Reduction Sensitive PEG Hydrogels for Codelivery of Antigen and Adjuvant To Induce Potent CTLs

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Supporting Information

ABSTRACT: Educating our immune system via vaccination is an attractive approach to combat infectious diseases. Eliciting antigen specific cytotoxic T cells (CTLs), CD8+ effector T cells, is essential in controlling intracellular infectious diseases such as influenza (Flu), tuberculosis (TB), hepatitis, and HIV/AIDS, as well as tumors. However, vaccination utilizing subunit peptides to elicit a potent CD8+ T cell response with antigenic peptides is typically ineffective due to poor immunogenicity. Here we have engineered a reduction sensitive nanoparticle (NP) based subunit vaccine for intracellular delivery of an antigenic peptide and immunostimulatory adjuvant. We have co-conjugated an antigenic peptide (ovalbumin-derived CTL epitope [OVA257−264: SIINFEKL]) and an immunostimulatory adjuvant (CpG ODNs, TLR9 agonist) to PEG hydrogel NPs via a reduction sensitive linker. Bone-marrow derived dendritic cells (BMDCs) treated with the SIINFEKL conjugated NPs efficiently cross-presented the antigenic peptide via MHC-I surface receptor and induced proliferation of OT-I T cells. CpG ODN-conjugated NPs induced maturation of BMDCs as evidenced by the overexpression of CD80 and CD40 costimulatory receptors. Moreover, codelivery of NP conjugated SIINFEKL and CpG ODN significantly increased the frequency of IFN-γ producing CD8+ effector T cells in mice (∼6-fold improvement over soluble antigen and adjuvant). Furthermore, the NP subunit vaccine-induced effector T cells were able to kill up to 90% of the adoptively transferred antigenic peptide-loaded target cell. These results demonstrate that the reduction sensitive NP subunit vaccine elicits a potent CTL response and provide compelling evidence that this approach could be utilized to engineer particulate vaccines to deliver tumor or pathogen associated antigenic peptides to harness the immune system to fight against cancer and infectious diseases.

KEYWORDS: nanoparticles, cancer vaccine, peptide vaccine, disulfide linkers, cytotoxic T cells (CTL)

1. INTRODUCTION

Vaccines are one of the major discoveries in modern medicine. Contributions include the almost complete elimination of polio, eradication of small pox, and a decrease by more than 95% of the incidence of diseases such as diphtheria, tetanus, pertussis, measles, mumps, and rubella.1 Vaccines have substantially decreased morbidity and mortality related to infectious disease and increased the average life span in the twenty-first century.

Despite these successes, there is a clear need to develop vaccines

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against pathogens like human immunodeficiency virus (HIV) and hepatitis C virus (HCV) and diseases such as malaria, tuberculosis, and cancer. Effective, traditional vaccines utilize live or attenuated pathogens which pose safety concerns due to the administration of unnecessary components of pathogenic microorganisms.2

New-generation subunit vaccines offer a safer and more specific approach to generate immunity, in which very specific components of pathogenic organisms (e.g., bacterial coat proteins, peptides, carbohydrates, or lipids; immunogenic determinants) are administered to protect against disease. Clinical success of subunit vaccines includes the use of Fluvirin (trivalent subunit protein) and Influvac (inactive purified surface antigens from influenza virus) against influenza, BioThrax (AVA-anthrax adsorbed vaccine, contains no whole cell or live bacteria) against anthrax, and Cervarix (contains a mixture of human papilloma virus protein antigens with adjuvants alum and monophosphoryl lipid-A) against cervical cancer caused by HPV.7 Because of our improved understanding of the immune system, more specific and safer peptide epitopes are being designed based on recombinant technology and epitope focusing.9

Peptide antigens are easily synthesized, stored, and transported. Specific antigenic peptide epitopes can induce an antigen specific cytotoxic T cell (CTL) response, which is of the utmost importance in the elimination of intracellular pathogens, as well as cancer cells. Successful intracellular delivery of peptides to professional APCs (mostly dendritic cells [DCs]) and their cross-presentation to T cells elicits CTLs. For cross-presentation, APCs can process endocytosed antigens by either the classical (also cytosolic pathway) or vacuolar pathway. In the classical pathway, protein antigens are processed into 8 to 12 amino acid fragments by proteosomal degradation machinery in cytosol and loaded onto MHC-I molecules in the endoplasmic reticulum (ER) or phagosome by antigen processing machinery (APM). In the vacuolar pathway, degradation of pathogens/antigens and loading of peptide fragments to MHC-I occurs in the endosome.5 However, soluble peptides suffer from fast degradation half-life (from a few seconds to minutes depending on their length)6 and low cellular uptake, resulting in poor cross-presentation by DCs and lower immunogenicity.

Many particulate subunit vaccine delivery systems such as liposomes, nanobeads, solid-lipid nanoparticles, polymeric nanoparticle and micelles, etc. have been investigated as a delivery vehicle to induce protection against HIV, influenza, and cancer. Particulate carrier systems can improve immunogenicity of antigens and adjuvants by mimicking the size, shape, and/or surface molecule organization of pathogens, in order to facilitate uptake by APCs.8 Additionally they prevent enzymatic degradation of antigens/adjuvants, and facilitate their intracellular delivery by increasing their local resident time.9 Depending on their degradation mechanism, the antigen can be released into the late endosome/lysosome or into cytosol. By optimizing the polymeric material of the nanoparticulate carrier, release of antigen can be triggered by changes in an intracellular pH,10,11 slower degradation rate of biodegradable polymeric carrier,12−14 or enzyme mediated release of antigen such as endolysosomal lipases15 or α-chymotrypsin.16

Efficient activation of DCs as well as induction and proliferation of T cells also requires costimulatory signals which can be provided by delivering adjuvants.7 Codelivery of antigens with adjuvants using particulate carriers can further boost the immune response.13,19 Many immune stimulating agents such as alum, oil in water emulsions, various TLR/NLR agonists, etc. are being explored in clinical as well as preclinical studies as vaccine adjuvants.20 TLR agonists such as monophosphoryl lipid-A (MPL-A), CpG, resiquimod, poly I:C, etc. have been investigated in the development of particulate vaccines.4 Antigen and adjuvant can be encapsulated, adsorbed, or conjugated to NPs. Sarti et al. demonstrated induction of IgA titers via codelivery of encapsulated MPL-A and ovalbumin (OVA) through PLGA (poly(lactic-co-glycolic acid) particles.21 De Titta et al. has shown generation of antigen specific CD8+ effector cells as well as memory cells in mice by delivering OVA protein and CpG conjugated particles via disulfide bond.22 Reversible cleavage and release of vaccine component can be made possible intracellularly by reduction of disulfide bond. Many others have also utilized disulfide linkers to design endosomal releasing nanovaccine carriers to induce in vivo CTL response.23−26

In this study, we have designed a PEG (polyethylene glycol) based PRINT (particle replication in nonwetting template) NP subunit vaccine to deliver a MHC-I epitope (SIINFEKL) of ovalbumin and a TLR-9 agonist, CpG oligonucleotide (ODN). PRINT offers the ability to mold biocompatible nanoparticles with complete control over particle size, shape, and chemical composition in a manner heretofore not possible with other particle technologies. High aspect ratio (80 × 80 × 320 nm, aspect ratio = 4) particles were chosen since the rod shape emulates known pathogens26 and increased aspect ratio is known to enhance cellular uptake.28 The antigenic peptide (CSINFEKL) and adjuvant (CpG ODN) were surface conjugated to NPs through reduction sensitive linkers, taking advantage of the intracellular reducing environment to trigger their release. Two cleavable linkers were investigated, a short SPDP (succinimidyl 3-(2-pyridyldithio)propionate) linker, and long NHS-PEG(2k)−OPSS linker. SPDP has been widely used to conjugate amine groups to thiol groups for intracellular delivery of siRNA−polymer conjugates,29,30 aptamer toxins, aptamer virus capsid conjugates,31,32 delivery of siRNA or DNA33−36 or delivery of model antigen ovalbumin via NPs.37−39 Though the NHS-PEG−OPSS linker has not been used as extensively in the literature, there is evidence to support the hypothesis that linker length is important. For example, Chen et al. showed higher immunostimulatory response with gold labeled CpG ODN by increasing the proximity of CpG ODN to gold nanoparticles.40 Furthermore, Singh et al. were able to demonstrate the effect of linker length on gene silencing efficiency of siRNA conjugated to quantum dots.41 We therefore hypothesized that linker length would play a significant role in the accessibility of biomolecules to their appropriate receptors. Our work demonstrates the significant role that linker length plays in the development of particulate based subunit vaccines. Herein we report the formulation of highly uniform and monodisperse hydrogel PRINT NPs co-conjugated with SIINFEKL and CpG, that are successfully taken up and processed by BMDCs, resulting in their efficient maturation and leading to SIINFEKL cross-presentation and subsequent induction of potent antigen-specific T cell proliferation and cytotoxic activity.

2. MATERIALS AND METHODS

2.1. Materials. Poly(ethylene glycol) diacrylate (M₄₇00), (PEG₇₀₀₀Da), 2-aminoethyl methacrylate hydrochloride (AEM), diphenyl (2,4,6-trimethylbenzoyl)-phosphine oxide (TPO), thiol modified CpG 1826 (C₆-S-S-C₆-tctagctgccttgacggt), dithiothreitol (DTT), sucrose, and DNase, RNase free sterile
water were purchased from Sigma-Aldrich. Tetraethylene glycol monoacrylate (HP₄A) was synthesized in house. Cysteine modified OVA₂₅₇₋₂₆₄ (CSINFEKL) were purchased from Peptide 2.0. Trifluoroacetic acid, methanol, dimethyl sulfoxide (DMSO), PTFE (polytetrafluoroethylene) syringe filters (13 mm membrane, 0.22-μm pore size), HPLC grade water, and acetonitrile were obtained from Fisher Scientific. Conventional filters (2 μm) were purchased from Agilent Technologies, and poly(vinyl alcohol) (Mw 2000) (PVOH) was purchased from Acros Organics. N-Succinimidyl 3-(2-pyridyldithio)propionate (SPDP) was purchased from Thermo Scientific. Ortho-pyridyl disulfide-PEG-N-hydroxysuccinimide ester (NHS-PEG(2k)-OPSS) was purchased from Creative PEGworks. PRINT molds (80 nm × 320 nm) were obtained from Liquida Technologies. DNA grade NAP-10 columns were purchased from GE Healthcare. RPMI1640 medium, penicillin and streptomycin, t-glutamine, and fetal bovine serum (FBS) were all from Life Technologies.

2.2. Methods. 2.2.1. PRINT Nanoparticle Fabrication. The PRINT particle fabrication process is described previously. Briefly, the preparticle solution was prepared by dissolving 3.5 weight percent (wt /wt) of the various reactive monomers in isopropyl alcohol (IPA). The reactive monomers included a cure-site monomer (an oligomeric PEG with a nominal molar mass of 700 g/mol terminally functionalized on both end groups with an acryl or acryloxy functionality); a hydrophilic monomer used to make up the majority of the particle composition (tetraethylene glycol monoacrylate, HP₄A); an amine containing monomer (aminoethyll methacrylate, AEM) which provides a chemical handle to conjugate various linkers and peptides; and a photoinitiator, TPO. Preparticle solution was composed of 69 wt %HP₄A, 10 wt % PEG₇₀₀DA, 20 wt % AEM, and 1 wt % TPO. Using a #3 Mayer rod, a thin film of preparticle solution was drawn onto corona-treated PET using roll-to-roll lab line (Liquida Technologies) running at 12 feet per minute. The solvent was evaporated by heat guns. Then an 80 × 80 × 320 nm, cylinder shaped mold was laminated to delivery sheet and passed through nip (80 psi, 12 feet per minute). After delamination, filled mold were cured by passing through a UV LED lamp (λmax = 395 nm, 30 psi of N₂, 12 feet per minute; Phoseon). Due to UV light initiated radical chain polymerization, monomers cross-linked into polymers to form a hydrogel. After cross-linking the hydrogels inside the mold cavities, the filled mold was laminated against a PVA harvesting sheet and passed through a heated nip (140 °C, 80 psi, 12 feet per minute). Particles were removed from the mold by splitting the harvesting sheet from the mold. Particles were then harvested by dissolving a sacrificial harvesting layer of PVA into water (2 mL of water per 10 feet of harvesting sheet). Particle suspensions were passed through 2 μm filter to remove additional scum layer. To remove excess PVA, particles were spun down at 14,000 rpm (Eppendorf Thermomixer R) for 25 min, and resuspended into sterile water. This purification procedure was repeated 3 times.

2.2.2. Thermogravimetric Analysis. Concentrations of particles were determined by thermogravimetric analysis (TGA) using TA Instrument’s Discovery TGA. Aluminum sample pans were tared before loading the sample. 20 μL of the stock nanoparticle solution was loaded onto the pan. Samples suspended in water were heated at 30 °C/min to 130 °C, followed by a 10 min isotherm at 130 °C. All samples were then cooled at 30 °C/min to 30 °C, followed by a 2 min isotherm at 30 °C.

2.2.3. Scanning Electron Microscopy. Particles were visualized by scanning electron microscopy (SEM) using a Hitachi S-4700 SEM. Prior to imaging, SEM samples were coated with 1.5 nm of gold–palladium alloy using a Cressington 108 auto sputer coater.

2.2.4. Dynamic Light Scattering. Particle size and zeta potential (ZP) were measured in sterile water by dynamic light scattering (DLS) on a Zetasizer Nano ZS (Malvern Instruments, Ltd.).

2.2.5. Conjugation of Linker to NPs. We utilized amine groups from AEM to conjugate cleavable linkers, SPDP and NHS-PEG(2k)OPSS. Theoretical numbers of −NH₂ groups contributed from 1 mg of nanoparticle suspension were calculated. Different molar ratios of linker to amine groups (such as 0.28, 0.55, and 2.30) were evaluated for the conjugation scheme (Table S1). For the optimized reaction scheme, particles were reacted with a 0.55 molar excess of linker, resulting in reacting 1 mg of NPs with 0.24 mg of SPDP or 6 mg of NHS-PEG-OPSS in 1 mL of 1x PBS + 0.1% PVA for 2 h. SPDP (or NHS-PEG(2k)OPSS) was dissolved in DMF. The volume of DMF was kept constant at 160 μL regardless of different linker mass used. Total volume of reaction was 1 mL. NPs were continuously agitated for 2 h at 1400 rpm (Eppendorf Thermomixer R). After 2 h, unconjugated linker was removed from the pyridyl dithiol decorated NPs via 2 centrifugation washes (Eppendorf Centrifuge 5417g) with sterile water. Efficiency of linker conjugation was evaluated by incubating 1 mg of NPs in 1 mL of DTT solution. Moles of DTT used was 10 times higher than the moles of linkers added during the conjugation procedure. Due to disulfide exchange reaction, pyridine-2-thione is released from pyridyl dithiol modified NPs, which can be detected by reading absorbance at 343 nm using Spectra Max M plate reader. Conjugation efficiency was determined by the following formula:

\[
\% \text{ conjugation efficiency} = \frac{\text{amount of linker conjugated}}{\text{amount of linker charged}} \times 100
\]

Conjugation efficiency of linkers was between 50% and 60%.

2.2.6. Conjugation of CSINFEKL to NPs. Once NPs were modified with SPDP or NHS-PEG(2k)OPSS, they were spun down for 25 min at 14,000 rpm (Eppendorf Centrifuge 5417g) and resuspended into 800 μL of sterile water. 1 mg/mL of cysteine labeled SIINFEKL (CSINFEKL) solution was made in sterile water. 200 μL of this solution was mixed with 800 μL of NP suspension and incubated overnight. NPs were continuously agitated at 1400 rpm (Eppendorf Thermomixer R). The following day, NPs were spun down for 25 min at 14,000 rpm (Eppendorf Centrifuge 5417g), supernatant was collected, and NPs were resuspended in 1 mL of sterile water. To remove unbound peptide, NPs were washed in sterile water 2 times. Due to disulfide exchange with cysteine labeled peptide, pyridine-2-thione is released from pyridyl dithiol modified NPs. By reading the absorption of pyridine-2-thione in the supernatant at 343 nm using the Spectra Max M plate reader, the amount of pyridine-2-thione released from NPs can be calculated and thus the amount of peptide conjugated was evaluated. Conjugation efficiency of peptide was evaluated as mentioned by eq 1. We found 60–70% conjugation efficiency of peptide to NPs.

2.2.7. Reduction and Purification of C6-S-S-C6 CpG 1826. CpG 1826 with phosphorothioate backbone was chosen because of slower in vivo degradation via nucleases as compared to CpG
with phosphodiester. C6-S-S-C6 CpG 1826 was reduced with 100 mM DTT solution in sodium phosphate buffer of pH 8.0 and purified by gel filtration chromatography using Sephadex NAP-10 column. Unreduced CpG was kept for an hour in the presence of 100 mM DTT solution in sodium phosphate buffer of pH 8.0. Sephadex NAP-10 column (DNA grade) was equilibrated by flowing through 15 mL of sterile water (DNase, RNase free). After equilibration, 0.75 mL of sample was loaded to column and allowed to pass through completely. Reduced CpG was eluted by passing through 1.2 mL of water. Concentration of CpG was measured by evaluating absorption at 260 nm by using a NanoDrop 2000 spectrophotometer.

2.2.8. Conjugation of Thiol-CpG 1826 to NPs. Once NPs were modified with SPDP or NHS-PEG(2k)OPSS, NPs were spun down for 25 min at 14,000 rpm (Eppendorf Centrifuge 5417g) and resuspended in 800 μL of sterile water. 40 μg of thiol CpG 1826 was mixed with 1 mg of linker modified NPs and kept for 8 to 10 h. Total volume of reaction was 1 mL. NPs were continuously agitated at 1400 rpm (Eppendorf Thermomixer R). Then NPs were spun down for 25 min at 14,000 rpm (Eppendorf Centrifuge 5417g) and resuspended in 200 μL of 100 mM DTT solution in sodium phosphate buffer. This washing procedure was repeated 3 times to remove adsorbed CpG. To evaluate conjugation of CpG, NPs were incubated in 100 mM DTT for 4 h. NPs were spun down, and supernatant was collected. Conjugation efficiency of CpG was evaluated as mentioned by eq 1. Evaluation of CpG was done by reading absorption at 260 nm using NanoDrop 2000 spectrophotometer. We found 70–80% conjugation efficiency of CpG to NPs.

2.2.9. Co-conjugation of CSINFEKL and CpG ODN to NPs. CSINFEKL peptide and CpG were co-conjugated in a 2 step process. For the co-conjugation process we used 0.39 mg of SPDP or 12 mg of NHS-PEG(2k)OPSS. After modification with linkers, NPs were conjugated to CSINFEKL by incubating 1 mg of NPs with 0.2 mg of CSINFEKL. Unbound peptides were removed by washing NPs in sterile water 2 times. After the final wash, nanoparticles were resuspended in 900 mL of sterile water and 0.04 mg of reduced CpG was added and left to react for 8–10 h at 1400 rpm (Eppendorf Thermomixer R). After co-conjugation, to evaluate peptide and adjuvant loading, CSINFEKL and CpG were cleaved from NPs by incubating 1 mg of NPs with 100 mM DTT for 4 h. After 4 h, nanoparticles were spun down at 14,000 rpm (Eppendorf Centrifuge 5417g) for 25 min, and supernatant was evaluated for released peptide and adjuvant through HPLC Agilent 1200 series and NanoDrop 2000 spectrophotometer, respectively. Co-conjugation of adjuvant resulted in a reduction of final peptide loading to 50% conjugation efficiency, while CpG loading remained at approximately 50–60% conjugation efficiency.

2.2.10. Peptide Evaluation via HPLC. Reverse phase high performance liquid chromatography (HPLC) was run on an Agilent 1200 series HPLC system using an Agilent C18 column. A Waters 2695 Alliance module equipped with quaternary pump, mobile phase degasser, temperature controlled auto sampler, and column thermostat were used for HPLC analysis. The separation was carried out on a Zorbax C18(2) column (150 mm × 4.6 mm i.d., 3 μm particle size, 100 Å pore size) from Phenomenex (Torrance, CA) at column temperature of 50 °C and sample temperature of 50 °C. Peptide was eluted using a mobile phase gradient, which consists of two solutions: solution A, water (0.1% TFA, trifluoroacetic acid), and solution B, acetonitrile with 0.1% TFA. The gradient method is shown in Table 1.

Binary linear gradients started from a mixture of 100% A and 0% B from 0 to 5 min. From 5 to 25 min B was increased gradually from 0% to 95%. Then, until 28 min the gradient was kept 5% for A and 95% for B. The mobile phase composition was then changed back to initial solvent mixture, and the column was equilibrated for 2 min before every subsequent run. The flow rate of the mobile phase was set to 2.0 mL/min. The samples for HPLC injections were prepared by cleaving off peptide in 100 mM DTT in sterile water. Peptides were detected via UV–vis detector at 210 nm, and concentration was determined by comparing area of peak with standard curve. Know amounts of peptides were dissolved with 100 mM DTT in sterile water to prepare a standard curve.

2.2.11. Animals. Female C57BL/6 mice and OT-I TCR transgenic mice were purchased from Jackson Laboratory and used at age 6–12 weeks. All experiments involving mice were carried out in accordance with an animal use protocol approved by the University of North Carolina Animal Care and Use Committee.

2.2.12. Preparation of Single Cell Culture from Mouse Spleens. Spleens were harvested from euthanized mice aseptically. Individual spleens were mechanically dissociated with the back of the sterile syringe plunger through a 100 μm cell strainer into RPMI 1640 medium. Red blood cells were lysed by ACK buffer. Cells were then resuspended in R-10 medium (RPMI 1640 medium supplemented with 10% FBS, 10 U/mL penicillin and 10 μg/mL streptomycin, 2 mM L-Glutamine) and filtered through a 70 μm cell strainer to remove any residual tissue fragments.

2.2.13. Preparation of BMDCs. Bone marrow was collected from mouse femurs and tibias as reported. Erythrocytes were lysed by ACK lysis buffer (Lonza). Bone marrow cells were then cultured at 2 × 10^6/mL in R10 supplemented with 50 μM 2-mercaptoethanol and 10 ng/mL each of IL-4 and granulocyte-macrophage-colony stimulating factor (GM-CSF). The culture was replenished with fresh medium on day 3 without removing old medium. On day 6 or 7, BMDCs were harvested and purified with Opti-Prep density medium (Sigma) to remove dead cells.

2.2.14. Antigen Presentation Assay in BMDCs. To evaluate the effect of linker length on antigen presentation, day 6 BMDCs (3 × 10^6 cells) were either untreated or treated with blank hydrogels (with and without linkers), CSINFEKL (5 μg/mL), CSINFEKL conjugated to PEG hydrogels via SPDP (short cleavable linker, 5 μg/mL), or CSINFEKL conjugated to PEG hydrogels via PEG(2k)OPSS (longer cleavable linker 5 μg/mL) for 4 h. After 4 h incubation, cells were washed with PBS of pH ~7.4 or citrate-phosphate buffer (pH 3.0) for 3 min on ice to strip off the MHC-I-peptide complex or NP-peptide/H-2Kb complex from cell surface. Additionally, cells were reincubated at 37 °C for an additional 20 h postwashes and then stained with CD11c-APC and 2S-D1.16-PE (anti-SIINFEKL/H-2Kb complex) antibodies (eBioscience), followed by flow cytometry analysis on Cyan ADP (Dako).

2.2.15. In Vitro T Cell Proliferation Assay. In vitro CD8+ T cell proliferation was done as previously reported. Briefly, day 6

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BMDCs were dosed with the samples described above for 24 h at 37 °C. OT-I CD8+ T cells were isolated from OT-1 mouse spleens using CD8α+ T cell Isolation Kit (Miltenyi Biotec) as per the manufacturer’s instructions and labeled with CFSE (5 μM) in PBS with 0.1% FBS for 10 min at 37 °C. BMDCs were then incubated with OT-I-CFSE cells in R-10 medium for 72 h at 37 °C. After incubation, cells were stained with CD8 and Vα2 antibodies. Division of OT-I cells as indicated by dilution of CFSE fluorescence in T cells was examined by flow cytometry.

2.2.1.6. Immunization Study. All formulations were prepared using low endotoxin grade reagents. To confirm the low endotoxin content of vaccines, formulations were routinely tested for endotoxin content using Pierce LAL Chromogenic Endotoxin Quantitation Kit following instructions. All formulations were prepared 24 h before injections, resuspended in an isotonic 9.25% sucrose, and subcutaneously administered in the right flank. SIINFEKL was given at a dose of 100 μg, and CpG was injected at a dose of 20 μg. On day 7, mice were sacrificed, and spleen and draining lymph nodes were harvested. All studies were repeated twice, and each experimental arm contained 4–6 mice.

2.2.17. ELISPOT Assay. Frequency of antigen specific IFN-γ producing T cells in spleen was evaluated using IFN-γ ELISPOT kit (BD Biosciences). Immobilon-P hydrophobic PVDF plates (Millipore) were briefly treated with 35% ethanol, washed 2 times with PBS, and coated overnight with anti-mouse IFN-γ antibody at 4 °C. The following day, plates were blocked with 200 μL of R-10 medium with for 2 h at room temperature (RT). 100,000 splenocytes in R-10 medium were plated in each well, with or without restimulation with 10 μg/mL of SIINFEKL peptide overnight at 37 °C. Spots were then developed following the manufacturer’s instructions.

2.2.18. In Vivo CTL Assay. In vivo CTL assay was performed as previously reported.47 Briefly, OT-1 T cells were isolated from spleen using CD8α+ T cell Isolation Kit (Miltenyi Biotec). C57Bl/6 mice were injected intravenously with 10,000 OT-I T cells on day −1. On day 0, mice were either untreated or immunized subcutaneously with a mixture of soluble SIINFEKL and CpG ODN, or NPs-SPDP-SIINFEKL-CpG, or NPs-PEG(2k)OPSS-SIINFEKL-CpG. Fifteen days after vaccination, mice were intravenously injected with 5 × 10^6 splenocytes as target cells. To prepare target cells, splenocytes from wild type C57Bl/6 mice were pulsed with SIINFEKL, peptide at 1 μg/mL in PBS for 1 h at 37°C, washed with PBS, and labeled with CFSE at 4 μM in PBS for 10 min at 37 °C (CFSEhi); or non-peptide pulsed cells were labeled with CFSE at 0.4 μM in PBS for 1 h at 37 °C (CFSElo). CFSEhi and CFSElo cells were mixed at 1:1 ratio to generate target cells for iv injections. On day 16, mice were euthanized, splenocytes were isolated, and cells were stained with anti-CD8 and Vα2 antibodies. Percentages of CFSEhi and CFSElo were determined with a flow cytometer. The ratio of unpulsed to pulsed target cells in the naive (unimmunized) mouse defined the 0% lysis level. The percent-specific lysis is determined by loss of the antigen-pulsed CFSEhi population compared with the unpulsed CFSElo control population using the formula [1 − (ratio in naive mouse/ratio in experimental mouse)] × 100.

3. RESULTS AND DISCUSSION

3.1. Conjugation of SIINFEKL and CpG to NPs. Pathogen inspired cylinder shaped (80 × 80 × 320 nm) cationic PEG hydrogel NPs were fabricated by the PRINT process. It has been established that particles with a net positive charge bind to the negatively charged plasma membrane of the cell surface, thus increasing cellular uptake, as well as enhancing endosomal escape of cargos.46 Our previous work demonstrated that cationic hydrogel particles were able to efficiently deliver siRNA to cytosol, resulting in efficient gene silencing.47 Furthermore,
Fromen et al. demonstrated significantly higher lung and systemic antibody titers when cationic particles were used to deliver ovalbumin, as compared to anionic particles. Therefore, amine groups on the NP surface were used to conjugate cysteine labeled SIINFEKL peptide via reduction sensitive, heterobifunctional linkers SPDP or NHS-PEG(2k)OPSS in a two-step process (Figure 1). First, the succinimidyl ester of the linker was reacted to the amine groups on the particle, forming an amide bond. The excess linker was then removed via centrifugation washes, followed by reacting the cysteine labeled adjuvant with the pyridine disulfide ring. Linker density was controlled by varying the linker-to-NP ratio (Table S1), in an effort to determine maximum linker density while maintaining the positive charge of the nanoparticles. Thus, during mon conjugation or dual conjugation of antigenic peptide and CpG, we aimed to keep the overall charge of NPs positive (as indicated by positive charge of the nanoparticles. Thus, during monodisperse nanoparticles with homogeneous distribution, size ranged from 250 to 280 depending on the surface modification.

3.2. In Vitro Antigen Presentation in BMDCs by Subunit Vaccine. In order for vaccines to generate an efficient CD8+ T cell response, DCs must first internalize antigens, process them into 8–12 amino acid peptides, and present them to T cell receptors as peptide/MHC-I complexes on the cell surface. T cells recognize MHC-I-peptide complex via T cell receptors and in the presence of other costimulatory signals subsequently proliferate. Since antigen presentation is key to developing an efficient T cell response, the antigen presentation efficiency of our PRINT NP subunit vaccine was evaluated using an in vitro assay. 25-D1.16, an antibody that recognizes SIINFEKL/H-2Kb on antigen presenting cells, was used to stain BMDCs treated with various samples to quantify antigen presentation. As expected, SIINFEKL peptide binds to H-2Kb on the cell surface directly upon 4 h pulsing, while treatment with blank NP-SPDP or NP-PEG(2k)OPSS and washes with PBS resulted in minimum staining for MHC-I-peptide complex (Figure 3A). Interestingly, cells treated with NP-SPDP-SIINFEKL or NP-PEG(2k)OPSS-SIINFEKL showed similar surface MHC-I-peptide staining to soluble peptide (Figure 3A and Supporting Figure 1I). When unbound peptide or NP-peptide was removed by washes with PBS, followed by another 20 h culture at 37 °C, it was observed that both NP-SPDP-SIINFEKL and NP-PEG(2k)OPSS-SIINFEKL led to significantly higher pMHC level than that of soluble peptide pulsed BMDCs (Figure 3B and Supporting Figure 1J). On the other hand, pMHC complexes on soluble peptide pulsed and washed BMDCs decreased over time, suggesting loss of pMHC (Figure 3E). These results indicate that NP conjugation of antigenic peptide increases the overall level of cell surface pMHC over time, possibly due to enhanced cell uptake of NP-peptides and more efficient delivery of antigenic peptide into the class I presentation pathway.

To further examine the capability of delivering antigenic peptide into MHC class I presentation pathway by NP-peptide formulations, cells were pulsed with soluble or NP conjugated peptides for 4 h, and washed with an acidic citrate-phosphate buffer (pH3.0) for 3 min to strip off the MHC-I peptide complex from the surface of the BMDCs. Cells were then incubated for another 20 h at 37 °C allowing for internalized antigens to be represented onto cell surface. As shown in Figures 3C and 3F, citrate-phosphate treatment completely removed SIINFEKL from pMHC complexes for cells treated with soluble SIINFEKL (down to 0.9%), but only partially removed cell bound NP-peptide (down to ~50%) (Supporting Figure 1K). This result may be due to the increased binding avidity of NP-peptide to cells as a result of the multivalence of peptide on the surface of NPs. As shown in Figure 3D, after acidic washes and an additional 20 h incubation, peptide/MHC I staining for NP-SPDP-SIINFEKL and NP-PEG(2k)OPSS-SIINFEKL pulsed cells were 92.7% and 77%, up from around 50% (Figure 3G), which was also significantly higher than soluble SIINFEKL (6%) (Supporting Figure 1L). Dur ing the reincubation period, cells treated with NP conjugated peptide had higher staining of pMHC complex, as compared to cells that did not go through a reincubation period (Figure 3G), which indicates higher uptake of NP and presentation of antigenic peptide. In conclusion, the conjugation of peptide to PRINT NPs via cleavable linkers is able to achieve not only higher but also longer lasting presentation of antigenic peptide by BMDCs as compared to soluble peptide. 3.3. In Vitro Proliferation of OT-I T Cells in BMDCs Treated with Subunit Vaccine. Next, we evaluated how cross presentation via BMDCs translated into activation and priming of CD8+ T cells. CD8+ T cells derived from OT-I TCR transgenic mice were used to evaluate OVA-specific T cell response for NP and soluble vaccine formulations. In this study, BMDCs, either untreated or treated with soluble SIINFEKL or

![Figure 2. Representative SEM image of 80 × 320 nm PRINT subunit peptide vaccine.](image-url)
NP-SIINFEKLNP conjugated peptides, were incubated with 5- (and 6-) carboxyfluorescein diacetate succinimidyl ester (CFSE) labeled OT-1 T cells for 3 days. Proliferation of T cells was evaluated by flow cytometry. A representative flow cytometry histogram for each group is shown in Supporting Figure 2. As shown in Figure 4, BMDCs pulsed with soluble peptide at a concentration of 5 μg/mL were able to induce proliferation of cognate CD8+ T cells. In comparison, much enhanced T cell proliferation was observed for BMDCs treated with NP conjugated peptide, which indicates higher priming and activation of CD8+ T cells. CSINFEKL conjugated to PRINT NPs via the longer cleavable linker resulted in approximately 95% of T cell growth, as compared to short cleavable linker (74.5%) and soluble peptide (32.6%). This is consistent with the enhanced antigen presentation by NP-peptide demonstrated above (Figure 4).

### 3.4. Maturation of BMDCs by CpG ODN Conjugated PEG Hydrogels.

It is well-known that activation of naive T cells and development of multiple effector functions depends not only on recognition of MHCI- peptide complex on APCs by TCRs but also on a second signal from engagement of costimulation receptors. Concurrent delivery of adjuvants has been shown to promote the second signals, further drive clonal expansion of naive T cells, and aid in their differentiation into armed effector T cells. The CD40 ligand and CD28 expressed on T cell surface bind to costimulatory molecules CD40 and CD80 on the surface of DCs, respectively. CD40 provides signals for activation, while CD80 provides signals for proliferation. The ability of PRINT NPs to deliver CpG ODN to provide costimulatory signals was tested in BMDCs. BMDCs were incubated with soluble CpG ODN or NP-CpG for 18 h. After incubation, cells were analyzed for expression of CD80 and CD40 via flow cytometry. Expressions of costimulatory molecules were presented as a ratio of mean fluorescent intensity (MFI) to untreated cells. Treatment with CpG conjugated PRINT NPs at 1 μg/mL or 10 μg/mL concentrations induced upregulation of CD40 and CD80 on BMDCs as compared to untreated cells, as potently as soluble CpG. Therefore, the functionality of CpG is well retained during conjugation to nanoparticles. No upregulation of either marker was found when BMDCs were treated with SIINFEKL conjugated NPs. Lack of response indicates the limitation of delivering antigen alone for maturation of BMDCs. CD40 and CD80 were both upregulated when CpG ODN were combined with antigenic CSINFEKL peptide either via co-conjugation of CpG ODN and SIINFEKL on a single NP or when delivered on separate NPs (NP-CpG + NP-peptide), indicating the helper effect of adjuvant in the presence of antigen (Figure 5A,B).

### 3.5. Induction of IFN-γ Producing SIINFEKL Specific CD8+ T Cells in Mice.

After evaluating the efficiency of antigen presentation and maturation of BMDCs as well as OT-I T cell proliferation, PRINT NP subunit vaccines were analyzed for their ability to induce IFN-γ producing antigen specific T cells in mice. Frequency of antigen-specific IFN-γ producing T cells in spleens of mice 7 days postimmunization was evaluated ex vivo by ELISPOT. First we evaluated whether co-conjugation of antigen and adjuvant are necessary by vaccinating mice with NP-SPD-CSIINFEKL + NP-SPDP-CpG and NP-SPDP-CSIINFEKL-CpG. We found that mice treated with NPs co-conjugated to antigenic peptide and CpG induced significantly higher CD8+ T cell response as compared to mice treated with separately conjugated NPs (Supporting Figure 3). Similar results were also reported by Schlosser et al. when they coencapsulated TLR ligands CpG or poly I:C and antigen in the same delivery system. Next we tested both the vaccine formulations for their ability to induce IFN-γ producing CD8+ T cells. As shown in Figure 6A, there is a significantly higher induction of IFN-γ producing CD8+ T cells when mice were immunized with NP-SPDP-CSIINFEKL-CpG and NP-PEG(2k)OPSS-CSIINFEKL-CpG as compared to mice receiving a mixture of soluble antigen and adjuvant. Co-conjugation of adjuvant and antigen resulted in 10 times higher induction of IFN-γ producing T cells as compared to NPs conjugated to antigen alone. IFN-γ producing T cell response was minimum in case of negative control-blank PRINT NPs.

Secretion of IFN-γ by antigen-specific CD8+ T cells into culture medium upon restimulation over 3 days was also examined by ELISA. Similar to T cell frequency analysis, the bulk production of IFN-γ was also significantly higher for mice treated with NP formulations with co-conjugated peptide and adjuvant as compared to NP conjugated with just antigen, or soluble antigen and adjuvant (Figure 6B), again confirming the importance of codelivery of CpG adjuvant for IFN-γ production.

### 3.6. In Vivo CTL Response in Mice after Immunization with SIINFEKL and CpG Co-conjugated NPs.

Induction of CTLs is critical in generating an antitumor immune response. We evaluated PRINT NP subunit vaccines for their ability to induce CTLs through an in vivo CTL assay. Fifteen days after immunizations with mixture of soluble SIINFEKL and soluble CpG or NP-SPDP-CSIINFEKL-CpG or NP-PEG(2k)OPSS-CSIINFEKL-CpG, mice were adoptively transferred with SIINFEKL pulsed, CSFE+ and control nonpulsed, CSFE+ target cells. On day 16, peptide pulsed and nonpulsed target cells in mouse spleens were differentiated by flow cytometry by two distinct population with high and low CFSE fluorescence (RS).
and R4, respectively in Figure 7A–D). CSFE<sup>hi</sup> population was decreased for peptide pulsed cells in mice immunized with a mixture of soluble peptide and soluble CpG (Figure 7B), NP-SPDP-CSIINFEKL-CpG (Figure 7C), and NP-PEG(2k)OPSS-CSIINFEKL-CpG (Figure 7D). Moreover, the percentage of OVA specific cell killing was 90% for mice treated with NP-PEG(2k)OPSS-CSIINFEKL-CpG, which is significantly higher than for mice treated with soluble antigen with soluble adjuvant (35%) and NP-SPDP-CSIINFEKL-CpG ODN (50%) (Figure 7E). Results from these assays reveal that the longer linker is more efficient at in vivo target cell killing as compared to short linker.

4. DISCUSSION

Here we have shown (i) successful co-conjugation via reduction sensitive linkers of model antigenic peptide-SIINFEKL and CpG
ODN to hydrogel NPs; (ii) delivery of CSIINFEKL via NPs to DCs that induced cross-presentation of SIINFEKL via MHC-I protein molecules and subsequently stimulated in vitro proliferation of OT-I T cells; (iii) delivery of CpG ODN via NPs provided “adjuvanted” effect by inducing maturation of DCs as demonstrated by upregulation of CD80 and CD40; (iv) codelivery of CSIINFEKL and CpG ODN induced IFN-γ producing robust CD8+ T cells; and (v) codelivery of CSIINFEKL and CpG ODN by reduction sensitive hydrogel system elicited potent CTLs that kill antigen specific target cells. Within the past decade PRINT NPs have evolved into a unique delivery platform for various agents such as chemotherapeutics, fluorescent dyes, quantum dots, siRNA, and proteins. Due to the versatility of the PRINT platform, it has been employed to study the impact of various particle parameters in a biological system such as biodistribution of intravenously administered NPs as well as lymphatic trafficking of particles after intramuscular administration. Utilizing immunologically inert materials to mimic size, shape, and surface functionallity of pathogens, PRINT provides an excellent platform to engineer subunit vaccines investigating various combinations of antigens and adjuvants to tune the immune response. Galloway et al. showed enhanced humoral response when mice were treated with trivalent influenza protein adsorbed cationic PLGA PRINT particles. In an effort to induce mucosal immune response, Fromen et al. showed higher antibody titers of ovalbumin protein when delivered via intranasal immunization by PRINT hydrogels. Building upon this platform, herein we have presented the first PRINT system to induce potent CTL response by codelivering antigenic peptides and CpG ODN through reduction sensitive linkers.

Disulfide linkers have been widely used in antibody drug conjugates (ADCs) to allow for release of chemotherapeutic drugs in intracellular reductive environment and delivering antigenic proteins. Swartz lab and Hubbell lab have utilized disulfide chemistry to deliver model protein OVA as well as TAA peptide, Trp-2 via nanoparticles for the induction of in vivo immune response. They have shown induction of potent effector and memory CD8+ T cell responses for the inhibition of tumor growth. Others have also shown higher humoral and cellular response by disulfide conjugated OVA via polymeric nanoparticles. One advantage of this technique is that triggered release of the cargo only happens upon cellular internalization, due to the presence of various reductive enzymes in the endosome, lysosome, and cytosol. Therefore, employing reduction sensitive linkers to conjugate CpG ODN and SIINFEKL to NPs, we were able to deliver CpG, resulting in stimulation of TLR-9 and the triggered release of costimulatory molecules to enhance T cell activation, while also delivering SIINFEKL, to be cross-presented to the surface of APCs (vacular pathway). More experimental studies are required to understand the mechanism of NP uptake, trafficking, and antigen release.

Employing peptide antigens has many advantages including cost-effective, enhanced safety, specificity, and stability. Short peptides for MHC-I/HLA binding domains can be readily synthesized in large quantities. Using defined peptide epitopes allows for the generation of very specific arms of effector T cells without the health risk of inducing autoimmune or adverse reactions which may occur in response to whole protein antigens. Moreover, peptides can be chemically modified to improve their solubility, stability, and antigenicity. In our studies, the cysteine modification added to the peptide for conjugation purposes did not interfere with MHC I binding to SIINFEKL. However, ensuring that peptide modification does not alter MHC binding will need to be investigated for each unique application to confirm that the appropriate antigen/peptide is being presented. Clinical application of peptide vaccine requires matching of patients’ HLA class I alleles to CTL epitopes to be able to induce T cell response. Multiple strategies, like multivalent peptide cocktail, hybrid peptides, and long peptides, have been adopted in recent clinical trials to enhance the applicability of peptide vaccines. Nanoparticle-based delivery platform with versatility in peptide association like ours would be able to help simultaneous delivery of multiple antigenic peptides.

Polyethylene glycol (PEG) is extensively used in biomedicine and cosmetics due to its biocompatibility. It is widely acknowledged that acrylate-derivatized PEG (PEGDA) hydrogels undergo slow degradation in vivo due to either hydrolysis of end group acrylate esters or oxidation of the ether backbone. Making PRINT nanoparticles with poly(ethylene glycol) diacrylate (PEGDA) allows us to tune many physicochemical parameters. Another important parameter of PRINT hydrogel vaccine design is the positive surface charge provided by AEM in the composition. Cationic formulations enhance NP cellular internalization and endosomal escape thereby delivering cargo in the reductive environment of cytosol. Being able to take advantage of these cellular events should favor antigen cross presentation through the classical pathway. Neumann et al. has also reported inflammation some activation by cationic charged particles alone, which may potentially provide adjuvanticity and contribute to the high potency of our particle vaccines. Additionally, positively charged particles remain trapped at the injection site because of

![Figure 4. T-cell proliferation assay. Peptide conjugated NPs induced higher T cell proliferation as compared to soluble peptide. Bone marrow derived dendritic cells (BMDCs) were incubated for 4 h with free peptide and nanoparticulate peptide conjugated via short and long cleavable linkers. Pulsed BMDCs were then incubated with CFSE-labeled OT-I CD8+ T cells for 3 days. T cells were surface stained, and proliferation of OT-1 T cells was assessed by flow cytometry. Percentage of T cell proliferation after treatment with different formulations is shown in Figure 5. Results are shown as mean ± SD, n = 3. Data were analyzed by one-way ANOVA, Bonferroni post hoc analysis, **p < 0.01, ***p < 0.001.](Image)
Figure 5. Upregulation of maturation markers by NP-CpG on BMDCs. BMDCs were treated with various samples for 24 h with 1, 4, or 10 μg/mL of CpG. Equivalent amounts of NPs were dosed in the case of NP-peptide. NP-SPDP-CSIINFEKL (1) has an equivalent dose of NPs to 1 μg/mL particulate CpG, and NP-SPDP-CSIINFEKL (2) has an equivalent dose of NPs to 10 μg/mL particulate CpG. The expression of costimulatory molecules was detected by flow cytometry analysis of fluorescence labeled CD40 and CD80 antibodies. Ratio of MFI from sample to MFI of untreated is plotted in panel A for CD40 and panel B for CD80. Results are shown here as mean ± SD, n = 3. Data were analyzed by one-way ANOVA, Bonferroni post hoc analysis, ***p < 0.001.

Figure 6. Induction of IFN-γ producing SIINFEKL specific CD8+ T cells in spleen. Mice were immunized with various samples containing 100 μg of peptide and/or 20 μg of CpG via sc injection in the left flank. One week later, splenocytes were isolated and restimulated with SIINFEKL for 20 h for IFN-γ ELISPOT assay (A), or for 72 h for ELISA analysis of secreted IFN-γ in medium (B). Results are shown as mean ± SD, n = 4. Data were analyzed by one-way ANOVA, Holm–Sidak’s multiple comparison test, *p < 0.05, ***p < 0.001.
collagen fibers and negatively charged proteins (glycosaminoglycans) of extracellular matrix (ECM). Our previous study suggested that 80 × 80 × 320 nm cationic particles have minimal lymphatic drainage. Thus, our NP vaccines most likely rely on the uptake by tissue resident DCs at the injection site for antigen presentation, such as migratory CD103+ DCs in peripheral tissues, a subset of DCs very potent in antigen cross presentation. These results demonstrate the ability to perform well-controlled mechanistic studies to investigate the effect of optimizing PRINT particulate vaccine parameters in terms of immune cell targeting, APC uptake/activation, and nanoparticle vaccine efficacy.

5. CONCLUSION

In summary, we have investigated the use of an engineered PEG hydrogel subunit vaccine to harness the power of our own immune system to generate CTLs to validate this approach in the fight against cancer. We have designed and developed a model platform NP subunit vaccine to codeliver antigenic peptide and CpG ODN. These NPs were successfully internalized and processed by BMDCs, resulting in BMDC maturation, subsequent cross-presentation of antigenic peptide, and induction of potent antigen-specific T cells. Taken together, results from this study provided a highly specific and effective platform to develop vaccines against infectious disease and cancer.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.molpharmaceut.6b00288.
(30) York, A. W.; Huang, F.; McCormick, C. L. Rational design of targeted cancer therapeutics through the multiconjugation of folate and...


(73) Zhan, X.; Tran, K. K.; Shen, H. Effect of the poly(ethylene glycol) (PEG) density on the access and uptake of particles by antigen-presenting cells (APCs) after subcutaneous administration. *Mol. Pharmaceutics* 2012, 9 (12), 3442−51.