignments).
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Fig. 4. The generated cartoon structure of Bap protein. A) The protein consisted of 19 antiparallel β-sheets with two small α-helices. The amino acids in α15, α16, and α18 are located in the C-terminal and anchored to the membrane. B) Bap protein regions and C-TM-RMSD scores of a highly accurate model of Bap protein. The model contained 4 regions where N-terminal forms oligomers and mediates biofilm development. This sequence was conserved in many bap proteins. In contrast, the C-terminal region was membrane bond and variable in amino acid composition. C) Domains analysis of Bap protein.

DISCUSSION

Bacteria can assume two modes of growth: unicellular or planktonic cells and multicellular forms or biofilms. The planktonic bacteria have better freedom of migration, are more prone to mutation but are more sensitive to the environment and antibiotics. In the multicellular life phase or biofilms, they are better protected from the harsh environment, are less subjected to mutation, and are more resistant to antibiotics and disinfectants [22]. Most of the available information on the Bap protein is related to Staphylococcus aureus, but the data can most likely be attributed to the other Bap orthologs. The A. baumannii bap gene encodes an extensive surface protein, comprising 8,621 amino acids homologous to the Bap protein first described in S. aureus [7].

The protein structure predicted in this study using homology modeling indicated model 1, the best Bap protein model in A. baumannii that shows strong biofilm. Multiple sequence alignment resulted in significant sequence identity to similar sequences that existed in the UniProtKB database. This close identity was observed mainly in conserved regions such as the N-terminal consensus sequence and surface domain through homophilic interactions between coiled-coil domains of opposing proteins in neighboring cells. The results were further supported by dendrogram analysis which suggested 99.6% homology with those Bap-like prefix domains containing protein. The protein that belonged to the superfamily Ca_tandamer contained Ig-like domains; these domains are found in a wide variety of extracellular bacterial proteins, often in multiple copies [15].
Stereochemical analysis of Bap protein revealed a monomeric high molecular weight protein that contained 4 regions well suited in X, Y, and Z axes. The high quality of the model structure was determined by I-TASSER software and approved by the ExPasy modeler. The amino acid residues position in segments from the logistic representation of high-fidelity Ramachandran plot data of the model protein structure showed mainstream $\delta^2$-region near 86% of the residues, representing the most favorable combinations of phi and psi values. The $\Phi$–$\Psi$ population density in the Ramachandran plot represents the total folding free

Fig. 5. A) Ramachandran plot for quality of the 3D structure of Bap protein. B) Predicted contact and distance map of Bap protein using the LOMETS program.
energy. Further analysis of amino acid residues of Bap protein in the present study revealed Glu with 9.2% as the most frequent amino acid, while Arg and Cyst were absent. In a similar study [7], analysis of the primary structure of the Bap protein showed absence of Cys and an abundance of Thr (1,389 residues; 16% of the total), Ala (1,176 residues; 14% of the total), and Val (1,109 residues; 13% of the total) [7].

Previously, analysis of the physicochemical characteristics of Bap revealed that the protein was acidic (estimated isoelectric point [pI] = 3.37) with a molecular weight of 220959.8 Da [23]. Furthermore, the core region of Bap represents 52% of the protein consisting of 13 successive nearly identical repeats, each containing 86 amino acids and two canonical calcium-binding EF-hand motifs able to control Bap functionality in response to the calcium present in the growth media [23]. Surprisingly, a large part of the protein is not required for the adhesion-accumulation biofilm phenotype, and that Bap-mediated biofilm development occurs in response to environmental cues [24].

Despite all the studies supporting the crucial role of Bap protein in biofilm formation, a few reports have described the detailed stereochemical structure of this protein in high biofilm-producing A. baumannii isolates [25]. This study, therefore, paved the way for better understanding the structure, amino acids similarities, modeling prediction, and domains analysis of Bap protein in A. baumannii isolates.
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CONFLICT OF INTEREST

The author declares there are no conflicts of interest associated with this manuscript.

REFERENCES


