Abstract: Leptospirosis is a wide spread bacterial zoonosis that is common worldwide. The disease symptoms are mild or acute. Leptospira has pathogenic and non-pathogenic species; it has a lot of surface antigens. Adenylate Guanylate Cyclase (AGC) is a membrane protein that is found only in pathogenic species. In this study, the complete coding sequences of AGC protein of 242 pathogen serovars were investigated by bioinformatics tools. A Pattern was selected as a target sequence based on high prevalence pathogenic serovars in Iran Antigen sites; moreover, B-cell and T-cell epitopes were predicted by IEDB web server. An antigen site amino acid (D259-R462) in complete coding sequence of AGC protein was selected. This nucleotide related sequence was cloned into the pET32a+ expression vector. Expression of recombinant protein was optimized in E. coli strain BL21-DE3 by 0.2mM IPTG after 16-hour incubation at 37°C and confirmed by 10% SDS-PAGE and western blotting. Antigenic peptide D259-R462 was highly expressed as Trx tag fusion protein. Recombinant peptide (rAcB) was purified by 6M urea from inclusion body with high extent yield 514.2 mg per 1000ml culture of E. coli. 20µg rAcB protein with montanide adjuvant was injected subcutaneously in BALB/c mice. Results showed that the recombinant peptide D259-R462 was produced...
significant antibody compared to adjuvant and PBS groups. The induced antibody in sera of immunized animal with Leptospira vaccine was detected by 250 ng of rAcB coated in ELISA microplate. This study demonstrated that antigenic region (D259-R462) of AGC protein might be useful for evaluation of antibody level in vaccinated animal.

Keywords: AGC; Leptospirosis; Recombinant protein.

INTRODUCTION

Leptospirosis is wide spread zoonosis bacterial disease caused by Leptospira spirochetes that is common all over the word [1,2]. This bacterium needs a humid environment to grow and survives in a muddy environment for months. The persistence of this bacterium in water depends on the salt concentration, pH and viscosity [3, 4]. Both of non-pathogen (bilinea serovar) and pathogenic (interrogans) species belong to the Leptospira genus [5]. The Leptospira was isolated from different hosts: mammals, birds, amphibians and reptiles [6, 7]. Human infection is caused by direct contact with the urine of contaminated animals or indirectly through the water contamination [1]. Leptospirosis is an acute febrile illness that encompasses a wide range of symptoms or without indication [8]. Symptoms can be observed from a mild form similar to influenza to severe ones such as hemorrhage, jaundice, myalgia, renal impairment, and aseptic meningitis [8,2]. In the meantime Of course, most patients exhibit moderate symptoms [1]. Also in domestic animals, leptospirosis causes, abortions, premature births, and stillbirths eventuate in world that reason economic detriment industry [1]. Leptospira has been consists of 24 sero groups and more than 300 serovars are close to 270 pathogenic [7] that can be segregated from different ecological niches and animal reservoirs. The genetic classification recently confirmed the existence of 35 species [9]. The prevalence of this disease is high among Ranchers and Smithers in Iran. According to studies in Razi Vaccine and Serum Research Institute, frequency of animal serum samples of suspected leptospirosis (icterohaemorrhagiae serovar) in two provinces Gilan and Qom were 57.1 and 20% respectively. Icterohaemorrhagiae is more prevalence serovar in Hormozgan, too, with 45.5% frequency. Autumnalis, Pomona, and Sejroe Hardjo were reported in low incidence. Frequence of human serum samples of suspected leptospirosis in Gilan province is 12.1 percent. Sejroe Hardjo serovar was dominant in Ardebil province [10].

Many virulence factors were involved in the pathogenesis and infection of Leptospira bacteria, such as hemolysin, membrane proteins, lipopolysaccharides (LPS) and other surface proteins and external molecules [11, 12]. The outer membrane proteins (OMP) of the bacterium, which are considered as antigen or binding targets for antibacterial antibodies, they act as receptors for different molecules of the host and thus an important role in the pathogenesis of these bacteria [13,14,15].

Diagnosis of leptospirosis has been confirmed by different methods such as Microscopic Agglutination Test (MAT) and Enzyme-Linked Immunosorobt Assay (ELISA). MAT is the standard technique for leptospirosis diagnosis which is carry out by incubating the patient serum with different live Leptospira serovars and recognizes by the grading of agglutination. The MAT needs to retain a collection of live strains. That is requires to alternative confirmation of each serovar in culture [16].

PCR and Real Time PCR techniques based on different genes such as 16srRNA and gyrB genes have successfully developed for the identification of pathogenic Leptospira [17].

Multi Locus Sequence Typing (MLST) was introduced to study strain taxonomy, global surveillance and in comparing evolution of the genus in Leptospira [18]. MALDI-TOF mass spectrometry was done for identification of Leptospira as a rapid and reliable method [19]. In different studies, several Leptospira antigens (rLipL32, rLipL41, rLigA, rLigB, rLipL21) were expressed in E. coli and evaluated in enzyme-linked immunosorobt assay [20, 21, 22]. Available commercial ELISA kits have a low sensitivity and specificity. On the other hand, there are several vaccines against different types of leptospira only for animals that can only reduce the risk of transmitting disease to human [23]. Therefore, finding new diagnostic methods is necessary to eliminate the limitations in standard tests. Enzyme-Linked Immunosorobt Assay (ELISA) using recombinant leptospiral antigens are sensitive techniques which have been expand as diagnostic methods for leptospiral infection [16,24,25, 26]. Adenylate Cyclase protein belongs to outer membrane lipoproteins, which are found only in pathogenic and do not exist in non-pathogenic Leptospira [35]. In this study, we attempted to introduce the rAcB as a part of AGC protein, for detection of antibody titers in immunized herds also evaluation of vaccine performance by using ELISA test.
MATERIAL AND METHODS

Bioinformatics studies

Multiple Sequence Alignment (MSA) of 242 complete coding sequences of AGC protein was done by MegAlign 5.00 DNASTAR Inc. software from 35 different serovars available in GenBank until 2.16.2019 conserved, variable regions, amino acid substitutions and other polymorphisms between different serovars were analyzed.

According to multiple sequence alignment of different serovars, one dominant pattern selected as a target sequence based on previous epidemiological studies of high prevalence pathogenic serovars (Icterohaemorrhagiae, SejroeHardjo) in Iran.

Immune informatics studies

T-cell and B-cell epitopes were predicted by Immune Epitope Database and Analysis Resource (https://www.iedb.org). Epitopes of MHC class I and II have been chosen according to HLA alleles (HLA-A*01:01, HLA-A*02:01, HLA-A*03:01 with Length 9) (DRB1*11, DRB1*13, DRB1*15, DRB1*4, DRB1*9) with consensus method and percentile rank of MHC-I <0.5 and MHC-II <1 were predicted respectively [27].

Kolaskar and Tongaonkar Antigenicity method was used for prediction. B-cell antigenic regions of AGC protein are intended to predict regions of protein that are likely to be recognized as epitopes in the context of a B cell response. Antigenic index, hydrophobicity and surface probability were analyzed. Based on mentioned criteria, one peptide (region D259-R462) selected and named AcB.

Codon optimization and DNA synthesis

Nucleotide coding sequence of antigenic region D259-R462 of Adenylate Guanylate Cyclase protein was codon optimized according to codon table usage of E. coli by GenSmart™ Codon Optimization Tool (https://www.genscript.com/gensmart-free-gene-codon-optimization.html). Two enzyme recognition sites EcoRI and XhoI were consider in 5’ and 3’ of DNA respectively. DNA synthesis was done by ShineGene Bio-Technologies, Inc., Shanghai, China.

Expression of fusion protein

Recombinant plasmid was transformed into the Escherichia coli strain BL21-DE3 and cultured in 50ml 2YT media in presence of 50µg/ml ampicillin. Induction was done when the optic density reached to 0.8. Concentration of IPTG between 0.1 to 0.5 mM and two incubation temperatures 22°C and 37°C were compared. Expression of recombinant protein (rAcB) was confirmed by SDS-PAGE and commassie blue staining.

Purification of rAcB

After induction, the cell from 50mL bacterial cell culture was harvested by centrifugation at 5000 rpm for 10 minutes. The cell pellets resuspend to 5ml PBS and sonicated 1 minute five times with one-minute interval (Hielscher Germany). Protease inhibitor PMSF in 0.1mM final concentration was added after sonication. Post sonication pellet (inclusion body) was washed five times by 1 to 5 molar Urea, pH=5.5 after 1 hour shaking at 37°C for each one to remove non-target proteins. Finally, the pellet suspended in 6 to 8 molar urea by shaking at 37°C for 5 hours. Dialysis in PBS buffer was performed for 16 hours at room temperature for the removal of urea from solubilized protein. Presence of rAcB in urea 6M, 7M and 8M were checked by SDS-PAGE and commassie blue staining.

Western Blotting

Purified protein was electrophoresed into the 10% reduced SDS-PAGE and transferred to a nitrocellulose membrane using the transfer buffer (194mM Glycine, 24mM Tris, 10% Methanol). The membrane was blocked with 2% (w/v) bovine serum albumin (Sigma, Germany) in PBST, and incubated with anti-His tag HRP conjugated antibody (1:5000) (Thermo Fisher, Germany) for 1 hour at room temperature. After three washing steps by PBST, the rAcB protein band was developed by adding 4-Chloro-1-naphthol (Sigma, Germany).
Determination of protein concentration

Concentration and yield of purified rAcB protein was achieved by Bradford method. Standard curve prepared based on BSA (bovine serum albumin) serial dilution from 0 to 10 µg on XURU’S web server regression tools (www.xuru.org) by online linear regression method.

Immunization of BALB/c Mice

Five BALB/c mice injected subcutaneously with 20 micrograms of purified rAcB formulated with adjuvant montanide (ISA 70) (SEPPIC, France) in 0.5 ml total volume three times in two weeks' interval. Four control groups PBS, Adjuvant-PBS, Adjuvant Alum, and Leptospira vaccine from Razi Vaccine and Serum Research Institute, Iran (www.rvsri.ac.ir) were considered. The blood samples were collected at before and 14, 28 and 42 days after injection.

ELISA test

96-well ELISA microplate was coated with 250 ng/well rAcB and carbonate coating buffer (pH=9.6), then microplate incubated overnight at 4°C. Also plate was washed three times with PBST. After that coated plate was washed and first, incubated for 1 hour at room temperature with immunized mice sera diluted 1:100 in PBST, pH 7.5. After another washing, 100µl Anti-mouse HRP-conjugated antibody 1:5000 in PBST was added to each well and incubated at 37°C for 1 hour then washed with PBST for five times. Then 50 µl TMB was added and incubated in dark place for 15 minutes and 2N H2SO4 was added to stop reactions. Quantification was performed using ELISA reader at wave length 450 nm.

Statistical analysis

Collected data from ELISA plate were analyzed by paired samples T-test (n=10, p-value 0.01) from five groups and box plot was designed by SPSS version 22.

RESULTS

Bioinformatics results

Twenty-two linear B-cell epitopes (score < 0.5), 21 MHCI (score < 50) and 67 MHCII (score < 1) linear T-cell epitopes which were located in two antigenic regions in complete coding sequence of AGC protein were found by using The Immune Epitope Database and Analysis Resource (www.iedb.org). An antigenic site D259 to R462 with 203 amino acids length was selected for DNA construction [Fig 1].
Expression of rAcB in E. coli BL21 (DE3)

Expression condition was optimized into Bl21 (DE3) strain of E. coli by 0.2mM IPTG after 16-hour incubation at 37°C. Expression of recombinant protein confirmed by 10% SDS-PAGE and commassie blue staining. Antigenic peptide rAcB was highly expressed as a fusion by Trx tag sequence successfully in E. coli. A 38 kDa Trx-AcB specific band was observed in bacterial lysate after induction [Fig.2]. Results demonstrated that rAcB was highly expressed as an inclusion body in non-soluble form [Fig.2].

Figure 2. Electrophoresis of cell lysates derived from induction of rAcB protein with 0.2mM IPTG in to the 10% SDS-PAGE in 10 mA for 16 hours. L: unstained protein MW marker (thermo scientific). Lanes 1 to 6 show the results of cell lysate from samples which were incubated for 0, 1, 2, 3, 4 and 16 hours at 37 C. A: post sonication cell lysates. B: post sonication Supernatant, C: post sonication pellet.

Purification and immune blotting of rAcB

High amount of recombinant AcB (514.2 mg) was purified from 1000mL culture of E. coli surprisingly the serial washing with urea 1 to 5 molar resulted in high purity protein purification. The rAcB protein was observed in urea 6 to 8 molar [Figure 3A, lane 1, 2, 3].

To verify successful production of the rAcB protein, its fusion with the His-tag was confirmed by Western blotting. The results indicated that recombinant AcB-His-tag fusion protein created a band of about 38 kDa with strong binding with the anti-His MAb. [Figure 3B, lane 2].
Figure 3. (A): SDS-PAGE of rAcB protein which was purified in gradient urea. Lanes 1: unstained protein MW marker (Thermo Fisher Scientific), 2: Purification of AcB protein with 6 molar urea, 3: Purification of AcB protein with 7 molar urea, 4: Purification of AcB protein with 8 molar urea. (B): Western blotting of purified rAcB by Anti His-Tag HRP conjugated antibody in nitrocellulose membrane. Lanes 1: prestained protein ladder (Thermo Fisher Scientific), 2: purified rAcB protein.

Yield of rAcB purification

Based on XURU’S online regression tools (www.xuru.org), with linear regression method $y=47.31697024X–0.788$ formula was identified. Concentration of purified protein was calculated by Bradford assay method. Concentrations were 12, 4.7 and 9.01mg/ml in purified recombinant protein from 6, 7 and 8 molar urea respectively. Totally, yield of purified recombinant protein was estimated 514.2 mg per each 1000ml cell culture.

Evaluation of rAcB in ELISA

Concentration of rAcB for coating of ELISA microplate was 250 ng/well. Antibody titer in blood samples of immunized mice were read in wave length 450 nm. The prominently increasing of antibody level (0.4 to 1.20) were observed in immunized mice sera with rAcB, while in control groups were 0.2 to 0.4 two weeks after last injection, [Figures 4, 1]. Mice that received rAcB significantly had higher titers compare to control groups. Results demonstrated rAcB can detect antibodies from immunized mice whit Leptospira vaccine. [Figures 4,2]
Figure 4. The diagram shows the ELISA results which was read at length at 450nm and analyzed by SPSS version 22 software.

(1): The graph indicates an increase in antibody level in blood samples of immunized mice by recombinant protein rAcB which was collected at the first day before injection to two weeks after the last injection.

(2): Comparison of antibody levels in different mice groups. AcB T4: group of immunized mice with recombinant AcB protein, ADJT4: group of injected mice with montanide adjuvant, APT4: the group of mice injected Adjuvant-PBS, PBST4: group of mice injected by PBS, VACT4: the group of immunized mice by leptospira vaccine. According to the graph, based on the rAcB it was able to have a significant response to the control groups and was able to detect the vaccine.

DISCUSSION

Leptospirosis is an acute illness and spreads rapidly [1]. Animals and human can be infected through the urine of infectious animals or contaminate soil and water [11,28]. MAT is standard test for diagnosis of Leptospirosis, nevertheless; this technique unfavorable as it will take time and need labor [1]. Keeping in view the difficult of the MAT test as a standard technique, sensitive methods such as ELISA, using recombinant leptospiiral antigens have been expand as diagnostic methods for leptospiiral infection indifferent animal species. In ELISA test several samples analyzing simultaneously and faster than MAT test. Several recombinant proteins such as rLipL32, rLipL41, LigB, produced for recombinant ELISA for sero diagnosis of bovine leptospirosis [16, 29, 26, 30, 31]. Also recombinant LigA for equine leptospirosis [32] and a Leptospira OMP (Outer membrane protein) antigen, rLipL21, rLoa22 and rLipL32 for canine leptospirosis were evaluated as antigens for ELISA [33,34]. The aim of this study was Expression of recombinant peptide based on antigenic region (D259-R462) of Adenylate Guanylate Cyclase protein of Leptospira in E. coli and Evaluation of vaccine performance also detection of antibody titers in immunized herds by using ELISA test. AGC is a very important protein in regulating cell metabolism and is the membrane lipoprotein present only in pathogenic Leptospira [35]. The AGC protein contains 758 amino acids. Antigenic site D259-R462 (AcB) according to immune informatics study was selected and to increasing yield of protein expression codon optimization was done based on E. coli codon table usage. Nucleotide sequence was inserted into pET32a (+) vector and rAcB was expressed as a fusion protein with thioredoxin (Trx). Fortunately, the rAcB protein from inclusion body was purified with high efficiency by the urea method. We found that urea purification method is a simple and low-cost with high efficiency technique. Several outer membrane proteins LipL32, rLipL21, rLipL21and LigANI of Leptospira were cloned and expressed in different hosts such as E. coli and P. pastoris. Purification yield of mentioned hosts were respectively 3 to 285 mg/L [36,37]. In this study, purification of the recombinant protein resulted in high extent yield of 514.2 mg per each 1000ml bacterial culture.

Results obtained from ELISA test showed that recombinant protein rAcB was able to produce high level of antibody in mice which were immunized (recombinant AcB protein receiving group) and was suitable immunogenic. Amazingly analyzing data from ELISA demonstrated that rAcB which was coated to ELISA
plate could be detected *Leptospira* antibody from immunized herds group with *Leptospira* vaccine. This is the first study to introduce AGC protein of *Leptospira* as a candidate antigen in ELISA to explore the feasibility of using them for detection of specific *Leptospira* antibodies in immunized animal.

**CONCLUSION**

The recombinant peptide AcB (D259-R462) which was produced in the *E. coli* has the ability to detect antibodies produced in vaccine-immunized animals, so this protein can be used to evaluate antibody levels in clinical trial of under developing vaccine and vaccination program.

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