Identification and Characterization of a Gene Encoding 18.5kDa Protein from *Mycobacterium avium* Sub species *paratuberculosis*

Dheeraj Pal* and P.P. Goswami

*Gene Expression Laboratory, Indian Veterinary Research Institute, Izatnagar, Bareilly, Uttar Pradesh - 243122, India.

(Received: 21 August 2015; accepted: 13 October 2015)

In this study the effort was assumed to clone and express coding sequences of *M.a.paratuberculosis* to study their immune reactivity. Primers were designed for ORFs retrieved from MAP complete genome strain k10 (locus tag MAP 0862 and MAP 1087). The PCR amplified product of each gene fragment was cloned into *E. coli* expression vector pQE-30 and the resultant constructs were designated as pQE 501. The positive recombinant clones on induction with IPTG expressed the protein bands corresponding to 18.5kDa protein on SDS PAGE. The His-18.5 protein was purified using single step Ni-NTA chromatography. The yield of the purified His-18.5 protein was about 15 mg/L and from induced *E. coli* cultures harbouring plasmid pQE-501. Antigenicity of these proteins were evaluated by western blot using sera from a small number of cattle infected with MAP. The immuno proteomic analysis of culture filtrate (CF) and cellular extract (CE) of MAP revealed that serological tests may be improved by the use of MAP proteins derived from culture filtrates and not from cellular extracts. Development of sensitive serological tests for the rapid identification of infected animals at subclinical stage requires expression and characterization of proteins or secreted early from post infection MAP. Polyclonal anti sera raised against purified His-18.5 protein reacted with induced *E. coli* whole cell lysate harbouring pQE 501 and also with purified recombinant 18.5kDa protein on western blot. The recombinant His-18.5 protein was recognized by rabbit hyper immune sera of the MAP culture filtrates and also by serum from a goat with clinical paratuberculosis.

Keywords: DNA-Star, serological, *Mycobacterium avium*, MAP.

Paratuberculosis (Johne’s disease), chronic granulomatous enteritis of domestic ruminants1 and wild animals2,3 caused by acid-fast, slow growing, fastidious *Mycobacterium avium* subspecies *paratuberculosis* (MAP). It is an enzootic disease on the B list of the Office des International Epizootes (‘OIE’ and causes huge economic losses to the dairy industry worldwide.

Detection of MAP, especially during the often-lengthy subclinical phase of the disease, remains difficult due to intermittent shedding of small numbers of bacteria and a lack of effective diagnostic reagents4. The fecal culture test requires 12-18.5 weeks for cultivation and its sensitivity level is estimated to be approximately 38%5. So there is a need to improve the sensitivity or specificity of currently used diagnostic tests. Entry of microorganisms in payer’s patches and infection of macrophages at this site allows presentation of antigens to immune cells and usually activates defense site allows presentation of antigens to immune cells usually activates defense mechanism. In paratuberculosis, as in other mycobacteria, Cell mediated immunity (CMI) is considered to be the principal mechanism for establishment or clearing
of infection. Antibody is assigned are often seen in advanced clinical cases. In mycobacterial infections, CMI, which is highly initially, is associated with higher TH1 responses leading to IFN-γ production.

Development of sensitive serological tests for the rapid identification of infected animals at subclinical stage requires expression and characterization of early secreted proteins of MAP as important antigens for the diagnosis of paratuberculosis6,7. Literature mining indicates the present work as the first of its kind involving epitopic strain from MAP 0862 from the MAP genome strain k10 encoding immune reactive 18.5 kDa protein.

Currently used diagnostic tests for Johne’s disease need to be improved due to deficiencies in their sensitivity or specificity. Problems of specificity are due to the high degree of similarity that exists between MAP and environmental mycobacteria, especially the closely related MAA. It is not surprising therefore that cell mediated immune assays such as the intradermal (skin) test, which uses a complex, undefined MAP secreted protein preparation to stimulate an immune reaction in the host, are prone to false-positive results due to cross-reactivity with similar proteins present in other mycobacteria.

MATERIALS AND METHODS

Materials and reagents

All the chemicals and biological used in the present study were of molecular biology grade. Agarose, Ammonium persulphate, Acryl amide, Bovine serum albumin, Bisacryl amide, â-mercapto- Life technology USA ethanol, Bromophenol blue, Calcium Chloride, Coomassie brilliant blue 250, Magnesium Chloride, Sodium Chloride, Potassium Acetate, Tween-20, Isopropyl thiogalactose-pyranoside (IPTG), X-gal, lysozyme, Sodium dodecyl sulphate, Diaminobenzidine, TEMED, Glycine, Ethidium Bromide, Proteinase K, RNase, Triton X-100, PMSF, Trypan blue, Fetal calf serum , GMEM, Goat anti-rabbit IgG HRP conjugate, Rabbit anti-goat IgG HRP conjugate, Dialysis tube and Antibiotics (ampicillin and kanamycin) were obtained from, Sigma Chemicals, USA. Potassium chloride, Disodium hydrogen orthophosphate, Potassium dihydrogen orthophosphate, Hydrogen peroxide, Certrimide, Agar, CTAB, Glucose and Glycerol, Sodium nitrite and Orthophosphoric acid, Sodium carbonate, Sodium bicarbonate, Isopropanol, Phenol, Chloroform, Ethylene diamine tetra-acetic acid, Boric acid, Glacial acetic acid, Isoamyl alcohol, Sodium hydroxide, Sodium acetate, Sodium dihydrogen phosphate, Magnesium sulphate were purchased from Qualigens & Merck, India respectively. QIAGEN, Germany supplied QIAEX II Gel Extraction Kit while DiCo, USA supplied Luria-Bertani Media, SOB Media and Middlebrook 7H10 agar and OADC salt. Restriction endo-nucleases and T4 DNA ligase were procured from New England Biolabs, UK and MBI Fermentas, Germany respectively. All other reagents were acquired from commercial sources.

Oligonucleotide primers

A set of specific oligonucleotide primers 862 F and 862 R from (locus tag MAP 0862 region884392-884892), were synthesized by integrated DNA technologies, inc.Coralville, IA USA, basedon sequence information of MAP strain k10 complete genome GeneBankAccession No:AE016958 Linkers with restriction endo-nuclease sites BamH1 at 5’ end and Pst1 at 3’ end were included in the forward and reverse primers respectively.

MAP 862 (F) 5’- TAC GGATC C ATG CGT CGT GGC ACT GTG GT 3’- 29 Mer
MAP 862 (R) 5’-TAC CTG CAG TCA GCA TCT GTA AAC CCC AG-3’- 29mer

Mycobacterial strain and antigen

Bacterial strain of M. a. paratuberculosis (MAP) strain 316F were obtained from Biological Products Division of IVRI, Izzatnagar and later maintained at Gene Expression lab, Division of Animal Biotechnology. IVRI, Izzatnagar. Antigen against MAP culture filtrate and clinical sera from the goat affected with paratuberculosis were available in the lab.

Plasmid and host strains

Prokaryotic expression vector pQE-30 was purchased from QIAGEN, Germany.

Laboratory animals

New Zealand white rabbits were obtained from Laboratory Animal Resource Section, IVRI, Izzatnagar. Standard prescribed guidelines for care and use of laboratory animals were followed during the experimentation with these animals.
METHODS

Culture and growth of *M. a. paratuberculosis* and *E. coli*

MAP organisms were grown on Middlebrook 7H10 agar enriched with 0.1% glycerol v/v and 10% oleic acid dextrose catalase (OADC) with additional supplementation of Mycobactin J (2mg/l) while *E. coli* cells were grown in Luria Bertani (LB) medium with shaking at 180 rpm. *E. coli* M15 cells containing pQE-30 vector were grown in presence of kanamycin @ 50 mg/ml and ampicillin @ 50 mg/ml. Both organisms were grown at 37 ºC.

Isolation of genomic DNA from *M. a. paratuberculosis*

The genomic DNA from MAP was isolated by the method of Portillo *et al.* (1991) with slight modifications. The bacterial colonies scrapped from two months old Middlebrook 7H10 agar slants were washed thrice with 1X TE and re-suspended in 500 ml of 1X TE. Lysozyme was added to the final concentration of 5 mg/ml. After incubation at 37°C for two hrs, SDS and proteinase K were added @1% and 250 μg/ml respectively) and incubated further at 65ºC for 30 min. To this, added 80 μl of 5M NaCl followed by addition of 64 μl of CTAB/NaCl solution and vortexed. The suspension was again incubated at 65°C for 30 min. To this, added 80 μl of 5M NaCl followed by addition of 64 μl of CTAB/NaCl solution and vortexed. The aqueous phase containing DNA was pelleted by centrifugation and washed with 80% ethanol, dried and re-dissolved in 200 μl of 1X TE. And ethanol precipitation. Contaminating RNA was removed from DNA by incubating with 100 μg/ml RNase. The treatment was given for one hour at 37°C, followed by phenol: chloroform and ethanol precipitation. The DNA was quantitatively estimated using the following formula, by recording the absorbance at 260 nm wave length.

\[
\text{Concentration of dsDNA (µg/ml) = A260 X 50 X dilution factor}
\]

Agarose gel electrophoresis

The DNA preparations were analyzed on 1% agarose gel and visualized under UV transillumination and documented by photography in Uvitech Gel Documentation System (Austria).

Polymerase chain reaction

The PCR was carried out in 25 ll reaction volume using 1 ll of genomic DNA (10 ng) as template, 2.5 ll of PCR buffer, 1 ll of MgCl2 (1.5 mM), 1 ll (25 iM) of each primers, 1 ll of dNTP mix (200 iM of each dNTP) and 1 U of Taq DNA polymerase. The volume was made up to 25 ll by adding DNase free water. The thermal cycling steps were carried out in PTC-200 thermo-cycler MJ Research Inc, USA with initial denaturation at 94°C for 5 min followed by 30 cycles with denaturation at 94°C for 1 min, annealing of 1 min at 58°C for 501bp followed by extension at 72°C for 30 Seconds and final extension at 72°C for 1 min. Size of the amplified product was confirmed by using DNA molecular weight marker in a 1.2% agarose gel and quantified by spectrophotometric analysis. The DNA fragment of 501 bp was eluted from agarose gel using QIAEXII gel extraction kit.

Cloning of 501bp in prokaryotic (*E. coli*) expression vector pQE-30

Ligation

Plasmid pQE 30 was isolated from *E. coli* cells by small scale alkaline lysis method followed by the ligation of eluted PCR product of 501 bp as follows:

a. pQE-30 : 1 μl (50ng)
b.501bp PCR product: 1 μl (50ng)
c. 2X ligation master mix : 5 μl
d. nuclease free water: 3 μl

The 10 μl ligation mixture was kept at 16°C overnight and the resulting plasmid was designated as pQE 501 which was stored at -20°C.

Transformation and Screening of recombinant pQE 501

10 μl ligation mixture was diluted to 200 μl TCM (1X), to which, 200 μl of the competent cells were added with gentle mixing and left at 0 ºC on ice for 1hr. Heat shock was given to this mixture for 2min at 45°C and then was rapidly chilled on ice for 10 min. Then 600 μl of SOC medium was added and incubated at 37°C for 1hr with shaking. The transformed cells were spread on LB agar plate containing ampicillin (100 μg/ ml) and kanamycin (25 μg/ ml). Plates were incubated at 37°C overnight and later stored at 4°C.

Screening of recombinant pQE501

About twenty randomly selected colonies were picked up containing pQE 501 clones and
grown on LB broth containing Ampicillin and Kanamycin and incubated at 37°C overnight in a shaker incubator at 180 rpm. Plasmid DNA was extracted & positive clones were identified by Colony PCR and restriction endonuclease analysis.

Expression and purification of the recombinant 18.5 kDa (His 18.5) Protein

Fresh *E. coli* culture harbouring pQE (OD600 of 0.6 was induced with 1mM IPTG to express the recombinant 18.5 protein and grown again for 4 hr at 37°C in a shaker incubator at 180 rpm. The *E. coli* cells were pelleted by centrifugation at 6,000 rpm for 10 min, & frozen at −20°C overnight followed by thawing at room temperature for 15 min. These pellets were resuspended in 1/25th volume of lysis buffer (buffer B-8M urea, 0.1M NaH₂PO₄, 1M TrisCl - pH 8.0) and incubated with agitation for 1 hr at RT. Removed the cell debris by centrifugation at 10,000 x g for 20-30 min at RT (20-25°C) and about 600ìl of the cleared lysate supernatant containing the 6X His-tagged protein was loaded onto a pre-equilibrated (with 600 ìl of buffer B) Ni-NTA spin column and centrifuged for 2 min at 700 x g. The column Ni-
resin bound 6X His-tagged proteins was washed twice with 600 il of wash buffer (buffer C - 8M urea, 0.1M NaH2PO4, 1M TrisCl-pH 6.3) and eluted twice with 200 µl of elution buffer (Buffer E - 8M urea, 0.1M NaH2PO4, 1M TrisCl- pH 4.5).

**Sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE)**

SDS-PAGE was carried out on a vertical slab mini apparatus (Atto, Japan). After electrophoresis the gel was stained with Coomassie brilliant blue G250 and distained in solution containing methanol and acetic acid. The fraction containing the purified recombinant 18.5kDa protein was resolved and approximate molecular weight of the protein was determined by comparison with protein molecular weight marker in Uvitec gel documentation system using Uvipro software.

**Dialysis of the recombinant 18.5kDa**

The selected fractions of each protein were pooled and extensively dialyzed at 4ºC against PBS (pH 7.4) in dialysis tubing with 8 kDa molecular weight cut-off, and stored in aliquots at −20ºC until used.

**Hyperimmunization for raising anti sera**

New Zealand white 8-10 weeks old rabbits
were immunized to raise antibody against the purified His-18.5 by inoculating with 150 µg of the immunogen in 0.5 ml PBS (pH 7.4) along with equal volumes of incomplete Freund’s adjuvant (IFA) subcutaneously. After three weeks, subsequent boosters of 100 µg immunogens were given intramuscularly at weekly intervals. The rabbits were bled a week after third booster and sera were separated and stored at −20 °C in aliquots.

**Sequence analysis**

DNA sequencing of the plasmid pQE 501 was carried out commercially.

**RESULTS**

**Expression of Recombination 18.5kDa Proteins in E. coli**

PCR amplified 501bp gene fragments each was each digested with BamHI/PstI restriction enzymes and inserted in-frame (Fig:1). The nucleotide and protein sequence analysis of plasmid pQE 501 containing 501bp gene fragments of MAP 316 F strain revealed about 100% homology with retrieved ORFs of the MAP k10 strain by laser gene software DNA star. The predicted primary protein composed of 166 amino acids with a mature protein of 18.5kDa respectively. Analysis of the deduced amino acids sequence of 18.5kDa protein according to Kyte and Doolittle (1982) algorithm showed presence of three and two highly hydrophobic region (Fig:2).

**DNA sequence and deduced amino acid sequence analysis**

The nucleotide and protein sequence analysis of plasmid PQE 501 containing 501 bp gene fragments of MAP 318.5F strain revealed about 100% homology with retrieved ORFs of the MAP k10 strain by laser gene software DNA star. The predicted primary protein composed of 166 amino acids with a mature protein of 18.5kDa. Analysis of the deduced amino acids sequence of MAP18.5kDa according to Kyte and Doolittle (1982) algorithm showed presence of three hydrophobic regions.

**DISCUSSIONS**

Johne’s disease with a potential of causing economic losses worldwide could not be eradicated successfully due to the lack of simple and specific diagnostic tests for the early detection prior to the appearance of disease signs’ such as fecal shedding of MAP in the environment. Early diagnosis is important to identify and remove potential fecal shedders of MAP to prevent the spread of John’s disease, and requires the development of sensitive and specific diagnostic tests. Indeed fecal culture the current gold standard tests, which can detect both clinical and subclinical stages is time consuming, requires more than 15 weeks and also labour intensive. Poor specificity of the intradermal skin test and interferon gamma (IFN-γ) release assay, agarose gel immunoblotting (AGID), complement fixation (CF) and ELISA use a complex ill-defined mixtures of proteins derived from either, whole cell or fractionated extracts of MAP. To date a number of antigens have also been identified but the data about the characterization of proteins secreted early post-infection from MAP are still lacking. Recently Bannatine et al. constructed the partial protein array and examined the antibody profiles of subclinically MAP infected animals and study has focused on the detecting antigens eliciting early antibody response. Since, proteins produced by the MAP are in very low concentration, purification becomes highly cumbersome and uneconomical. Therefore, for high level expression of proteins, E. coli has been widely used for the expression of large number of genes at level sufficient for structural and biochemical analysis and even for product development.

The recombinant proteins will further be useful in the development of diagnostic reagents for subclinical detection paratuberculosis in animals. While further studies using sera from experimentally as well as naturally infected animals are needed to lay down the solid foundation towards the development of feasible ELISA based assay for subclinical detection of MAP infected animals. Although current serological tests are useful in detecting animals with clinical paratuberculosis, the application of this procedure in identifying animals in early stages of infection or in subclinical stages has proven to be of limited value. Also the sensitivity of the commercial ELISA might at least in part be due to heterologous nature of the antigen they are apparently based on derived from M. avium sub sp.

Therefore, in the present work keeping in
view the importance of identifying MAP specific antigens through the cloning and heterologous expression of 501 BP coding sequences found in the MAP genome encoding 39.7kDa proteins was chosen for expression and immuno-reactivity. Based on the sequence information of two genes encoding 18.5 kDa from MAP strain k10, complete genome Gene Bank Accession No. AE016958 (tag MAP 0862 and MAP 1087 coding for hypothetical proteins) and also the information about multiple cloning sites of the expression vector pQE-30, restriction sites BamHI and PstI were incorporated into the oligonucleotide primers to facilitate directional cloning. The resulting plasmid pQE501 containing an open reading frame encoding successively 6X histidine polypeptide, 18.5kDa. The recombinant pQE 501 clones were confirmed by release of the insert by double digestion with BamHI and PstI restriction enzymes and on induction with IPTG for 4 hours appearance of 18.5 kDa protein band. Expression of the recombinant proteins was induced by IPTG which bind to the lac repressor protein, inactivating it leading to transcription of sequences downstream of the promoter. Using pQE-30 vector, Goswami et al.14 and Basagoudanavar et al.15 have got maximum expression of the recombinant protein after 4-6 hrs post induction. Therefore, it was decided to purify the recombinant proteins 4 hours post induction from E. coli culture. The presence of 6X histidine tag of 840 Dalton at the N-terminal of the recombinant proteins facilitates single step affinity purification, which is poorly immunogenic and may not interfere with the protein immunogenicity its functional structure; hence the tag was not removed by protease cleavage. The polyclonal antisera raised in rabbit against the recombinant His-18.5proteins strongly reacted with the E. coli expressed recombinant 18.5 kDa and 18.5 kDa proteins on immunoblot.

To date a number of antigens have also been identified but the data about the characterization of proteins secreted early post infections from MAP are still lacking. The partial protein array and examined the antibody profiles of subclinically MAP infected animals and study has focused on the detecting antigens eliciting early antibody response. For functional studies and diagnostic purpose large amount of pure protein is required. Since, proteins produced by the MAP are in very low concentration, purification becomes highly cumbersome and uneconomical. Expression of the recombinant proteins was induced by IPTG which bind to the lac repressor protein, inactivating it leading to transcription of sequences down stream of the promoter.

This has not only confirmed the heterologous expression but also suggested that the recombinant proteins retained their antigenicity. Reactivity of the His-18.5 proteins with polyclonal sera against culture filtrate of MAP on Dot ELISA revealed the secretory nature of the recombinant proteins. Further reaction of these recombinant proteins with clinical sera from par tuberculosis affected goat suggested that these proteins have retained their antigen city even after purification. The result of these studies suggested that these recombinant proteins will further be useful in the development of diagnostic reagents for subclinical detection par tuberculosis in animals. While further studies using sera from experimentally as well as naturally infected animals are needed to lay down the solid foundation towards the development of feasible ELISA based assay for subclinical detection of MAP infected animals.

ACKNOWLEDGMENT

The authors are thankful to the Director, Indian Veterinary Research Institute (IVRI) for providing the facilities to carry out the work.

REFERENCES


