In Silico Investigation for Evaluation of the Potential of the SclA Protein in Streptococcus pyogenes

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Background: Streptococcus pyogenes is an important pathogen that is associated with a range of infections in humans, and causes common and severe invasive diseases. Currently, antimicrobial therapy is the first choice for the treatment of S. pyogenes; however, the emergence of antimicrobial resistance and side effects of antibacterial drugs is increasing. Consequently, there is an increased demand for novel drug targets and vaccine design.

Objectives: To develop an effective vaccine against Streptococcus pyogenes (group A streptococcus), we described a novel collagen-like surface protein of S. pyogenes which is important virulence factors

Materials and Methods: In this study, we focused on the SclA protein of S. pyogenes and characterized it using bioinformatic tools to introduce it as a candidate novel drug as a candidate for use in vaccine design. The secondary structure was determined and the 3D structure was modeled using SWISS-MODEL workspace. The immune epitope database analysis (IEDB) resource was used to predict regions of SclA that are likely to be recognized as epitopes.

Results: The SclA protein is present on the cell surface of the cell and has interact with a common ligand by its hypervariable NH2-terminal regions. The IEDB showed that the maximum peptide length that is likely to be predicted as an epitope is of 6 amino acids, from amino acid 26 to 31, with a score of 4.791. This epitope can be considered for use in Antibody and drug design.

Conclusions: Data from this study about SclA were not sufficient and further studies are needed; however, the information here suggests that SclA could be a candidate for further research on the design of drugs and vaccines against S. pyogenes infections.

Keywords: Computer Simulation; Streptococcal Collagen-Like Protein A; Vaccine Candidate; Streptococcus pyogenes

1. Background

Streptococcus pyogenes (group A streptococcus, GAS) is an important pathogen associated with a range of infections in humans (1). This organism causes common diseases such as pharyngitis, erysipelas, and impetigo, but sometimes can also cause severe invasive diseases such as toxic shock-like syndrome and necrotizing fasciitis (2, 3). Currently, antimicrobial therapy is the first choice for the treatment of GAS; however, antimicrobial resistance is a common problem worldwide and the consumption of antimicrobials has been identified as a driving force for the emergence and transmission of antimicrobial resistance in GAS (4). Streptococcus pyogenes has very complex virulence factor (5). The cell wall of S. pyogenes is associated with a range of proteins, a large number of which are encoded by the emm gene family: M proteins are the members that contain the most important structures for virulence (5-7).

Other cell wall proteins in S. pyogenes include streptococcal collagen-like surface protein (ScIA) (8). ScIA is transcriptionally regulated by Mga and has a highly conserved signal sequence; in a previous study, ScIA was found in 12 S. pyogenes strains tested by PCR (9). Since the M protein is a major virulence factor of GAS, it could be a candidate for use in vaccine design (10). However, previous studies have shown that some epitopes of M proteins can induce autoimmunity and cross-react with human tissues (9, 10), because of which they may not be suitable for use in vaccine design.

2. Objectives

In this study, we characterized the ScIA protein of S. pyogenes, using online and off-line computational tools for the development of new vaccines to prevent and treat pyogenesis.

3. Materials and Methods

3.1. Retrieval of the Target Sequence

The candidate organism for the present study was S. pyogenes. The protein sequence of ScIA (GenBank: CAC33784.1) was obtained from NCBI (http://www.ncbi.nlm.nih.gov), and a BLAST alignment was performed against Protein Data Bank (PDB) entries to find similar sequences.
3.2. Functional Characterization of SclA

Functional characterization of the SclA protein sequence was performed using Uniprot (http://expasy.org). Primary structure prediction was computed using the ExPASy ProtParam server (http://expasy.org/cgi-bin/protparam).

3.3. Protein Topology Prediction and 3D Structure of SclA

The secondary structure of the SclA protein was obtained using the PSIPRED Protein Sequence Analysis Workbench (http://bioinf.cs.ucd.ie/psipred/). Using the TMHMM database, we predicted transmembrane helices based on a hidden Markov model and predicted the locations of transmembrane intracellular and extracellular regions. Garnier- Osguthorpe- Robson (GOR) IV was used to predict alpha helices, beta sheets, and random coil secondary structures. The three-dimensional structure of SclA was obtained using http://www.ncbi.nlm.nih.gov/Structure/VAST.

3.4. B Cell Epitope Prediction

The immune epitope database analysis resource (IEDB-AR, http://tools.immuneepitope.org) provides a collection of web-based tools for the prediction and analysis of immune epitopes. In this study, IEDB was used to predict regions of SclA that are likely to be recognized as epitopes in the context of a B response. Conservancy analysis for epitopes and the percentages of protein sequence matches at identity ≥ 100% were also calculated.

4. Results

In silico analysis can be used for identifying drug targets, designing vaccines, and choosing a rapid strategy for disease control. The sequence of the SclA protein was obtained from the NCBI protein sequence database and a BLAST search was performed against Protein Data Bank (PDB) entries for sequence similarity. SclA was found to be associated with 3HQV_B, a protein from Rattus norvegicus. The score for protein alignment between SclA and 3HQV_B was 30.0, with a 46% identity. PROSITE analysis showed that SclA was a surface-localized protein. Primary structure prediction showed that SclA had 123 amino acid residues and an estimated molecular weight of 13137.6. The total number of positively charged residues (Arg + Lys) was 22 and the total number of negatively charged residues (Asp + Glu) was 21. The very high aliphatic index (48.54) and instability index (36.76) provided an estimate of the stability of protein in a test tube. The Grand Average Hydropathicity (GRAVY) value was -0.956.

The secondary structure prediction showed that the extended strand (10.57%) and alpha helix (36.59%) were the least frequent, whereas the random coil was more frequent at 52.85%. As the 3D structure of SclA had not been reported in Protein Data Bank (PDB), a BLASTp similarity search was performed against the PDB database and 3HQV_B, where a complete query coverage was obtained (Figure 1). By using PSIPRED, we showed the location and spatial arrangement of each amino acid separately (Figure 2). TMHMM predicted the locations of transmembrane intracellular and extracellular regions and showed that the SclA protein was present on the outside of the cell (Figure 3). The results of the IEDB showed that the maximum length of a peptide that is likely to be predicted as an epitope was of 6 amino acids between positions 26 and 31, with an epitope sequence of QERTRK and a score of 4.791 (Figure 4).
stains of the same serotype and different genotypes. Unfortunately, an effective vaccine to prevent GAS infection is yet to be developed. Secreted or surface proteins or lipoproteins are favored candidates for vaccines because they are the most easily accessible microbial antigens to the immune system. As SclA is a surface protein, it may be a suitable candidate for vaccine design.

The focus of the current study was therefore to characterize the SclA protein of *S. pyogenes* using computational tools. SclA had 123 amino acid residues with a molecular weight of 13,376. The total number of positively charged residues was more than that of negatively charged residues. The N-terminal sequence is considered to contain Alainin (Ala), Garnier-Osguthorpe-Robson (GOR) IV comparative analysis showed that a random coil is predominantly present, followed by an extended strand and an alpha helix. TMHMM determined that the SclA protein is present on the outside of the cell. The results of IEDB showed that the maximum length of the peptide that is likely to be predicted as an epitope is of 6 amino acids, from amino acid 26 to 31, with a score of 4.791 and a protein sequence identity match of 100.00% (1/1). These epitopes can be considered for antibody and drug design.

The information on the characterization of the SclA protein of *S. pyogenes* obtained from these studies alone is not sufficient. Nonetheless, this information suggests that SclA could be a candidate for more rigorous studies on drug and vaccine design against *S. pyogenes*.

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**Authors’ Contributions**

Maryam Pourhajibagher developed the original idea and the protocol, abstracted and analyzed data, and drafted the manuscript. Abbas Bahador contributed to the development of the critical revision of the manuscript for important intellectual content, and is the guarantor.

**References**


