Assessment of recombinant plasmid expressing fusion antigen Ag85B-Rv3425 in management of acute tuberculosis infection in mice

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Abstract. The emergence of drug-resistant tuberculosis (TB) and HIV-TB co-infection fuels an urgent need to develop novel therapeutic approaches, including therapeutic vaccines. Therapeutic vaccines have been proven to be a good strategy by inducing antigen specific immune responses against TB infection. In the present study, a recombinant plasmid based on lentiviral vector expressing fusion antigen Ag85B-Rv3425 (A3), and was constructed the immunogenicity and treatment effects in TB mice were assessed. The results showed that A3 delivered by the plasmid could be expressed appropriately in vivo and induced higher production of tumor necrosis factor-α and interleukin-2 compared with A3 recombinant protein in mice. Moreover, the recombinant plasmid expressing A3 conferred resistance to acute TB infection in mice, characterized by a reduction in the bacterial load in the lungs and spleen, as well as attenuated TB lesions in lung tissues. These results implicated that the recombinant plasmid based on lentiviral vector expressing A3 is a potent and promising therapeutic agent to treat acute TB infection.

Introduction

Tuberculosis (TB), a worldwide infectious disease caused by Mycobacterium tuberculosis (Mtb), continues to be a major threat to human health (1,2). Unfortunately, BCG (bacillus Calmette-Guérin), the only vaccine widely used against TB, provides varied protective efficacy (0‑80% in randomized control trials) (3-5). Recently, the emergence of drug-resistant Mtb strains as well as HIV-TB co-infection poses numerous complexities for TB control (6,7). More effective vaccines and more potent immune strategies against TB are now in urgent need.

Therapeutic vaccines shed light on TB control by inducing antigen specific immune responses against Mtb in vivo (8), and Mtb potent antigen encoding genes are delivered into host cells via plasmid, adenovirus or lentivirus for in vivo gene expression (9,10). As a novel immunotherapy strategy, therapeutic vaccine has been successful for TB control not only in latent TB infection model but also in acute infection model (11,12).

Ag85B-Rv3425 (A3) is a fusion protein of Ag85B and Rv3425, a member of the PE (Pro-Glu) and PPE (Pro-Pro-Glu) family, located in RD11 region which is absent in BCG strains (13,14). It has been reported rBCG::Ag85B-Rv3425 vaccine, formed by co-expressing Rv3425 and Ag85B in BCG, provides better protective efficacy against Mtb challenge compared with BCG (15). Recently, it was reported that A3 delivered into mice via lentivirus with one single dose administration confers post-infection resistance to acute TB infection (11).

In this study, we constructed a recombinant plasmid based on lentiviral vector expressing fusion antigen Ag85B-Rv3425 (A3) and assessed its immunogenicity and treatment effect in TB mice. We found that in vivo expression of A3 could induce more IL-2 as well as TNF-α compared with immunization using A3 purified from E. coli, which is recombinant fusion

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protein of Ag85B-Rv3425 purified from recombinant E. coli. Moreover, in vivo expression of A3 provides immunity to acute Mtb infection characterized by reduced Mtb burden in lung and spleen and attenuated pathology in lung tissue.

**Materials and methods**

**Animals.** A total of 6-8 week-old female C57BL/6 mice were purchased from SLAC Laboratory Animal Center (Shanghai, China). All mice were maintained under specific-pathogen-free (SPF) conditions in animal facilities at Animal Biosafety Level (ABSL)-III lab of Wuhan University and given sterile water, mouse chow and bedding. All mice experiments were performed in accordance with recommendations in the Wuhan University Research Council Guide for Care and Use of Laboratory Animals. Animal study protocols were also reviewed and approved by the Wuhan University Institutional Animal Care and Use Committee.

**Plasmid and proteins.** pLenti 6.3 plasmid was purchased from Biomiga (San Diego, CA, USA). The cloning of ag85b-rv3425 fusion gene into pLenti6.3 plasmid was our described previously (11). A3 protein were purified as our described previously (13). Endotoxins concentration of A3 protein was tested using the commercially available Quantitative Chromogenic End-point Tachypleus Amebocyte Lysate reactivity endotoxin kit (Chinese Horseshoe Crab Reagent Manufactory, Xiamen, China). A total of <5.0 EU/mg endotoxins was observed and purified A3 protein was subsequently subjected to animal immunization.

**Immunization.** C57BL/6 mice were divided into four groups (20 mice per group). At week 0, mice in PBS group received a subcutaneous injection of empty plasmids in PBS and mice in A3-Pro group were injected subcutaneously with 50 µg recombinant A3 protein in 250 µg dimethyl dioctadecyl-ammonium bromide (DDA) adjuvant (Sigma). Mice in A3-Vec group were immunized by an intramuscular injection of 50 µg recombinant plLenti6.3 plasmid harboring ag85b-rv3425 fusion gene. Mice in Vec group were immunized by an intramuscular injection of 50 µg plLenti6.3 empty plasmid vector. All mice received 3 injections at a two-week interval. 4 weeks after the last injection, mice were sacrificed for immune response evaluation (8 mice per group).

**Separation of spleen lymphocytes.** Spleen from each mouse was collected under aseptic condition immediately after sacrifice. The single-cell suspension (4 ml) was obtained by gently grounding spleen through a 75 µm cell strainer and under-layed by 4 ml pre-warmed Lymphocyte-M (Cedar Lane Lab, Burlington, VT, USA). Density-gradient centrifugation was performed at 1,250 x g for 25 min to isolate splenocytes (16,17). Cells were counted and diluted to a final concentration of 5x10⁶ cells/ml in RPMI-1640 medium (Invitrogen; Thermo Fisher Scientific, Inc., Waltham, MA, USA) supplemented with 10 U/ml penicillin, 10 µg/ml streptomycin and 10% fetal bovine serum (Invitrogen; Thermo Fisher Scientific, Inc.).

**Enzyme-linked ImmunoSpot (ELISPOT) assay.** Filtration plates (96-well; Merck KGaA, Darmstadt, Germany) were coated with monoclonal antibodies against mouse TNF-α (U-CyTech, Utrecht, The Netherlands) for 16 h at 4°C. After being washed with phosphate-buffered saline (PBS), wells were blocked with 200 µl blocking buffer R for 1 h at 37°C. Spleen lymphocytes (2.5x10⁵ cells) were mixed with 10 µg/ml A3 protein and added to each well. Stimulation was completed by incubating cells at 37°C for 36 h. After being washed with PBST for five times, each well was filled with 100 µl of biotinylated detection antibodies, and incubated at 37°C for 1 h. Then, wells were washed five times and incubated with 100 µl of streptavidin-HRP for 1 h at 37°C. After being washed, 100 µl of AEC substrate solution was added to each well and plates were incubated for 25 min at room temperature in the dark. Color development was stopped by rinsing both sides of the polyvinylidene fluoride membrane in 96-well filtration plates with demineralized water. Finally, the plates were air dried and the amount of positive spots was counted by a dissecting microscope. The number of TNF-α expressing cell was calculated by spot forming units per million cells.

**Cytokines assay.** 2.5x10⁶ cells were stimulated with 10 µg/ml A3 fusion protein at 37°C, 5% CO₂ and high humidity for 36 h as described before (18). The supernatant from stimulated spleen lymphocytes was collected, and the concentrations of TNF-α and IL-2 were measured by BD™ Cytometric Bead Array kit (BD Biosciences, Franklin Lakes, NJ, USA) according to the manual in the kit.

**Mtb infection and immunotherapy.** For immunotherapy, mice were infected intravenously via lateral caudal vein with 6.8x10⁵ live Mtb strain H37Rv. At 4 weeks after infection, immunotherapy was applied and injection strategy was the same as immune responses evaluation listed above. At 4 weeks later at week 12, mice were sacrificed for colony-forming unit (CFU) counts (6 mice per group) and histopathological analysis in lung and spleen (6 mice per group).

**Histopathology analysis.** Each lung was excised and fixed in 4% neutral-buffered paraformaldehyde solution for 24 h. Then lung tissue was embedded with paraffin. Series of sections with a thickness of 4-7 µm were then cut and stained with haematoxylin and eosin under standard methods. Double blind analysis was then made by board-certified pathologists and more than 10 slides of each lung were evaluated.

**Statistical analysis.** Statistical analysis was performed using SPSS Statistics 17.0 for Windows software package. Results were subjected to one-way Anova test followed by multi-comparison testing. P<0.05 was considered to indicate a statistically significant difference.

**Results**

**Fusion antigen A3 delivered by plasmid pLenti6.3 could be expressed in mice and induce immune response.** To make fusion Mtb antigen A3 expressed appropriately in eukaryotic cells, we constructed a recombinant plasmid expressing A3 (A3-Vec) by cloning ag85b-rv3425 fusion gene fragment
and inserting it into pLenti6.3 vector, whose CMV promoter upstream could drive the transcription of ag85b-rv3425 fusion gene. Then, 50 µg endotoxin-free recombinant plasmids (A3-Vec) were injected into mice for 3 times at a two-week interval. As controls, 50 µg endotoxin-free empty plasmids (Vec) and 50 µg recombinant proteins A3 (A3-Pro) purified from E. coli were also injected for 3 times at a two-week interval, respectively. In the 4th week after the last immunization, mice were sacrificed for immune response analysis. We have detected A3 fusion protein specific antibody IgG response in A3-Vec group and found that injection of recombinant plasmids A3-Vec, as well as recombinant proteins A3, induced A3 specific antibody Immunoglobulin G (IgG) production (Fig. 1). It indicates that Mtb fusion gene ag85b-rv3425 delivered by pLenti6.3 plasmid could be expressed and translated into fusion protein in mice and then recognized by immune system.

A3 delivered by pLenti6.3 plasmid increased the production of Th1-type cytokines. Next, we want to know whether A3 delivered by plasmid could induce protective immune response since it could be expressed in mice. As Th1-type immune responses plays important role in protection against Mtb (18,19). To identify whether A3-Vec plasmid could induce Mtb-specific Th1-type immune responses, we evaluated the production of Th1-type cytokines in mice. ELISPOT results show that there are more TNF-α producing cell in mice of A3-Vec group than those in A3-Pro group, Vec (empty plasmids) group and PBS group (Fig. 2A). And we found that the concentration of TNF-α in culture supernatant of spleen lymphocytes from A3-Vec immunized mice was higher than that from other groups determined with enzyme linked immunosorbent assay (ELISA) methods with recombinant A3 protein stimulation in vitro (Fig. 2B).
Moreover, we detected a much higher concentration of IL-2 production in spleen lymphocytes from A3-Vec group upon A3 protein stimulation (Fig. 2C). It seems that A3-Vec immunization exhibited higher ability in inducing Th1-type cytokines production than immunization of A3-proteins or empty plasmids.

We also compared the percentages/numbers of T cell subsets (CD3+, CD4+ and CD8+ T cells subsets) in spleen lymphocytes among these groups, and found no significant difference (data not shown).

Administration of A3-Vec plasmids effectively reduces Mtb burdens in lungs and spleens of Mtb infected mice. Since A3-Vec could induce antigen specific Th1-type cytokines production, we want to know whether A3-Vec could relieve symptoms of TB in mice. To address this, we evaluated the therapeutic effect of A3-Vec injection in Mtb-infected murine model. Mice were challenged by intravenous injection with Mtb H37Rv to establish acute infection. After 4 weeks of infection, we began to inject A3-Vec plasmids, A3-Pro, Vec (empty plasmids) and PBS control into TB mice according to the schedules listed in Table I. At the 12th week, mice were sacrificed, and bacteria burdens in lungs and spleens were counted. We found that there was a statistically significant CFU burden reduction in lung, as well as spleen, of A3-Vec

Table I. Schedule of therapy experiments against Mtb challenge.

<table>
<thead>
<tr>
<th>Group</th>
<th>0 week</th>
<th>4th week</th>
<th>6th week</th>
<th>8th week</th>
<th>12th week</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBS</td>
<td>Infection</td>
<td>PBS</td>
<td>PBS</td>
<td>PBS</td>
<td>Necropsy</td>
</tr>
<tr>
<td>A3-Pro</td>
<td>Infection</td>
<td>A3-Pro</td>
<td>A3-Pro</td>
<td>A3-Pro</td>
<td>Necropsy</td>
</tr>
<tr>
<td>A3-Vec</td>
<td>Infection</td>
<td>A3-Vec</td>
<td>A3-Vec</td>
<td>A3-Vec</td>
<td>Necropsy</td>
</tr>
<tr>
<td>Vec</td>
<td>Infection</td>
<td>Vec</td>
<td>Vec</td>
<td>Vec</td>
<td>Necropsy</td>
</tr>
</tbody>
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Pro, protein; Vec, vector; Mtb, Mycobacterium tuberculosis; PBS, phosphate-buffered saline.
treated mice comparing with Vec and PBS controls or even A3-Pro treated mice (Fig. 3A). And mice in A3-Vec present the lowest CFU counts in spleen (Fig. 3B). Acid-fast staining results showed that many Mtb bacteria aggregating were found in lung tissue from control groups (PBS and empty plasmid-Vec) as well as from A3-Pro treated group, while fewer Mtb bacteria were seen in slide from A3-Vec group (Fig. 3C). These results demonstrate that A3-Vec administration could provide immune protection to acute Mtb infection through inhibiting Mtb growth in vivo.

Administration of A3-Vec plasmids confers immune resistance to TB lesions. To identify whether administration of A3-Vec plasmid could decrease TB lesions in lungs caused by Mtb infection, we performed histopathological analysis to compare gross pathology in lungs of mice in different groups. TB lesions area percentage was calculated by double blind pathologists. We found that A3-Vec group exhibits the least lesion area among these four groups (Fig. 4). More detailed histopathological images illustrated mild lesion with small area of epithelioid cells and lymphocytes hyperplasia in A3-Vec group, while there were large area of lymphocytes, eosinophils and macrophages hyperplasia in pulmonary alveoli of the other three groups (data not shown). It means that administration of A3-Vec in mice conferred immune resistance to lesions caused by TB acute infection.

Discussion

The emergence of drug-resistant TB strains and HIV-TB co-infection makes global TB control a real challenge, it’s in urgent need to facilitate worldwide control of TB (20). Immunotherapy is regarded as a potential approach to eliminate Mtb (21). In this study, we constructed a recombinant plasmid based on lentiviral vector expressing multiple antigens and assessed its immune response and treatment effect in TB mice.

As our previously reported, fusion protein Ag85B-Rv3425 is proved to be an effective multiple epitopes antigen (11,13). In our study, we found that recombinant plasmid A3-Vec, like lentivirus delivered A3 (11), could induce A3 fusion protein specific antibody IgG response, indicating that it could be expressed in mice and then recognized by immune system. Moreover, we have observed immunization of plasmid A3-Vec could induce high Th1-type cytokines production, such as IL-2 and TNF-α, than immunization of A3-Pro recombinant protein, though A3-Pro can induce high IgG in mice. And A3-Vec can decrease the CFU counts in lung as well as in spleen and alleviate tuberculous lesions in acute Mtb infected mice comparing to A3-Pro. It implies that plasmid vector might be a more effective method in antigen presentation than protein subunit vaccine.

In summary, our results show a good therapeutic effect of recombinant plasmid A3-Vec treatment against acute TB infection in mice. Further study might be conducted in non-human primates or even in humans to assess therapeutic effect of this recombinant plasmid.

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