Immune-potentiation of Pneumococcal Capsular Polysaccharide Antigen using Albumin Microparticles

Bernadette D’Souza1, Prathap Nagaraja Shastri2*, Gabrielle Hammons3, Ellie Kim3, Lakshmi Prasanna Kolluru2, George M Carlone3, Gowrisankar Rajam2 and Martin J D’Souza2

1McWhorter School of Pharmacy, Samford University, Birmingham, AL 35229, USA
2College of Pharmacy and Health Sciences, Nanotechnology Laboratory, Mercer University, Atlanta, GA 30341, USA
3Division of Bacterial Diseases, Centers for Disease Control and Prevention, Atlanta, GA 30333, USA

Abstract

Microparticles (MPs) offer several advantages as unique vaccine delivery system, including the ease to manufacture, targeted delivery of antigen payload, sustained antigen release and possible role as an immune-adjuvant. In this study, we evaluated albumin matrix for pneumococcal (Pnc) serotype specific capsular polysaccharide (PS) antigen MPs. Microencapsulation of Pnc PS was successful with a product yield of >72%. The MP size, 1-5 μm, and negative zeta potential (~26.5 mV) were optimized to ensure effective uptake and presentation of Pnc PS antigens to immune cells. In mice, ST 19F and 23F MPs exhibited >10-fold increase (P<0.01) in ST specific IgG response over PS in solution given with or without alum. Relatively higher immune response was observed for ST 6B MPs when compared to PS solution; however, ST6B PS solution along with alum resulted in an overall higher response when compared to ST6B MPs. Microencapsulation may offer a simple and effective mechanism for the immune-enhancement of poorly immunogenic antigens such as Pnc PS.

Keywords: Pneumonia; Vaccine; Microparticles; Spray drying; Pneumococcal polysaccharide

Introduction

The field of vaccine research has made great strides over the last decade. In the past, vaccines were usually comprised of either live attenuated or killed whole cells of target pathogens. Even though such vaccines most often provided an immune response comparable to natural infection in terms of duration and stimulation of immune system, they caused several unwanted local or systemic immunological complications that at times outweighed benefits. Furthermore, some of the chronic pathogens such as Mycobacterium tuberculosis, HIV and HCV require alternate vaccine strategies to address their intra-cellular parasitic nature [1-3]. Development of recombinant, synthetic or purified immunogenic component(s) instead of whole cells of pathogen has shown promising results and steered vaccine development away from live and attenuated vaccines. The poor immunogenic potential of some of these components necessitates the inclusion of an adjuvant to enhance or prolong an antigen specific immune response [3]. Addition of adjuvants to the vaccine formulations in turn resulted in adverse events and undesirable immunological complications [4].

Microparticle and nanoparticle based drug delivery system is an emerging technology which has shown great promise in the delivery of small molecule anti-cancer drugs and in particular, large molecule vaccines [5-8]. Microparticle vaccines have the ability to release antigens over time resulting in a robust immunity with a single dose thereby eliminating the need for multiple doses along with reducing the chances of vaccine dropouts [9]. As discussed earlier, duration and simulation of the immune response are two critical aspects of vaccine formulations [10,11]. Microparticle based vaccine delivery systems not only enhance the immune response by stimulating antigen presenting cells (APCs) that preferentially take up particulate antigens in comparison to antigen solution but also prolong antigen exposure by creating a depot effect resulting in modified/sustained release of antigens [12-14].

Polysaccharides (PS) on their own are known to be poor immunogens, since PS are T-independent antigens that do not result in long term memory response [15,16]. Conjugation of PS to a carrier protein converts them to T-dependent antigens that confer immune-memory [17,18]. Streptococcus pneumoniae is a commensal bacterium found in the upper respiratory tract. Globally, pneumococcal disease due to this pathogen is prevalent in all ages [19-21]. The PS capsular chemistry and serological reactions has so far yielded greater than 103 pneumococcal (Pnc) serotypes (STs). There are conjugate and non-conjugate vaccines available for S. pneumoniae, including the 23-valent non-conjugated polysaccharide vaccine, Pneumovax–23 (Merck for adults > 65 years of age); and the 13-valent conjugate vaccine, Prevnar13 (PCV13, Wyeth, for patients of all ages). Pneumovax–23 is composed of purified Pnc capsular PS from 23 different STs. Pneumococcal capsular PS in their native form are poor immunogens [22,23] therefore the 23-valent Pnc PS vaccine (Pneumovax) are less effective in children and most effective in adults [23,24]. The major limitation of Pnc PS vaccines is that it does not protect from Pnc STs that are not in the vaccine [25]. Due to large number of circulating Pnc STs, targeted removal of only a limited number of circulating STs by the current available vaccines creates a need for those STs that are not protected against in the marketed vaccines. Therefore, Pnc disease due to serotype deletion or replacement is inevitable [26]. Moreover, protein conjugation of PS antigens is a complicated process that increases the production cost of the vaccine. Considering the

*Corresponding author: Prathap Nagaraja Shastri, College of Pharmacy and Health Sciences, Nanotechnology Laboratory, Mercer University, Atlanta, GA 30341, USA
Tel: +1 678 547 6353; Fax: +1 678 547 64230; E-mail: prathap.shastri@ambrx.com

Gowrisankar Rajam, Division of Bacterial Diseases, Centers for Disease Control and Prevention, Atlanta, GA 30333, USA, Tel: 404 639 2630; E-mail: rajamshanks@gmail.com

Received: May 22, 2018; Accepted: May 29, 2018; Published: June 05, 2018


Copyright: © 2018 D’Souza B, et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.
potential benefits of particle-based drug delivery systems, we worked on development of novel microparticulate polysaccharide antigen prepared using a one-step spray drying technology. This manuscript summarizes microparticles effectiveness in resulting immunogenic response to S. pneumoniae serotype specific to capsular polysaccharide antigens.

Materials and Methods

Preparation and characterization of microparticle formulation

Microparticle formulation optimization: The lyophilized Pnc capsular PS (ST 6B, 19F and 23F; ATCC, Manassas, VA) were solubilized to make a stock solution of 2 mg/mL using de-ionized water with overnight agitation at 4°C. Pnc PS MPs were formulated using a one-step spray drying technique [27]. Briefly, the MPs were prepared with mouse serum albumin (Thermo Fisher, Waltham, MA) matrix which was cross linked for 8 hours using glutaraldehyde. Sodium bisulphite was used to neutralize the excess glutaraldehyde, and 1% w/v chitosan was added to the resulting mixture to optimize the formulation. All the excipients added were Federal Drug Administration (FDA) Generally Recognized as Safe (GRAS) certified reagents. Pnc ST specific PS was maintained at 1% (v/v) concentration and spray dried in a Buchi Mini Spray dryer (B-191, Buchi Corporation and New Castle, DE). In the spray dryer, the solution passes through an atomizer, which disperses the solution into a drop size spray in a controlled manner. The small sized drops encompass a large surface area and dries up quickly to form microparticles upon transient exposure to elevated temperatures.

The percentage product yield of MPs was calculated using the formula given below:

\[
\text{Percentage product yield} = \frac{\text{Weight of microparticles obtained after spray drying}}{\text{Total amount of all polymers along with antigen prior to spray drying}} \times 100
\]

Microparticle size distribution: The uptake of MPs by antigen presenting cells (APCs) including macrophages and dendritic cells can be greatly influenced by their particle size. Microparticle suspension (10% w/v) was made in 10 mL deionized water and sonicated for 1 minute. Particle size of the suspension was measured using Zetasizer Nano ZS, Malvern Instruments, Worcester, UK), adopting the methodology previously described [28].

Zeta potential: Achieving a well dispersed suspension without aggregation is key for successful administration of MP based formulation, and Zeta potential provides the measure of the same. Aqueous suspension (10% w/v) of Pnc PS loaded and blank MPs were tested for their Zeta potential using Malvern Zetasizer (Malvern Instruments, Worcestershire, UK), adopting the methodology previously described [28].

Scanning electron microscopy of microparticle: The JEOL scanning electron microscope was used to study the size, surface morphology, and uniformity of the MP formulation. Briefly, the formulations were evenly spread on metal stubs and coated with gold and dried. These gold-coated MPs were vacuum dried and then imaged under a nitrogen airflow stream.

Antigen epitope integrity

The epitope integrity of the antigen is critical for the success of formulation process and loading efficiency is crucial to calculate the dosage for in vivo studies. The phenol-sulphuric acid assay is generally used to quantify polysaccharides [29]. The use of chitosan (polysaccharide) as an excipient in this MP formulation warranted an alternate technique due to its interference in the PS quantification with the phenol-sulphuric acid technique. To address this challenge, an alternate technique that utilizes the principle of homologous competition between an antigen and specific antibody was implemented. Extraction of antigens from the MPs was done by treating with 50 mM Tris HCl, 10 mM EDTA buffer, and followed by incubation with 0.2N NaOH and protease overnight at 37°C. The diluted extract was further incubated with a Pnc reference standard serum, 89SF (1:20) overnight at 4°C with head to toe rotation. Additional tubes of standards were prepared containing the reference standard and range of concentrations of the Pnc ST PS (23F, 6B, and 19F) to generate Pnc Ps calibration graph. Following overnight incubations, the mixtures were centrifuged at 8000 x g for 5 minutes and the supernatant was tested for Pnc ST specific IgG using microsphere based multiplex technique [30]. Pnc PS in MPs was calculated from the 89SF-Pnc PS calibration graph. Being an Pnc PS specific antibody capture technique, this also provides valuable information regarding the epitope integrity in the MP formulations.

Test for microparticle cytotoxicity

The epitope integrity of the antigen is critical for the success of formulation process and loading efficiency is crucial to calculate the dosage for in vivo studies. The phenol-sulphuric acid assay is generally used to quantify polysaccharides [29]. The use of chitosan (polysaccharide) as an excipient in this MP formulation warranted an alternate technique due to its interference in the PS quantification with the phenol-sulphuric acid technique. To address this challenge, an alternate technique that utilizes the principle of homologous competition between an antigen and specific antibody was implemented. Extraction of antigens from the MPs was done by treating with 50 mM Tris HCl, 10 mM EDTA buffer, and followed by incubation with 0.2N NaOH and protease overnight at 37°C. The diluted extract was further incubated with a Pnc reference standard serum, 89SF (1:20) overnight at 4°C with head to toe rotation. Additional tubes of standards were prepared containing the reference standard and range of concentrations of the Pnc ST PS (23F, 6B, and 19F) to generate Pnc Ps calibration graph. Following overnight incubations, the mixtures were centrifuged at 8000 x g for 5 minutes and the supernatant was tested for Pnc ST specific IgG using microsphere based multiplex technique [30]. Pnc PS in MPs was calculated from the 89SF-Pnc PS calibration graph. Being an Pnc PS specific antibody capture technique, this also provides valuable information regarding the epitope integrity in the MP formulations.

Test for microparticle cytotoxicity

Blank MPs were tested for cytotoxicity before animal immunization. Cell cytotoxicity studies were done with blank MPs using murine macrophages, RAW 246.7. About 2 x 10^4 cells/mL of RAW 246.7 macrophage cells were plated in a 96 well plate. Microparticle suspension in the concentration range of 10-1000 µg/mL was added on to the cells. Atropine sulphate was used as the positive control at a concentration of 10 mg/mL. The plates were incubated for 12 hours at 37°C with 5% CO2. Upon completion of incubation, about 20 µL well of Alamar blue dye (Life Technologies, Grand Island, NY) was added and the plate was read at 490 nm in a microplate reader. Alamar blue is a redox indicator and color change from blue to purple to red is directly proportional to the cellular respiratory activity and thus, viability.

In vivo immunization studies

Mouse strains: Female Swiss Webster mice (Charles River Laboratories, Wilmington, MA) 6 to 10 weeks of age were used in
this study. All experiments were approved by the Institutional Animal Care and Use Committee (IACUC) and conducted according to the institutional ethical guidelines for animal experiments and safety guidelines. Alum (Al₂(OH)₃) was used as an adjuvant. Each Pnc serotype testing, six groups (n=10/group) of mice were included. This includes, control groups that received blank MPs, and Pnc ST specific PS in solution with and without alum. Apart from Pnc PS MPs, another group of animals received Pnc PS in solution with blank MPs separately to test the role of MPs as an external adjuvant. Two doses of 50 μg of MP/Pnc PS were administered via subcutaneous administration with a 15 day dosing interval. Blood was collected on day 0 prior to immunizations, at day 14 and one week after the last dose.

**Pnc ST specific IgG quantification**

Serum samples were analyzed using multiplex Luminex technique [30] to specifically detect IgG with Pnc STs with some modification. The filter plates (MABVN1250, Millipore Corp, Billerica, MA) were pre-wet with 200 μL assay buffer (0.1% BSA in PBS). Serum dilutions were carried out in a 96-well round bottom titer plate (Dilution plate, CLS3799, Sigma, St. Louis, MO). Test serum samples were diluted 2-fold for 7 dilutions starting at 1/50. For reagent controls, each assay included buffer blanks. Replicates were maintained for unknown serum and blanks. The Pnc ST specific PS conjugated fluorescent microspheres were added to the plates at 25 μL/5000 beads/well/PS after 2X PBS pre-wash. To this, 25 μL/well of serum sample was transferred from the dilution plates and incubated at 37 ºC for 30 minutes with agitation. After 3 washes with 100 μL/well PBS, a secondary PE labeled anti-mouse antibody (Moss Pasadena, MD), was added and incubated for 20 minutes with agitation. On completion, plates were washed three times with 100 μL/well PBS and the beads were resuspended in 130 μL of luminex sheath fluid before reading in the luminex reader (L200, Luminex Corp, Austin, TX).

**Statistical Analysis**

Animal studies were carried out in two independent cycles. Masterplex QT (Miraibio, Hitachi, Irvine, CA) was used for the analysis of mouse serum IgG data extracted from the luminex reader. One-way ANOVA and Tukey test was performed on in vivo data using graph pad Prism. For all the data collected, results were tabulated as significant at p values <0.05. In the absence of reference standard calibrated for mouse sera, animal experiment groups and study time points were compared based on their respective Median Fluorescence Intensity (MFI) and immune response was expressed as fold changes.

**Results**

**Characterization of albumin microparticles**

The percentage product yield was measured as 76% ± 3.54 and 75% ± 4.18 for blank and antigen loaded MPs, respectively. The size of the MPs ranged from 1.2-2.4 μm in majority with overall size range of 1.5 μm (Table 1). No significant difference was seen in the particle size between the blank and the PS antigen loaded MPs. The zeta potential was found to be -24 mV and -26.5 mV for blank and antigen loaded MPs respectively (Table 1). Scanning electron microscopy revealed uniform spherical shape in MPs with each particle separated distinctly (Figure 1).

**Loading efficiency of microparticle and antigen epitope integrity**

Antigen epitope integrity was verified by quantifying Pnc ST specific PS using Luminex as described in section 2.2. 89SF, the reference serum has ST specific IgG assignments that react to Pnc ST specific Ps. The quantification assay demonstrated 72.5%, 80.76%, and 83.64% antigen loading efficiency (Table 1).

**In vitro microparticle cell cytotoxicity**

**In vitro** cell toxicity test was performed to determine the effect of MPs on eukaryotic cells using murine macrophages, RAW 246.7 model. Data presented in Figure 2 show that MPs have little or no toxic effect (p<0.05) on RAW 246.7 cells.

**In vivo immunization studies**

Bioactive levels of Pnc STs 19F and 23F were determined from serum of mice dosed with the MPs containing antigens. Both ST specific PS MP treatment groups resulted in an apparent greater immune response with 19F and 23F MPs resulting in 10- and 20-fold
statistically significant increase in immune response as compared to other groups including antigen solution control and with adjuvant alum (Figure 3a and 3b). Although, in case of 6B serotype, vaccine solution with alum showed significantly better immune response when compared to 6B MPs. However, when compared to 6B solution alone, MP vaccine produced higher immune response (Figure 3c).

Discussion

This research project was focused on optimizing a novel particulate formulation to improve the efficacy of pneumococcal antigens in a vaccine as well as to extend vaccine coverage [18,22]. We formulated pneumococcal PS MPs having a homogenous size range of 1-5 µm (90% MPs 1-2 µm). The MPs at this size range ensures effective uptake and processing of Pnc PS by antigen presenting cells (APC) and dendritic cells (DC) of the immune system. Buchi-190 was used to spray dry the MPs containing the ST specific Pnc Ps. Spray drying is a commonly used technique to prepare small to medium sized batches of particulate delivery systems including antigens and large molecules [31,32]. This technique ensures the formulation of MPs with size and surface properties critical for the uptake and stimulation of innate immune cells. Another critical physical characteristic of MPs is its ability to stay non-aggregated. Aggregated MPs lead to gross increase in their size leading to a sharp decline in the immunological benefits. The surface charge of MPs made by the spray drying technique were > ± 30 mV
which would impede aggregation and make the MPs more suitable for uptake by immune cells for an effective vaccination. In this study, optimal formulation condition yielded Pnc PS MPs at -25mV in water that would discourage particle aggregation resulting in heightened immunological benefits. The smooth spherical surface morphology of the MPs as seen in SEM images suggests the possibility of uniform release of antigens in vivo. This would allow slow and sustained release of antigen encapsulated within the MP when taken up by antigen presenting cells (APCs) and dendritic cells (DCs).

A luminex-based bioassay was developed to quantify Pnc PS in MPs as the use of polysaccharide excipients impeded the use of the traditional phenol-sulfuric acid assay for PS quantification. This bioassay was able to quantify the Pnc ST specific PS antigens as well as determine the bioactivity and epitope integrity of PS in MPs after formulation. In its multiplex format, this bioassay also offers the potential for the simultaneous quantification of multivalent MP formulation. Using this technique, the MP loading efficiency for all 3 Pnc ST was found to be >70%. This also reiterates the epitope integrity of the encapsulated Pnc PS as the analytical basis of this bioassay is the reaction between the antibody and the complementary epitope on the Pnc PS. This proves that the MP formulation technique used in this study is effective and has minimal impact on the epitope integrity of the antigen in MPs. Encouraged by the successful encapsulation of Pnc ST Ps resulting in MPs with desirable physicochemical properties and safety profiles, the Pnc PS MPs were compared for their immunogenicity in vivo to PS in solution. The animal experiments yielded a superior immune response with ≥10-fold increase in Pnc ST specific IgG for 19F and 23F in MPs as compared to the PS in solution. The serotype 6B in the MPs did show a robust IgG response, however the increase was not as significant as the 19F and 23F serotypes in MPs versus their solutions. This was not surprising since serotype 6B is one of the least immunogenic in the spectrum of serotypes used in the 23-valent Pnc PS vaccine (Pneumovax) formulation for adults. Further, ST 6B in the Prevnar-13 pneumococcal conjugate-vaccine has also been reported to be less immunogenic as compared to the other STs in the vaccine [33-35]. Higher loading concentration and/or alternate MP formulation strategies may help to improve the PS 6B immunogenicity.

The enhanced ST specific immunity can be attributed to the optimized size and surface morphology of the PS MPs that is critical for uptake by immune responsive cells. It has been shown that 2-8 μm MPs loaded with hepatitis B surface antigen elicited higher IgG response given intra muscularly [28]. Kalainidhi et al., suggested that MPs with a size distribution around 500 nm and up to roughly 3-5 μm were taken up mainly by phagocytosis and promote humoral immune response [36]. As suggested by Balasse et al., the 1-5 μm size for the optimized polysaccharide MPs help in increasing duration of exposure to DCs resulting in an increase of antigen specific immune response due to particulate nature over solution form [37].

Interestingly, blank MPs failed to exert an adjuvant effect on the Pnc PS solution form. This reiterates that steady state release of MPs entrapped antigens is crucial for the continuous insult of the immune system resulting in improved immune response. Several researchers have reported the usefulness of the blank MPs as a non-specific adjuvant but there are multiple factors that may govern this response. This property could be determined by the chemistry of the polymer or the reactivity of the polymer with the animal species tested to name a few. In one study, ionic polyphosphazene or PLGA were shown to have demonstrable immune adjuvant activity on their own [38,39].

We have shown that optimized microparticles with serotype specific antigens provide an opportunity for formulating pneumococcal vaccines that can be customized to specific geographical and demographic requirements using a cost-effective manufacturing process. This process is vital for diseases due to multiple virulent serotypes/serogroups in circulation. In our earlier studies, oral MPs with several vaccines including pertussis, typhoid and tuberculosis vaccine have shown promising outcome in animal models [40-43]. The present study aligns with the previous observations and holds promise as the next generation vaccine formulation strategy for pneumococcal disease. Pnc PS MPs have been extremely efficacious in humoral immune induction and ST specific antibody response in vivo. Additional studies to deduce the effect of MP immunization on the cellular immune response, Pnc carriage and disease in vivo is recommended to consolidate these MPs as Pnc PS vaccine delivery systems.

**References**


