Designing of potential vaccine candidates of fused cathepsin L and tropomyosin genes of Rhipicephalus (Boophilus) annulatus tick larva

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Abstract

Introduction: Regarding emerging tick resistance against acaricides, researches have been shifted toward alternative approaches such as immunologic methods. Vaccine preparation is an alternative way in which choosing appropriate protein with high immune induction potency is a prerequisite. In addition according to studies, using more than one protein could better enhance the immune induction and antibody production. Choosing immunogenic epitopes from selected proteins and adjoining them with a suitable linker is one of the novel approaches in vaccine design.

Materials and methods: Based on the fact that both cathepsin and tropomyosin proteins of Rhipicephalus tick were previously recognized as potent immunogenic antigens, we predicted the immunogenic epitopes of these proteins by immunoinformatic methods. Among studied epitopes, those that were met by multiple bioinformatics tools were used.

Results: Finally, the polytopic construction was designed by assembling the selected epitopes and connecting them with linkers.

Conclusion: Using immunoinformatic tools, we predicted the characteristics of two genes of Rhipicephalus annulatus tick larva as fused potent vaccine candidates namely, cathepsin and tropomyosin.

Keywords: Tropomyosin, Cathepsin, Rhipicephalus annulatus, vaccine design, Iran

Introduction

The cattle tick, Rhipicephalus (Boophilus) annulatus (Family Ixodidae), is one of the most abundant tick species worldwide specifically in tropical and subtropical territories (1). This species is mainly found in North of Iran (2). Cattle is the principal host for this obligate ectoparasite, but it has also been found occasionally in horse and deer. Rhipicephalus is considered as a potent vector for some bacterial and parasitic pathogens such as Anaplasma and Babesia (3). Additionally, tick infestation usually provokes a notable financial burden in livestock industry through blood
loss, weight reduction, decreased lactation, anorexia, irritation, and toxic conditions such as paralysis and allergies (4). The tick control programs have frequently been relied on the utilization of anti-tick chemical compounds, i.e. acaricides. However, the emergence of acaricide-resistant tick populations and chemical residuals persistence in animal productions and environment justify the demand for other prevention strategies (5). In this case, vaccination against tick infestation would be an alternative substitute; therefore, it is recommended to study and characterize different tick antigens with likely suitable immunogenicity properties (6).

One of the suggested anti-tick vaccine candidates is the actin-binding tropomyosin (TRM) protein. The molecular structure of this protein includes a coiled-coil dimer which is composed of two parallel-oriented tropomyosin molecules with α-helical arrangement (7). This molecule assists in different cell functions, such as intracellular trafficking, cell morphology and cellular contractile force. In invertebrates, several isoforms of tropomyosin have been recognized in muscular and other tissues. Nevertheless, this protein plays a substantial role in immunogenicity and allergic reactions against ticks and mites so it is an appropriate vaccine candidate against the mentioned parasites (8, 9).

Cathepsins (CT) are lysosomal proteases with several members such as cysteine, aspartic, and serine proteases. The physiologic actions of cathepsins are enclosing degradation of intracellular and endocytosis-introduced extracellular proteins. Among them, cysteine proteases in particular are involved in blood degradation that is one of the most important biological processes in ticks. Ticks provide their energy for laying eggs, molting, and other related processes from blood digestion. CT L is one of the members of cysteine protease in Rhipicephalus (Rh) microplus which is able to hydrolase tick vitellin and bovine hemoglobin at acidic PH (10, 11).

Currently, some polytopic vaccines are evaluated in clinical trials using specific epitopes against infectious agents, cancers, and autoimmune disorders (12-14). In this study, we aimed to predict the characteristics of two genes of Rh.annulatus tick larva as a fused potent vaccine candidates namely CT and TRM, using immunoinformatic tools.

Materials and methods

Protein sequence retrieval: Since the TRM and CT protein sequences of Rh. annulatus did not exist in reference databases, the desired sequences of a close species, Rh. microplus TRM, containing 284 amino acid (O97162) and CT containing 332 amino acid (Q9NH5) were retrieved from UniProtKB database (http://www.uniprot.org/) and saved in FASTA format.

Homology modeling and its validation: The three dimensional (3D) structure of CT and TRM of Rhipicephalus or even other species of ticks was not found in the protein data bank (PDB), so we performed protein modeling. Two most reputable servers in this field are the online servers of the ITASSER (https://zhanglab.ccmb.med.umich.edu/I-TASSER/) and SWISS-MODEL (https://swissmodel.expasy.org/). In order to check the potential errors in modeled 3Dstructure, we used Ramachandran plot analysis at http://mordred.bioc.cam.ac.uk/rapper/rampage.php.

Secondary structure prediction: Based on PSIPRED server (http://bioinf.cs.ucl.ac.uk/psipred/) and GOR IV server (https://npsa-prabi.ibcp.fr/cgi-bin/seqpred_gor4.pl), secondary structure was predicted and analyzed.

Prediction of discontinuous B-cell epitopes: Discontinuous B-cell epitopes were predicted from 3D structure of
proteins epitopes in PDB format by IEDB database at http://tools.iedb.org/main/analysis-tools/.
In order to enhance the reliability, the epitopes were predicted with others bioinformatics servers. After determining B-cell epitopes, we aligned the sequences of predicted epitopes of CT and TRM of the tick with these proteins in cattle using NCBI Blast server for annotating and final selecting of epitopes based on dissimilarity as much as possible.

Prediction of physico-chemical characteristics: These profiles were illustrated using BcePred servers, IEDB, and ExPAsy-ProtScale (http://www.expasy.org/). The considered factors included hydrophathy, availability, flexibility, and exposed surface. The hydrophilic and hydrophobic profiles were estimated on an algorithm introduced by Kyte and Doolittle. (15) Also, surface availability and flexibility of different protein regions were appraised by Emini algorithm and Karplus and Schulz algorithm respectively. Among the predicted epitopes, those with more availability and the so-called exposed surface were selected using Chimera software. In the end, the construct was investigated in terms of allergenicity using the AllerTOP v. 2.0 server (http://www.ddg-pharmfac.net/AllerTOP/). The online server SOLpro (http://scratch.proteomics.ics.uci.edu/) was used to investigate Solubility and Antigenicity.

Protein structure analysis: Prosa was used to analyze protein structure at https://prosa.services.came.sbg.ac.at/prosa.php. The overall model quality is shown by the z-score. In fact, z-score in such models contains the z-scores of all protein chains that have been experimentally determined in current PDB. In this plot, different colors are used to distinguish groups of structures from sources (X-ray, NMR).

Construct designing: Sequences of selected epitopes were linked using suitable linker and then the final construct was back translated to DNA at http://web.expasy.org/translate/ and codon of final sequences were optimized in E.coli by jcat online server (expasy/codon optimization tools) (http://www.jcat.de/). Then restriction enzymes (EcoRI/XhoI) were added to 5´ and 3´ ends of sequences and synthetic DNA fragment was cloned into the pET32a. This vector is appropriate for cloning and high-level expression. This is due to the fact that its peptide sequence has been fused with a tag (109aa Trx•Tag thioredoxin protein).

Result

Selection of epitopes: Finally using BcePred and IEDB servers, two high immunogenic epitopes from each TRM and CT were selected as B-cell epitopes as shown in Table 1. Selection of these epitopes was according to immunoinformatics analysis and all of them had the highest score. To maintain the three-dimensional structure of epitopes, the epitopes of TRM and some amino acids between them were chosen as a larger size epitope.

Table 1. Predicted B-cell epitopes from target antigens.

| Target antigen | Position | Sequence |
|----------------|----------|----------|
| Tropomyosin    | 168-188  | KLAMVEADLERAEARTGET |
|                | 197-222  | RVVGNLKSLEVSEEKALQEETYE |
| Cathepsin      | 18-72    | SQEILRTQWEAFKTHKKSYQSHMEELRRFKIFTENSLLIAKIHNAKAYAKGLYSY |
|                | 86-142   | FARIIFGHHHTRKGTGGSTFLPPANVNDSSLPKVDWKKGAFTVLPKDKQGCSCWA |

Linker: Since our epitopes are based on B cells and they require a certain degree of flexibility, we used a flexible linker (GGGGS) to connect epitopes. In the end, the schematic diagram of the construct including epitopes and likers is shown in Figure 1.
The physico-chemical characteristics:
The molecular weight, number of amino acids, and theoretical P.I of this construct were 19.37, 175, and 7.15, respectively. The calculated aliphatic index and Grand average of hydropathicity (GRAVY) by Protparam were determined 70.23 and -0.660 respectively. The total number of negatively charged residues (Asp + Glu) and positively charged residues (Arg + Lys) are the same and equal to twenty-five.

Secondary and Tertiary structures prediction: Predicted secondary structure using online servers PSIPRED was represented in Figure 2. Results obtained from GOR IV server shows that our designed protein was composed of 52.00% alpha helix (H), 8.57% extended strand and 39.43% random coil (C) secondary structural elements. Also, 3D structure of our construct was produced by ITASSER server and is illustrated in Fig 3. C-score is a confidence score for estimating the quality of predicted models by I-TASSER. It is calculated based on the significance of threading template alignments and the convergence parameters of the structure assembly simulations. The normal range of the C-score is -5 to 2 and the higher the C-score value is, the more confident a model will be and vice-versa. The resulted C-score in the present model was 0.14 signifying a high degree of confidence. A (template modeling) TM-score less than 0.5 shows a correct topology for a model and a TM-score <0.17 indicates a random similarity. TM-Score in the model of this paper was 0.73±0.11. In addition, root-mean-square deviation (RMSD) for this model was 4.8±3.1Å.

Validation of tertiary structure and allergenicity prediction: Analysis of the results by Ramachandran plot indicated that number of residues in favored and allowed region were 207 (96.7%) and 7 (3.3%) respectively. In addition, there was no amino acid in in outlier region. The designed construct was determined as non-allergen using AllerTOP v. 2.0 server.

Solubility and Antigenicity of protein: Obtained results of SOLpro server indicated that this protein is Soluble with the probability of 0.87% and is Antigenic with the probability of 0.93%.

Protein structure analysis: Based on PSIPRED and GOR IV server results, Z-score of the protein vaccine was -2.51.

Discussion

One of the frequent ticks in the world is *Rhipicephalus* and the species of *R. annulatus* is abundant in north of Iran (16-17). This tick leads to economic losses due to sucking blood and transmission of pathogens such as *Anaplasma* and *Babesia* in cattle industry (18-19). Although the chemical control is one of the oldest and the most powerful methods of tick control (20), such method is unendurable, time-consuming, and costly; thus, attempts are toward designing vaccines based on immunogenic peptides with prediction methods based on amino acid sequence and physico-chemical characteristics (accessibility, flexibility, hydrophobicity, antigenic propensity, exposed surface, turns) (7). Biological and molecular methods are the most important control methods in recent years (21-22). Attempts have been made to produce vaccine against ticks and some were successful (23-24), but because of the diversity of antigens in species of tick, attempts are still ongoing for producing an effective vaccine (25).
Figure 1. Schematic shape of Final designed CaTro construct. E1: CT epitopes, E2: TRM epitopes, E3: CT epitopes.

Figure 2. Secondary structure of final Catro construct using online server PSIPRED.
Accordingly, novel strategies are essential to reinforce immune response in cattle and protect them against infestation of tick. Nowadays, in the production of vaccine, allergenicity and concentration of immune response are used from polytopic vaccines due to safety that can produce immune response specifically targeted against the ticks. So far several immunoinformatic researches have been accomplished related to parasitic immunogenic peptides such as TSOL18 in *Taenia solium* (26), apical membrane protein (AMA-1) in *Plasmodium vivax* (27), cathepsin D-derived epitope in *Schistosoma mansoni* (28), EG95 oncosphere protein in *Echinococcus granulosus* (29), *Echinococcus multilocularis* (30) and T- and B-cell epitopes of SAG1, GRA2, GRA7 and ROP16 in *Toxoplasma gondii* (31) and etc.

In this study, we designed a polytopic vaccine from B-cell epitopes of cathepsin and tropomyosin of *Rh. annulatus*. The epitopes were chosen by bioinformatics approaches based on high flexibility, hydrophobicity, high accessibility and strong antigenicity. So we used B-cell epitopes to produce an effective vaccine. Flexible, rigid, and cleavable linkers are three types of experimental linkers for connecting constructed fusion protein. The flexible linkers were composed of small and non-polar amino acids such as Glycine or polar amino acids such as serine or threonine. These features allow them mobility and flexibility. One of the most common flexible linkers widely used is in the form of GGGGS. In the present study, the mentioned linkers were used to connect the epitopes of B cells (32).

Various evaluations of vaccine properties revealed that our designed vaccine is a highly potential antigen and non-allergen. It should be mentioned that this predicted construct has a high aliphatic index which indicated its stability in a broad range of temperatures.

One of the significant factors in epitope mapping is the presence of proper protein secondary structures such as alpha helix and beta strand, as well as intermediate compositions such as turn and coil. The turn and coil structures have major importance due to more accessibility (33).

**Conclusion**

Since, the previous studies shown that CT and TRM are suitable immunogenic proteins as vaccine candidate (34-35), in this study, we designed the potent vaccine candidate of CT and TRM proteins of *Rh.annulatus* tick larva. Also, we decide to evaluate the protective immunity induced by this vaccine candidate construct against infestation of *Rh. annulatus* in rabbits.

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