4-2-2012

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Petersen, L. K.; Phanse, Y.; Ramer-Tait, Amanda E.; Wannemueler, M. J.; and Narasimhan, B., "Amphiphilic Polyanhydride Nanoparticles Stabilize *Bacillus anthracis* Protective Antigen" (2012). Faculty Publications in Food Science and Technology. 205. [http://digitalcommons.unl.edu/foodsciefacpub/205](http://digitalcommons.unl.edu/foodsciefacpub/205)

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Amphiphilic Polyanhydride Nanoparticles Stabilize Bacillus anthracis Protective Antigen

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Abstract

Advancements towards an improved vaccine against Bacillus anthracis, the causative agent of anthrax, have focused on formulations composed of the protective antigen (PA) adsorbed to aluminum hydroxide. However, due to the labile nature of PA, antigen stability is a primary concern for vaccine development. Thus, there is a need for a delivery system capable of preserving the immunogenicity of PA through all the steps of vaccine fabrication, storage, and administration. In this work, we demonstrate that biodegradable amphiphilic polyanhydride nanoparticles, which have previously been shown to provide controlled antigen delivery, antigen stability, immune modulation, and protection in a single dose against a pathogenic challenge, can stabilize and release functional PA. These nanoparticles demonstrated polymer hydrophobicity-dependent preservation of the biological function of PA upon encapsulation, storage (over extended times and elevated temperatures), and release. Specifically, fabrication of amphiphilic polyanhydride nanoparticles composed of 1,6-bis(p-carboxyphenoxy)hexane and 1,8-bis(p-carboxyphenoxo)-3,6-dioxaoctane best preserved PA functionality. These studies demonstrate the versatility and superiority of amphiphilic nanoparticles as vaccine delivery vehicles suitable for long-term storage.

Keywords

Protective antigen for anthrax; protein stability; polyanhydrides; nanoparticles

Introduction

Anthrax is a significant public health concern due to its potential as a bioterrorism and biowarfare agent. Use of the currently licensed alum-adjuvanted anthrax vaccine adsorbed (AVA) poses many concerns due to its high reactogenicity, multi-dose (five) immunization schedule followed by yearly boosters, and painful side effects¹. The recombinant protective antigen (PA) from Bacillus anthracis, the key element of the AVA vaccine that confers immunity against an anthrax infection, has been the focus for new vaccines². PA is the receptor-binding subunit for both lethal factor (LF) and edema factor (EF), which upon binding, lead to the formation of lethal toxin (LT) and edema toxin (ET), respectively³. Although highly immunogenic, PA is unfortunately a labile recombinant protein, especially in low pH and high temperature environments⁴–⁷. These stability concerns have led to manufacturing roadblocks during a Phase I clinical trial for a new anthrax vaccine based on PA⁸.

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In addition to stability concerns during the vaccine manufacturing process, there are other obstacles for PA-based vaccines. PA circulating in the bloodstream is eliminated within six hours, meaning its limited in vivo half-life may fail to induce a protective immune response. Studies by Zinkernagel and others have shown that antigen persistence is a key element associated with the development of a protective immune response. Several methods have been employed to increase PA availability to the immune system and improve its immunogenicity, including engineering recombinant PA, incorporating other antigens or adjuvants in the vaccine, evaluating alternate delivery routes, and using controlled delivery systems. Currently, the AVA vaccine is administered intramuscularly and many murine studies have focused on subcutaneous delivery of PA. As an alternative, intranasal vaccination has been reported as the best route for induction of a protective immune response. While PA is a relatively good immunogen, it may be essential to use an adjuvant; however, well known mucosal adjuvants (e.g., cholera, pertussis, and edema toxins and CpG DNA) have limitations due to their potential toxicity when delivered by this route in humans.

This work focuses on the development and refinement of a controlled delivery system for PA based on biodegradable polyanhydride nanoparticles. Polyanhydrides are a class of biomaterials with excellent biocompatibility and have been extensively studied as vaccine delivery vehicles and adjuvants. These polymer formulations provide safe parenteral delivery, enhanced adjuvanticity, sustained release of encapsulated antigens, and a stabilizing environment for protein antigens during fabrication, storage, and release. Specifically, micro- and nanoparticles made of copolymers based upon sebacic anhydride (SA), 1,6-bis(p-carboxyphenoxy)hexane (CPH), and 1,8-bis(p-carboxyphenoxy)-3,6-dioxoaestane (CPTEG) are suitable for inhalation or injection. Moreover, they possess tunable antigen release kinetics that provide an in vivo antigen depot, thereby allowing for sustained antigen exposure and eliminating the need for multiple administrations. These surface eroding polymers slowly release encapsulated antigen, minimizing exposure to unfavorable aqueous environments.

The overall goal of this work was to design nanoparticle-based formulations capable of encapsulating and releasing biologically functional PA that results in a robust, high avidity, neutralizing antibody response after a single administration. In this regard, nanoparticles based on 50:50 CPTEG:CPH, 20:80 CPTEG:CPH, 50:50 CPH:SA, and 20:80 CPH:SA were investigated for their ability to stabilize and release biologically active and immunogenic PA. In contrast to CPH:SA nanoparticles, amphiphilic CPTEG:CPH nanoparticles better preserved the functionality and immunogenicity of PA during nanoparticle fabrication, storage, and release. The use of amphiphilic polyanhydride nanoparticles as delivery vehicles presents a viable solution for long-term vaccine storage.

**Experimental Section**

**Materials**

Chemicals needed for polymerization, nanoparticle fabrication, and buffer preparation included acetic anhydride, chloroform, methylene chloride, petroleum ether, pentane, monobasic potassium phosphate, and dibasic potassium phosphate; all were purchased from Fisher Scientific (Fairlawn, NJ). Monomer synthesis utilized the following chemicals: 1,6-dibromohexane, triethylene glycol, 4-hydroxybenzoic acid, and 1-methyl-2-pyrrolidinone; these were purchased from Sigma Aldrich (St. Louis, MO); 4-p-fluorobenzonitrile was purchased from Apollo Scientific (Cheshire, UK); and sulfuric acid, acetonitrile, dimethyl formamide, toluene, and potassium carbonate were obtained from Fisher Scientific. Deuterated chemicals for NMR (nuclear magnetic resonance) analysis, chloroform and dimethyl sulfoxide were purchased from Cambridge Isotope Laboratories (Andover, MA).
Polymer synthesis, nanoparticle fabrication, protein encapsulation, and characterization

The CPH and CPTEG monomers were synthesized as described previously. The SA monomer was purchased from Sigma Aldrich (St. Louis, MO). Pre-polymer and polymers were synthesized as described before. The CPTEG:CPH and CPH:SA copolymers were formed into nanoparticles as described previously. Briefly, PA (NIH Biodefense and Emerging Infections Research Resources Repository, Manassas, VA) loaded nanoparticles (2% (w/w)) were fabricated by dissolving the desired polymer in methylene chloride at a concentration of 25 mg/mL. The desired amount of lyophilized PA was added to the dissolved polymer, the solution sonicated (40 Hz for 30 s), poured into a non-solvent (pentane) at a solvent to non-solvent ratio of 1 to 200, and then vacuum filtered to recover the PA-loaded nanoparticles (20–150 mg batch size). Polymer molecular mass, chemical structure, and chemical composition were determined with 1H NMR spectroscopy using a Varian VXR 300 MHz spectrometer (Varian Inc., Palo Alto, CA). Specifically, the molecular mass was determined by end group analysis of NMR spectra. Nanoparticle size and morphology were characterized with scanning electron microscopy (SEM; JEOL 840 A, JEOL Peabody, MA).

In vitro protein release kinetics

PA-loaded nanoparticles at a concentration of approximately 50 mg/mL were incubated with 0.1 M PBS buffer (pH 7.4) in capped microcentrifuge tubes. Samples were sonicated to ensure uniform nanoparticle distribution and placed in a shaker/incubator at 37°C. Sample supernatants were removed (entire volume) incrementally to measure the amount of released PA (PA\textsubscript{rel}), which was analyzed using a micro-bicinchoninic acid (BCA) assay. Once supernatants were removed, equivalent volumes of PBS buffer were added back to maintain perfect sink conditions. Data is presented as cumulative percentage of PA released, which was determined by normalizing the cumulative amount released at each time point by the total amount encapsulated into the nanoparticles.

In vitro pH change upon nanoparticle degradation

Blank nanoparticles at a concentration of approximately 100 mg/mL were incubated with 0.1 M PBS buffer (pH 7.6) in microcentrifuge tubes. Samples were sonicated to ensure uniform nanoparticle distribution and placed in a shaker/incubator at 37°C. The pH measurements were carried out at various time points with a pH probe to monitor the buildup of acidic degradation products.

In vitro release of PA for analysis of bioactivity, antigenicity, and structure

Treatment groups tested for their effect on PA stability were 20:80 CPTEG:CPH, 50:50 CPTEG:CPH, 20:80 CPH:SA, and 50:50 CPH:SA nanoparticles, native PA (i.e., positive control stored at −20°C), and protein exposed to nanoparticle fabrication conditions (NFC) which included sonication (40 Hz for 30 s), solvent exposure (methylene chloride and pentane), and vacuum drying. All groups were sonicated and incubated with 0.1 M PBS buffer (pH 7.4) in microcentrifuge tubes in a shaker/incubator at 37°C. In order to accumulate enough of the PA\textsubscript{rel} for characterization, PA-loaded nanoparticles were incubated for two days. Following release, supernatants were removed and the concentration, antigenicity, bioactivity, and structural alterations of the PA\textsubscript{rel} were assessed.

Circular dichroism: secondary structure

Far UV circular dichroism (CD) (190–250 nm) was used to study the changes in protein secondary structure of the PA\textsubscript{rel}. All CD spectra were collected for samples on a Jasco J-170 Spectropolarimeter (Easton, MD). The emission spectrum from 200–260 nm was analyzed for shifts in molar ellipticity and wavelength, which are indicative of alterations in protein...
secondary structure. The experiments were performed in triplicate using a 200 μL sample volume.

**Fluorescence spectroscopy: tertiary structure**

Fluorescence spectroscopy was used to study the changes in tertiary structure of the PArel with a Cary Eclipse fluorescence spectrometer (Varian, Inc., Palo Alto, CA). Samples were excited at a wavelength of 280 nm and a voltage of 800 V, which resulted in an emission spectrum that corresponds to the tryptophan and tyrosine residues of the protein. The emission spectrum from 300–500 nm was analyzed for shifts in peak intensity and wavelength that are indicative of alterations in protein tertiary structure. The experiments were performed in triplicate.

**In vitro release of PA for in vivo administration of the PArel**

PA-loaded nanoparticle (2% w/w) samples (200 mg) were incubated in 0.1 M PBS (pH 7.4) for 14 days at 4°C in dialysis cassettes (4,000 Da MWCO) to accumulate enough of the PArel for immunization. This release was carried out at 4°C to prevent temperature induced instability. Protein concentration was determined with a micro-BCA assay and all samples were adjusted to a final concentration of 200 μg/mL. Equal amounts of PA released from 20:80 CPTEG:CPH, 50:50 CPTEG:CPH, 20:80 CPH:SA, and 50:50 CPH:SA nanoparticles and native PA were adsorbed to Imject Alum (Fisher Scientific). A/J mice (Jackson Laboratories, Bar Harbor, Maine) were immunized subcutaneously in the nape of the neck with a dose of 10 μg of the PArel or native PA (3 or 4 mice per group). Each mouse was given a second immunization on day 15 with 10 μg of the PArel or native PA corresponding to the primary immunization regimen, euthanized on day 21, and serum collected. All animal procedures were conducted with the approval of the Iowa State University Institutional Animal Care and Use Committee.

**Enzyme-linked immunosorbent assay (ELISA): PA-specific serum antibody response**

PA-specific serum antibody response (total IgG [H+L]) and PArel antigenicity were analyzed via ELISA using a previously described method with modifications. High-binding 96 well ELISA plates (Fisher Scientific) were coated in PBS buffer (pH 7.2, 0.05M) with 0.5 μg/mL native PA or PArel overnight at 4°C. After washing with buffer (PBS containing 0.5% Tween 20 (PBS-T)), plates were blocked for 2 h (PBS containing 0.5 % Tween 20 (PBS-T) and 2% gelatin) at room temperature, washed three times in PBS-T, and incubated with mouse serum (from individual mice, not pooled) for 24 h at dilutions from 1:1,000 to 1:1,000,000 overnight at 4°C. The following day, plates were washed three times in PBS-T, incubated with alkaline phosphatase-conjugated goat anti-IgG [H+L] antibody (Jackson ImmunoResearch Laboratories, West Grove, PA) for 2 h at room temperature, washed again, and incubated with alkaline phosphatase substrate buffer (Sigma Aldrich) for 2 h at room temperature. Absorbance was measured at an optical density (OD) of 405 nm using a Spectramax 190 Plate Reader (Molecular Devices, Sunnyvale, CA). Each sample was analyzed in triplicate.

**ELISA for serum antibody avidity**

Serum antibody avidity was determined by modifying an ELISA method described previously. The avidity assay was performed similarly to the serum antibody response protocol described above. Following serum incubation for 24 h at a dilution of 1:1,000,000, the chaotropic agent, sodium thiocyanate, was added to the wells using incremental increases over a concentration range from 0 to 5 M. Plates were incubated for 20 min at room temperature and then washed four times in PBS-T. The detection of bound serum antibody was performed as described above. Relative avidity index is defined as the
concentration of sodium thiocyanate required to decrease the optical density by 50% of that obtained for each control well (i.e., no sodium thiocyanate). Antibody titers were too low to measure the avidity of the antibody response for serum samples obtained from mice immunized with PA released from the CPH:SA nanoparticles.

**Lethal toxin in vitro cytotoxicity assay**

The lethal toxin *in vitro* cytotoxicity assay was modified from a previously described method. The murine macrophage-like cell line RAW 264.7 obtained from ATCC (Manassas, VA) was maintained in culture at 37°C in a 5% CO₂ humidity using high glucose DMEM medium (Invitrogen, Carlsbad, CA) supplemented with 10% heat-inactivated FBS (Valley Biomedical, Winchester, VA), 100 IU/ml penicillin and 10 μg/ml streptomycin (Mediatech, Herndon, VA). For the lethal toxin *in vitro* cytotoxicity assay, RAW cells were transferred to a 96-well plate (100,000 cells per well) grown to approximately 90% confluency in 100 μL medium over two days. Approximately 12 μL of each treatment (PA<sub>rel</sub>, PA-loaded nanoparticles, blank nanoparticle controls, medium only control (negative control and 100% cell survival), native PA (positive control and 100% cell death), LF (NIH Biodefense and Emerging Infections Research Resources Repository, Manassas, VA) control, and cell lysis buffer) diluted in RAW medium were added to the cells with 0.3 μg/mL LF for no more than 6 h. The amount of PA released from the nanoparticles was calculated from the release curve in Figure 1. This assay was also utilized to investigate antibody neutralization and shelf life (i.e., PA stability) of the PA-loaded nanoparticles. For antibody neutralization assessment, individual serum samples (dilutions ranging from 1:100 to 1:204800) from the different immunized mouse groups (described in the in vitro release of PA for in vivo administration of the PA<sub>rel</sub> section) were added to the cells along with native PA (0.8 μg/mL) and LF (0.3 μg/mL). For antibody neutralization assessment, the positive control was medium only (100% cell survival) and the negative control was native PA (100% cell death). Samples were then allowed to incubate for up to 6 h, after which the CellTiter 96® Non-Radioactive Cell Proliferation Assay (MTT) (Promega, Madison, WI) was used to determine cell viability. Optical density (OD) was measured at 570 nm with a background (690 nm) subtraction as recommended by the manufacturer’s protocol. Antibody neutralization was considered significant when measured OD values fell below 50% of the positive control OD value. Data are presented as % residual activity, as determined by Equation 1:

\[
\%\text{ residual activity} = \frac{OD_{\text{sample}} - OD_{\text{neg control}}}{OD_{\text{pos control}} - OD_{\text{neg control}}} \times 100
\]

**Thermal stability of stored dry powder PA-loaded nanoparticle formulations**

Bioactivity was tested with the lethal toxin *in vitro* cytotoxicity assay to determine the ability of each formulation to preserve and release active PA. The thermal stability upon storage was studied in formulations that included PA loaded into the nanoparticles (dry) (labeled by polymer chemistry), PA dissolved in PBS buffer (labeled “PBS”), or PA absorbed to alum (labeled “Alum”). All formulations were stored under desiccant at four different temperatures, 40, 25, 4, and -20°C, for four months. Following storage, the formulations were tested at prescribed times for their ability to preserve PA activity using the lethal toxin *in vitro* cytotoxicity assay as described previously. PA-loaded nanoparticle bioactivity was evaluated at a concentration of 125 μg/mL for 6 h. This nanoparticle dose was chosen because after 6 h of release (assay incubation time) the resultant PA concentration was the same as that administered in the positive control (i.e. 0.8 μg/mL native PA). It is important to note that nanoparticle formulations investigated in this study were stored dry and were not added to aqueous buffers or medium until initiation of the
cytotoxicity study, which lasted for 6 h. This is in contrast to the PA controls (i.e., PBS and Alum), which were stored in solution to mimic the current storage strategy of the AVA vaccine. The nanoparticle storage conditions in this study are distinctly different from those in the studies investigating PA release, activity, structure, antigenicity, and immunogenicity (Figures 1–4) in which the nanoparticles were incubated in an aqueous environment for extended time periods (see previous methods) to enable sufficient release of the encapsulated protein.

Statistical analysis

Statistical analyses performed using JMP® 7 (Cary, NC) and comparisons between treatments were made with Tukey’s HSD (Honestly Significant Difference) to determine statistical significance and \( p \)-values \( \leq 0.05 \) were considered significant.

Results

Polymer and nanoparticle characterization

Consistent with previous work, polymer characterization by \(^1\)H NMR indicated that the polymer molecular weights were within the desired ranges of 9,000–15,000 Da\(^{31, 33, 34}\). The structure and composition of the polymers were also confirmed using NMR. SEM images (data not shown) indicated that protein-loaded and blank nanoparticles resulted in similar size \((212 \pm 43 \text{ nm})\) and morphology as previously reported\(^{16, 19, 27, 29, 35}\). Particle yield was approximately 80% (recovery after filtration).

In vitro PA release and pH of nanoparticle degradation were dictated by chemistry

Release of PA and change in pH due to nanoparticle degradation were evaluated as a function of time. As shown in Figure 1, polymer chemistry-dependent protein release kinetics and buffer acidification were observed. Increasing the content of SA or CPTEG relative to the CPH content resulted in a more rapid release of encapsulated PA (Figure 1A). The 20:80 CPTEG:CPH particles provided the slowest PA release profile. This is consistent with previously published studies demonstrating chemistry-dependent release profiles of other proteins from polyanhydride nanoparticles\(^{22–25, 27, 28, 30, 36}\). The effect of the polymer chemistry is also reflected in the pH profile of the degrading nanoparticles, as shown in Figure 1B. The acidic monomers produced from the degradation of CPH:SA nanoparticles have been shown to be water soluble\(^{25}\) and result in a greater decrease in pH \((\Delta \text{pH} = 2.5)\) as compared to that of the CPTEG:CPH monomers \((\Delta \text{pH} = 0.6)\). In these experiments, an even greater pH decrease \((\Delta \text{pH} = 5.0)\) was observed upon degradation of poly(lactic-co-glycolic acid) (PLGA) nanoparticles (data not shown).

Protein activity and structural stability were preserved by amphiphilic CPTEG:CPH nanoparticles

To evaluate PA stability upon release from nanoparticles, a detailed structural evaluation of the PA\(_{\text{rel}}\) was performed (Figure 2). In this study, we analyzed the structural integrity of the PA\(_{\text{rel}}\) that accumulated over 48 h. During this time, the protein was continually exposed to the acidic microenvironments caused by the degradation of the polymers in an aqueous environment. The CD spectrum for native PA revealed a minimum at approximately 208–210 nm and a shoulder at 215–217 nm (Figure 2A). These findings were consistent with the known CD spectra reported previously for the secondary structure of PA, and indicate that PA consists mainly of a mixture of \(\alpha\)-helices and \(\beta\)-sheets\(^6, 37\). Only the PA exposed to NFC or released from 50:50 CPTEG:CPH nanoparticle formulations maintained the native secondary structure (Figure 2A). PA released from all other formulations (20:80
CPTEG:CPH, 20:80 CPH:SA, and 50:50 CPH:SA) resulted in a significant increase in molar ellipticity (Δε), indicating a loss of secondary structure.

Tertiary structure analysis, performed with fluorescence spectroscopy, revealed a red shift in peak position for the PA released from all the nanoparticle formulations (Figure 2B). This peak shift indicates exposure to a more polar environment indicative of protein unfolding. However, PA peak intensity was best preserved by encapsulation into and release from the amphiphilic 50:50 CPTEG:CPH nanoparticles and to a lesser extent by encapsulation into and release from 20:80 CPTEG:CPH nanoparticles. The nanoparticle fabrication conditions appeared to have no detrimental effects on the tertiary structure of PA. In contrast, the encapsulation of PA into and release from CPH:SA formulations resulted in a loss of tertiary structure.

Following the structural evaluation, the relative antigenicity and bioactivity were assessed for the PArel. The results of this study, which were used to identify lead candidate formulations for stabilizing PA, are summarized in Figure 3. PA released from amphiphilic 50:50 CPTEG:CPH nanoparticles retained almost 80% of its antigenicity (Figure 3A) and functional activity (Figure 3B). PA released from 20:80 CPTEG:CPH nanoparticles retained 52% of its activity but only 20% of its antigenicity, while CPH:SA-containing nanoparticles preserved very little PA activity and antigenicity. Nanoparticle fabrication conditions had little or no detrimental effect on the antigenicity or activity of PA (Figure 3).

In vivo immunization with the PA released from amphiphilic CPTEG:CPH nanoparticles resulted in high titer, high avidity, neutralizing antibody

Upon completion of the structural, antigenicity and bioactivity studies, it was hypothesized that the amphiphilic CPTEG:CPH nanoparticles would be the most viable candidates for preserving the PArel, thus enabling a primed PA-specific antibody response when administered in vivo. In order to determine the immunogenicity of the released protein, separate groups of mice were immunized with alum adjuvanted PArel from the four nanoparticle formulations. We hypothesized that if PA was degraded upon release from a particular nanoparticle formulation, as seen for the PA released from the CPH:SA nanoparticle chemistries (Figures 2 and 3), there would be a loss in the immunogenicity of the PA. Antibody responses induced by the released protein were characterized in terms of titer (Figure 4A), relative avidity (Figure 4B), and neutralizing antibody titer (Figure 4C). The data indicated that the PA released from CPH:SA nanoparticles induced a weak humoral response, while the PA released from the CPTEG:CPH nanoparticles or PA exposed to the nanoparticle fabrication conditions resulted in the induction of high titer and avid antibody responses that were statistically similar to those induced by native PA. Overall, these data indicate that the immunogenicity of PA released from the amphiphilic CPTEG:CPH nanoparticles was better preserved than that of PA released from CPH:SA nanoparticles, as evidenced by the 4 log₁₀ difference in antibody titer.

Shelf life stability of PA encapsulated into 50:50 CPTEG:CPH, 20:80 CPTEG:CPH, and 20:80 CPH:SA was superior to that of PA conjugated to alum

The ability of a PA-based vaccine to retain full immunogenicity upon storage is a major concern. As a measure of PA stability, an in vitro cytotoxicity assay was employed to assess the preservation of the biological activity of PA upon storage and immediate release from polyanhydride nanoparticles. This study was also an assessment of the thermal stability of the nanoparticles at various temperatures. During optimization of the cytotoxicity assay, we determined that co-incubating 125 μg/mL of 0.8% PA-loaded nanoparticles for 6 h with RAW 264.7 cells resulted in a concentration of PArel that was equivalent to that of the positive control (i.e., 0.8μg/mL native PA). Because there was no statistical difference in the
amount of PA released between the different nanoparticle chemistries in the first 6 h (Figure 1), direct comparisons between the various polyanhydride formulations can be made. In this study, the shelf life storage capabilities of i) PA loaded into polyanhydride nanoparticles (dry), ii) PA in PBS buffer (pH 7.6), or iii) PA adsorbed to alum were investigated at multiple time points following storage at 40, 25, 4, and −20°C. It is known that adjuvant solutions containing alum have acidic bulk conditions (pH = 5.6)38, so a PBS buffer group was included to investigate the effect of storage temperature at a neutral pH on the bioactivity of PA.

In order to assess the preservation of the biological activity after dry storage for the desired amount of time (1, 7, 14, 30, 60, and 120 days – only long term data are presented in this manuscript), PA-loaded nanoparticles were directly applied to cultures of RAW 264.7 cells at the initiation of a lethal toxin cytotoxicity assay. By employing these assay conditions, any of the PArel was immediately available to bind to the host cell receptor, thus avoiding any detrimental effects of accumulated acidic degradation products on the PArel. This is an important difference between these studies and the stability studies described in Figures 2–4 in which the PArel was accumulated over several days in an aqueous environment before assessing structural and antigenic integrity, and biological activity. After one week of storage at 40°C, the biological activity of PA adsorbed to alum and PA stored in PBS buffer was undetectable (Supplementary Figures 2A and B). Following two weeks and four months of storage at 25°C, PA adsorbed to alum and PBS retained only half of its original bioactivity respectively. After one month at 25°C, PA adsorbed to alum lost all of its bioactivity (Supplementary Figure 2A). After two months of storage at 40°C, PA encapsulated into 20:80 and 50:50 CPTEG:CPH or 20:80 CPH:SA nanoparticles retained from 60 to 90% of its biological activity (Supplementary Figure 1). After four months of storage, PA encapsulated into 20:80 CPTEG:CPH and CPH:SA retained a significant portion of its bioactivity (Figure 5). PA adsorbed to alum lost all its bioactivity after two months of storage at all temperatures except −20°C (Figure 5 and Supplementary Figure 2A). When stored as a dry powder, 20:80 CPH:SA nanoparticles were capable of preserving nearly 100% of the bioactivity of PA at all temperatures studied for 4 months. In contrast, dry storage of 50:50 CPH:SA nanoparticles was ineffective at stabilizing PA under most of the storage conditions tested. It is important to note that all nanoparticle chemistries were successful in preserving PA bioactivity when stored at −20°C and could be viable vaccine carriers when cold storage is available.

Discussion

Next-generation anthrax vaccines need to possess several attributes, including storage stability and improved immunogenicity to reduce the need for multiple booster injections. In this work, we have demonstrated that polyanhydride nanoparticle-based delivery systems can stabilize PA and enhance the shelf life of PA-based vaccines. In particular, these studies have shown that amphiphilic polyanhydride nanoparticles provide controlled antigen release over several weeks, the release of immunogenic protein capable of inducing high titer, neutralizing antibody, and extended shelf life storage at tropical and/or desert temperatures (e.g., 40°C).

Long term or repeated antigen exposure (depot effect) is essential for the development of immunological memory, which is a part of the rationale for the current immunization regimen for the AVA vaccine (i.e., five immunizations with yearly boosters)3. In the event of a bioterrorism attack, this protracted vaccination strategy may not be viable for treating a naïve population. A need therefore exists for a vaccine capable of inducing levels of neutralizing anti-PA antibody similar to that induced by the current immunization regimen, but in a single dose, that will also improve patient compliance. The nanoparticle-based
delivery system described in this work demonstrated sustained release kinetics for over two months, dictated by polymer chemistry (Figure 1A), thereby offering an improved method for delivery of biowarfare vaccines.

Although controlled release kinetics enable long-term antigen exposure, it is also essential that the antigen remain stable during fabrication and long-term storage to maintain its immunogenicity. Alum, an adjuvant that is currently approved for human use in the U.S., is known to produce an acidic bulk-environment that can be detrimental for pH-sensitive proteins. While the adsorbed antigen would experience a less acidic microenvironment on the surface of alum, many pH-sensitive protein immunogens in the bulk would be denatured and rendered ineffective by the acidic conditions (pH~5.6). PA is a highly labile protein that is not only sensitive to acidic environments, but also to increased temperature - an effect that is exacerbated as the conditions become more acidic. In this work, we demonstrated the superior PA stabilization capabilities of a nanoparticle-based delivery system that preserved the biological activity of PA during fabrication, delivery, and storage/thermal stability, thereby overcoming several of the problems associated with the use of alum-based vaccines.

Because of the absence of water in the process, nanoparticle fabrication conditions (solvent exposure, sonication, and vacuum drying) had no detectable effect on the structure, antigenicity, or activity of PA. This demonstrates the utility of the polyanhydride nanoparticle fabrication process for labile proteins, including PA, and is in agreement with similar results for other proteins exposed to these conditions. However, low pH and elevated temperature have been reported to have more detrimental impact on the structural stability and bioactivity of PA. As shown in Figure 1B, the pH, due to CPTEG:CPH nanoparticle erosion, only decreased to approximately 7.0 and was maintained at that value for over 30 days. In contrast, the erosion of CPH:SA nanoparticles decreased the pH to 5.2, which was similar to pH changes caused by alum. The acidic micro-environment caused by degradation of the CPH:SA nanoparticles correlated with the losses in structure (Figure 2), antigenicity and bioactivity (Figure 3) of PA released from these nanoparticles. The PA released from CPH:SA nanoparticle formulations also failed to prime an immune response capable of producing high titer, neutralizing serum antibody (Figure 4). Detailed structural analysis demonstrated that significant alterations in the secondary and tertiary structure of the PA were important determinants associated with the loss in immunogenicity, bioactivity, and quality (i.e., avidity) of the antibody response (Figure 4). Consistent with these results, it has been reported that thermally induced ellipticity changes in PA occur in acidic environments. In addition, at low pH and high temperature conditions, PA is structurally modified to a molten globular state. As observed in this work, this loss of structural integrity rendered PA biologically inactive and poorly immunogenic.

Amphiphilic CPTEG:CPH nanoparticles, specifically 50:50 CPTEG:CPH, provided an effective stabilizing environment for PA. While nearly 80% of PA antigenicity and activity was preserved upon release from 50:50 CPTEG:CPH nanoparticles (Figure 3), less antigenicity and activity were preserved (20% and 50%, respectively) upon release from 20:80 CPTEG:CPH nanoparticles (Figure 3). From the detailed structural analysis, it is likely that the alterations in the secondary and tertiary structure of PA released from the 20:80 and 50:50 CPTEG:CPH nanoparticles (Figure 2C) may be responsible for the loss in antigenicity and bioactivity (Figure 3). These alterations may result from the slight decrease in pH at 37°C, consistent with what was reported by Jiang et al., which occurred as the acidic degradation products built up over the duration of the in vitro stability study. However, these polyanhydride chemistries did not prove to be detrimental to the immunogenicity of PA. The total and neutralizing antibody titers as well as the antibody
avidity for mice immunized with PA released from 20:80 or 50:50 CPTEG:CPH nanoparticles were statistically indistinguishable from those induced by native PA (Figure 4). Thus, even in such “worst case” conditions (i.e., buildup of an acidic micro-environment in a microcentrifuge tube), the amphiphilic polyanhydride chemistries were not deleterious to PA.

One of the most difficult challenges to overcome in vaccine design and development is long-term storage and maintenance of bioactivity. This is an immediate concern in developing countries with limited infrastructure for cold storage. In this work, we tested the thermal stability of PA-loaded nanoparticles, PA adsorbed to alum, and PA in PBS for four months over a 60-degree temperature range (−20 to 40°C) while under dry storage. The changes observed in PA stored with alum are hypothesized to be a result of the acidic pH environment combined with the storage temperature. In contrast, PA stored in PBS buffer at a neutral pH is only likely to be affected by storage temperature. After two months of dry storage, all polyanhydride chemistries, except for 50:50 CPH:SA, outperformed the alum group in preserving the stability of PA at 40, 25, and 4°C (Supplementary Figure 1). After four months of dry storage, PA released from 20:80 CPTEG:CPH and CPH:SA nanoparticles retained its bioactivity (Figure 5). In these studies, it is important to note that dry formulations were evaluated while the studies described in Figures 2–4 tested PA-loaded nanoparticles that were incubated in an aqueous environment (to obtain the PA_rel). The dry storage conditions largely eliminate the build-up of acidic degradation products. This may explain why the 20:80 CPH:SA nanoparticles preserved PA bioactivity under these conditions.

Furthermore, these results indicate that the thermal properties of the copolymers, mainly the glass transition temperature (T_g), may play an integral role in particle shelf life over prolonged periods at elevated temperatures. Nanoparticles with T_g values close to or lower than the storage temperature may lose their particle structure due to their transition to a rubbery phase. The T_g values of 20:80 CPTEG:CPH, 50:50 CPTEG:CPH, 20:80 CPH:SA, and 50:50 CPH:SA have been reported to be 18, 8, 50, and 11°C, respectively. This may explain why the polymers with the two lowest T_g values (i.e., 50:50 CPTEG:CPH and 50:50 CPH:SA) are unable to release bioactive PA upon extended storage at 25 or 40°C. In addition, the hydrophobic 50:50 CPH:SA chemistry may induce non-covalent hydrophobic interactions with PA that may lead to aggregation and loss of bioactivity. The 20:80 CPTEG:CPH and 20:80 CPH:SA nanoparticles revealed similar abilities to preserve PA at 40 and 25°C for 4 months, demonstrating enhanced nanoparticle thermal stability and subsequent protein stabilization similar to lyophilized PA stored with stabilizers after only one month. While protein released from the 20:80 CPH:SA nanoparticles was not able to induce a robust antibody response following immunization, these formulations provided a suitable dry storage formulation at elevated temperatures. In contrast, the amphiphilic 20:80 CPTEG:CPH nanoparticles were able to induce robust antibody responses following immunization and provide for the preservation of a labile protein during long-term storage at tropical temperatures. The preliminary studies described herein have demonstrated a novel strategy for the design and development of a next-generation anthrax vaccine formulation with long-term stable storage capabilities.

Conclusions

Together, the studies reported herein demonstrate the potential use of a polyanhydride nanoparticle platform in next-generation vaccines employing recombinant proteins. These polyanhydride nanoparticles provide the ability to control PA antigen release, which would provide an in vivo antigen depot capable of long-term antigen presentation. Additionally, amphiphilic polyanhydride nanoparticles demonstrated superior thermal shelf life stability.
and the ability to withstand “worst case” conditions, thus making them capable of preserving PA bioactivity upon fabrication and storage at elevated temperatures. Of the four formulations tested, the amphiphilic 20:80 CPTEG:CPH nanoparticles demonstrated the best combination of characteristics to facilitate the stabilization of labile proteins, such as PA, making them excellent candidates for next-generation single-dose anthrax vaccines.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This material is based upon work supported by the National Institutes of Health (R03 AI076855-01A1). The LF and PA were obtained through the NIH Biodefense and Emerging Infections Research Resources Repository, NIAID, NIH.

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Figure 1. PA was released from polyanhydride nanoparticles in a chemistry dependent manner. A) Release of PA from polyanhydride nanoparticles over 60 days and B) the change in pH of PBS buffer (pH 7.6, 0.1 M) from nanoparticle degradation over 30 days. Error bars represent standard deviation of three replicates performed in three independent experiments.
Figure 2.
50:50 CPTEG:CPH best preserved the structural integrity of PA while CPH:SA nanoparticle chemistries cause a loss in secondary and tertiary structural integrity. A) Secondary structure as measured by circular dichroism (CD) and B) tertiary structure as measured by fluorescence spectroscopy after two days of release from polyanhydride nanoparticles.
Figure 3.
CPTEG:CPH nanoparticle chemistries released biologically active and antigenic PA. A) Residual antigenicity and B) residual activity of PA after two days of release from polyanhydride nanoparticles. The direct ELISA was performed with 0.5 μg/mL PA_{rel} from the nanoparticles. The biological assay was performed by incubating 6.4 μg/mL PA_{rel} from polyanhydride nanoparticles with 0.3 μg/mL LF with RAW cells for 6 h. Error bars represent standard deviation of 3 replicates. Treatments with different letters are significantly different from one another at p < 0.05.
Figure 4.
Mice immunized with PA released from 50:50 and 20:80 CPTEG:CPH, adjuvanted with alum, developed the highest antibody titers, the most avid antibody response, and highest neutralizing antibody titers. A) Antibody titers, B) antibody avidity (ND (not determined) for 20:80 and 50:50 CPH:SA because antibody titers were too low), and C) antibody neutralization titers of serum samples from immunized mice. Data is presented as the mean ± SEM of three replicates. All serum samples were evaluated individually. Treatments with different letters are significantly different from one another at p < 0.0001.
Figure 5.
Dry powder formulations of PA-loaded 20:80 CPTEG:CPH and CPH:SA polyanhydride nanoparticles were capable of preserving the activity of PA for four months when stored at 40, 25, 4, and −20°C. The biological assay was performed by incubating 125 μg/mL nanoparticles with 0.3 μg/mL LF with RAW cells for 6 h. Error bars represent standard deviation of 3 replicates. Treatments with different letters are significantly different from one another at p < 0.05.