Silylated Precision Particles for Controlled Release of Proteins

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Supporting Information

ABSTRACT: With the recent advances in the development of novel protein based therapeutics, controlled delivery of these biologics is an important area of research. Herein, we report the synthesis of microparticles from bovine serum albumin (BSA) as a model protein using Particle Replication in Non-wetting Templates (PRINT) with specific size and shape. These particles were functionalized at room temperature using multifunctional chlorosilane that cross-link the particles to render them to slowly-dissolving in aqueous media. Mass spectrometric study of the reaction products of diisopropyldichlorosilane with individual components of the particles revealed that they are capable of reacting and forming cross-links. Energy dispersive spectroscopy (EDS) and X-ray photoelectron spectroscopy (XPS) were also used to confirm the functionalization of the particles. Cross sectional analysis using focused ion beam (FIB) and EDS proved that the functionalization occurs throughout the bulk of the particles and is not just limited to the surface. Circular dichroism data confirmed that the fraction of BSA molecules released from the particles retains its secondary structure thereby indicating that the system can be used for delivering protein based formulations while controlling the dissolution kinetics.

KEYWORDS: silylation, protein delivery, controlled release, functionalization, bovine serum albumin, diisopropyldichlorosilane

INTRODUCTION

Controlled delivery of protein based formulations to the site of interest is of high practical importance, due to the development of novel biomolecules with therapeutic potential.1 With the recent advances in biotechnology, monoclonal antibodies (mAb) are also being used as protein based therapeutic agents for treating different diseases.2 However, low bioavailability of these molecules makes their systemic delivery through the oral route quite challenging. This can be attributed to large molecular weight and hydrophilicity of these macromolecules that makes their absorption through biological membranes difficult.3,4 The ability to protect the active compound from the harsh conditions of the gastrointestinal tract and enzymatic barrier are other challenges in oral delivery of proteins, since the delicate structure of these molecules is prone to different modes of instabilities.4,5 Due to less than 1% oral bioavailability of the initial dose, proteins are normally administered by either injection or infusion in aqueous solutions.4,6–8 Local delivery to a specific site using intratumoral,9,10 intravitreal,11 epidural,12 and intra-articular13 routes is desirable as it significantly reduces the therapeutic drug dose and adverse side effects associated with the undesirable distribution in various tissues and organs.14 Other noninvasive alternative routes suffer from the fact that, in the absence of a membrane permeation enhancer, the absorption is much less than that of parenteral administration while toxicity is a major drawback of using these enhancers in the formulation.14,15

According to Almeida and Souto,1 the established method of protein delivery consists of attaching the drugs to a proper particulate carrier, where the fate of the drug molecule is determined by the properties of the carrier rather than those of the protein. Therefore, use of an appropriate delivery vehicle that can protect proteins from enzymatic degradation and control the release rate is a promising approach for prolonged retention while saving the biological activity of the drug.16

There are several studies that address different systems for protein/antibody delivery. Lee and Yuk16 review both polymeric hydrogel carriers and liposomal systems for delivery...
of proteins. One of the obstacles in the development of a suitable polymeric carrier for protein drugs is low drug loading.10 A possible solution to this problem is preparation of particles that are made from a protein based carrier instead of a polymer. Several researchers have tried to prepare particulate systems that do not rely on a carrier polymer for delivering the protein. Yang et al.17 have developed methods to produce crystalline suspensions of approved therapeutic mAbs. Miller et al.18 studied highly concentrated suspensions of lysosome powder in a nonaqueous mixture of safflower oil and benzyl benzoate. α-Amylase crystals have been suspended in sucrose acetate isobutyrate plasticized with ethanol as a potential in situ formable protein depot.6

Preparation of cross-linked albumin nano/microparticles has been reported in the literature using suspension technology,19,20 layer-by-layer assembly,21,22 spray-drying,23 and desolvation/coacervation.24,25 These methods rely on heat treatment for self-cross-linking through formation of interchain amide links or chemical treatment with a stabilizer such as glutaraldehyde (GA) and butanedione to confer dimensional stability. GA reacts with amino groups of the protein while butanedione reacts with the guanidinyl moiety of arginine residue to form uncleavable chemical cross-links. In addition to the irreversible cross-linking reaction of these stabilizers, the shapes of the fabricated particles are limited to spheres, capsules, and rods. Bogdansky26 gives a detailed insight into the use of albumin based particles for therapeutic applications. The advantage of the albumin based particulate system is the lack of toxicity and degradation into natural products. Albumin based particles containing proteins have been examined as potential drug delivery systems by other researchers. Urokinase has been immobilized in GA cross-linked bovine serum albumin (BSA) particles by Bhargava and Ando,27 and GA cross-linked insulin-BSA microbeads were prepared by Goosen et al.28 Human serum albumin (HSA) nanocapsules were prepared by Rollett et al.29 for targeted drug delivery to activated macrophages. Application of albumin microspheres (3–4 μm) to pro-inflammatory cytokine inhibition was reviewed by Oettinger and D’Souza,30 and versatility of albumin as a drug carrier was evaluated by Kratz.31 Doshi and Mitragotri32 demonstrated that both size and shape of polymeric particles critically control their elimination by macrophages. Therefore, precise control over size and shape is desirable for controlling the clearance time of the particles upon local delivery.

According to Lee et al.,33 injectable beads prepared by cross-linking BSA under mild conditions using GA should yield a nonimmunogenic and biodegradable device for drug delivery and have been used for sustained release of progesterone. Furthermore, Ratcliffe et al.34 tested the in vivo biocompatibility of poly(lactic acid), poly(butyl cyanoacrylate), gelatin, and GA cross-linked homologous albumin microspheres (1.9 ± 1.5 μm) with rabbit’s synovial tissues upon intra-articular injection. They concluded that albumin microspheres are the most acceptable polymer among the ones studied. The slight hyperplasia observed by the authors in the synovial membrane was attributed to denaturation of albumin with GA.

Numerous pH cleavable linkers such as acetics, ketals, ortho esters,35,36 bis-pyridoxal polyphosphates,37 and the derivative of 2-methylmaleic anhydride38 have been developed for cross-linking proteins. Silyl ethers have also been successfully used for reversibly cross-linking precision particles made from hydroxyethyl acrylate and controlling the release kinetics based on the pH of the environment and size of the substituents on the silicon atom.39

The objective of this work was to fabricate suspensions of protein particles with controlled size and shape capable of slowly releasing the protein based therapeutic in aqueous media exploiting the pH cleavability of the silyl ether bond in cross-linked particles. This carrier system can be used for local delivery of the biomolecules to the site of interest while maintaining protein concentration within the therapeutic window. The outcome of this work demonstrates the ability to effectively increase the retention time of the protein based therapeutics deliverable through the parenteral route by fine-tuning the silylation reaction and can potentially maximize the efficacy of the drug.

**EXPERIMENTAL SECTION**

**Materials.** BSA, disopropylchlorosilane (DIDCS), and formic acid (FA) were obtained from Sigma-Aldrich. Extra-dry acetonitrile (ACN), trifluoroacetic acid (TFA), α-D-lactose monohydrate (lactose), glycerol, and extra-dry isopropanol (IPA) were all ACROS products obtained through Fisher Scientific. Standard buffers of pH 7.4 and 10.0, as well as HPLC grade water and acetonitrile, were all obtained through Fisher Scientific. The Alexa Fluor 555 conjugated BSA was purchased from Life Technologies.

**Particle Fabrication.** Particle Replication in Non-wetting Templates (PRINT) platform has been previously used to fabricate monodisperse, shape-specific albumin particles.40 In this work, PRINT was used to make particles from a mixture of BSA, lactose, and glycerol (40/40/20 wt %) in deionized water (18.2 MΩ-cm). The mixture was rolled and dried into the form of a solid flat sheet on the surface of a poly(ethylene terephthalate) film using Mayer rod on EZ Coater EC-200 (Chem Instruments) which is then used to fill the donut-shaped micron-sized molds. The filling process was carried out using a commercially available hot roll laminator (Chem Instruments) at 414 kPa and 138 °C as measured by Thermex Range C (TMC Hallcrest) temperature indicator films. Lactose was added to the formulation to lower the softening temperature of BSA and to make it moldable in the PRINT process. Glycerol merely serves as a processing aid during the filling stage. The particles were then transferred to a harvesting film made from Luvitec VA 64 (BASF) in a heated laminator (total lamination time of less than 30 s). Particles were freed from the harvesting film by dissolving the adhesive in extra-dry ACN. Due to the fact that the solubility of BSA in ACN is minimal (0.02 mg/mL),41 it was possible to separate these particles from the Luvitec layer through consecutive washing without any recognizable changes to the structural integrity of the particles. Furthermore, the secondary structure of the protein was expected to remain intact in the absence of water when using extra-dry ACN.42

**Particle Composition.** The composition of particles was determined using a high-performance liquid chromatography (HPLC) instrument (1260 Infinity by Agilent) equipped with an evaporative light scattering detector. The HPLC method used was based on separating BSA from lactose and glycerol using an Ultracel-30K Amicon Ultra centrifugal filter (Millipore) and then running them on two different columns based on reversed phase chromatography using a Poroshell 300SB-C8 column (Agilent) for the BSA in the mixture and ion exchange chromatography using a Hi-Plex Ca column (Agilent) for lactose and glycerol (Table S1, Supporting Information).

**Functionalization (Silylation).** In this work, silylation based on chlorosilanes was carried out on particles after harvesting them in ACN. The mass of the harvested particles was determined using thermogravimetric analysis (TGA Q5000-TA Instruments). Particles (5 mg) were dispersed in ACN (4.7 mL), and then DIDCS (1.27 mL) was added while allowing the reaction (rx) to proceed for 6, 12, and 24 hour (hr) in separate vials. At the end of the treatment with the chlorosilane, the particles were washed with IPA which quenched the reaction at the desired time point and enabled the removal of unreacted DIDCS. The silylation process on the particles and
introduction of a silyl group in place of the labile hydrogen atoms upon exposure to DIDCS result in the formation of cross-links (Figure S1, Supporting Information). The amino acid composition of BSA contains multiple residues with labile hydrogen atoms (e.g., Tyr, Ser) that can serve as an anchor point for attachment of diisopropyl groups. Therefore, it is expected that multiple residues participate in the reaction. The byproduct of the reaction is hydrochloric acid which is expected to behave as a weak acid in pure ACN (acid dissociation constant (pK_a) of 8.9).45

**Chemical Characterization. Mass Spectrometry (MS).** To understand the interaction of chlorosilane with the components of the particles, mass spectrometric analysis using electrospray ionization (ESI) was performed on the individual ingredients of the particles. BSA powder was pulverized in an agate pestle and mortar combination and passed through a 400 mesh sieve (38 μm nominal opening) to get a fine dry powder. This powder (20 mg) was then dispersed in extra-dry ACN (12 mL), and DIDCS (5 mL) was later added to the mixture. The silylation reaction was allowed to proceed for 12−24 hr and then neutralized with IPA at the end of the experiment. BSA crystals were then spun down and washed with extra IPA to remove unreacted and physisorbed DIDCS. Intact and silylated samples were all purified and isolated by centrifugal filtration of the aqueous solution through Amicon Ultracel-30K filters. Silylated samples were dialyzed (Slide-A-Lyzer dialysis cassette, Pierce, 20 000 MWCO) overnight in HPLC grade water. Samples (5 mg/mL) were then directly injected into the LTQ-FT mass spectrometer (Thermo Fisher Scientific) and analyzed in positive ion trap mode (600−2000 m/z). Typically, the sample (0.5 mL) was mixed with HPLC grade ACN (1 mL) and FA in HPLC grade water (3 v/v%, 0.5 mL). The instrument was externally calibrated with a resolution setting of 100 000. ProMass (Novatia) software was used for deconvolution of the intact and modified protein spectra to obtain the molecular weights. High baseline removal (1.2) was used in the deconvolution algorithm, and the output range was set to 40 000−100 000 Daltons (Da).

Reaction products of glycerol with DIDCS were analyzed using the same mass spectrometer in Fourier transform (FT) mode. Glycerol (1 mL) was mixed with DIDCS (1 mL), and the reaction was allowed to proceed for 3 days when it was finally neutralized with IPA (3 mL). The reaction mixture was then directly injected to the mass spectrometer without addition of water to prevent desilylation of the reaction products.

**EDS/FIB/XPS.** The nature of functionalization at the individual particle level was analyzed by energy dispersive spectroscopy (EDS) using INCA (Oxford Instruments) on the FEI Helios 600 Nanolab Dual Beam System which was also used for cross sectional study. Individual particle analysis was done by drying the particles over the surface of copper tape and then coating them with 3 nm gold (Au)/palladium (Pd) (Cressington sputter coater 108 auto). Three different random particles per each sample were analyzed to quantify the amount of silicon. For cross sectional analysis using focused ion beam (FIB), particles were dried over the surface of a carbon planchet (Electron Microscopy Services) and coated with carbon [10.5 nm] in a
high resolution ion beam coater (Gatan 681) to prevent repositioning of Au/Pd upon cutting the particle at 30.0 kV with 9.7 pA Gallium ion (Ga⁺) current. Linescans at 5.0 kV were then performed on the particle to measure the variation of elements across the particle cross section.

X-ray photoelectron spectroscopy (XPS) using the monochromatic Al Kα source (1486.6 eV) of Kratos Axis Ultra (150 W) was used to investigate the structural changes and to follow the cross-linking chemistry. Particles were deposited on Au [65 nm]/titanium (Ti) [15 nm] covered silicon wafers, and two different points on each sample were analyzed; then, the measured values were averaged. During these experiments, the charge neutralizer was on. Four sweeps of 800 ms were performed for each peak. Individual peaks were fitted with Gaussian–Lorentzian shape (GL[30]) curves, and locations of the components were compared against the values reported in the literature after shifting the peaks for +0.24 eV to fix the aliphatic carbon peak at 285.0 eV.

**SEM/TEM/Optical Microscopy.** Scanning electron microscopy (SEM) images were obtained on a Hitachi S-4700 Cold Cathode Field Emission Scanning Electron Microscope by coating the samples (SEM) images were obtained on a Hitachi S-4700 Cold Cathode Field Emission Scanning Electron Microscope by coating the samples with 3 nm of Au/Pd alloy. Transmission electron microscopy (TEM) was carried by first staining the particles with osmium tetroxide through vapor deposition, and then, an image was obtained using JEOL 2010F-FasTEM on a copper grid. Postfabrication particle size measurements were carried on a dry sample by performing 28 different measurements on SEM images using ImageJ. Optical images of the particles were obtained with a LSM 700 (Zeiss) microscope in fluorescent channel.

**Circular Dichroism.** Circular dichroism (CD) spectra were obtained with Chirascan Plus spectrometer (Applied Photophysics) in a cell with a path length (d) of 0.1 cm at room temperature from 190 to 240 nm. Coomassie plus colorimetric assay (Thermo Scientific) was used to determine the protein concentration (c g/L). Solutions (15 μL) were mixed with assay reagent (300 μL), and the absorbance was measured at 595 nm (25–500 μg/mL). Concentrations obtained from the protein assay were then used to convert the CD data (θλ, millidegrees at specific wavelength λ) to Δε molar differential extinction coefficient per residue (M⁻¹cm⁻¹) assuming a mean residue weight (MRW) of 114.15 Da for BSA prior to and postmodification (Δε = ((MRW × θλ)/(c × 10 × 3298 × d))). CDPro software (Colorado State University) was used to analyze the CD spectra and quantify the structure. The spectra were fitted with CONTINLL subroutine using 7 bases, and the secondary structure of the protein was estimated by deconvoluting the peaks and calculating the percentage of helical, beta, turn, and random components of the molecule. Modified particles reacted for 24 h, and unmodified control particles were dissolved in pH 7.4 buffer for a period of 18 h at 4 °C and then diluted in 10 mM K₂HPO₄. For control particles (202 μg/mL) and those reacted for 24 h (82.5 μg/mL), concentrations were determined with coomassie plus assay. Melting experiment of BSA was done by heating a solution (350 μg/mL) using QNW temperature control from 25 to 95 °C to determine the structure of the heat denatured molecule.

**Alexa Fluor 555 BSA Conjugate Loading.** Alexa Fluor 555 conjugated BSA (5 mg) was loaded into the preparticle solution (per 250 mg of BSA) to fabricate fluorescent particles for studying the dissolution mechanism and kinetics. To understand the dissolution/desilylation process, particles were reacted with DIDCS for 6, 12, and 24 h, and then, the resulting material was exposed to pH 7.4 buffer to measure the fractional release of Alexa Fluor 555 conjugated BSA. Several different aliquots containing particles (3.13 mg) were prepared for each specific time point (15, 60, 150, 300, and 600 min). Each aliquot was then reconstituted in pH 7.4 buffer (600 μL), and at each
time point, the particles were spun down and the supernatant (100 μL) was analyzed in triplicate by measuring the fluorescence intensity (Excitation: 545 nm; Emission: 575 nm) using SpectraMax M5 (Molecular Devices) in Nunc 96 microwell plates (Thermo Scientific).

The intensity of soluble unfunctionalized particles at 600 min served as the control reference in calculation of the fractional dye release from the particles.

**RESULTS AND DISCUSSION**

**Particle Fabrication and Composition.** A SEM image of sieved BSA powder is shown in Figure 1A while that of the particles as fabricated with PRINT on the surface of the harvesting layer is displayed in Figure 1B. Harvested free particles of Figure 1C are obtained by washing them with ACN. A TEM image of the free particles is presented in Figure 1E. The composition of the particles postfabrication was determined to be 55.7 wt % BSA, 43.1 wt % lactose, and 1.2 wt % glycerol as schematically shown in Figure 1D. These particles were further loaded with Alexa Fluor 555 conjugated BSA as seen in Figure 1G. The mean external and internal diameters plus the thickness of the particles were measured as 2.88 ± 0.21, 0.7 ± 0.06, 0.56 ± 0.17 μm, respectively (Figure 1F).

**Component Reactivity.** Silylation is extensively used in organic chemistry to block certain reaction centers by introducing a silyl group in place of a labile hydrogen atom. Silylation reactions are reversible and proceed via a mechanism involving bimolecular nucleophilic substitution at the silicon atom. In many cases, the mechanism and rate of silylation are determined by the steric factors associated with the silylating agent or the compound being silylated. Rogozhin et al. have derivatized amino acids (Pro, Tyr, Trp, Val, Asp, Lys) and dipeptides (Gly-Ala, Val-Val, Gly-Tyr, Gly-Trp) with different silylating agents (trimethylsilyldiethylamine, bis(trimethylsilyl) acetamide, N-trimethylsilyl-N-methylacetamide, and hexamethyldisilazane) in organic solvents at 20 °C.

Silylation with chlorosilanes has been previously used by several authors on nonprotein/peptide based materials to alter the surface properties or incorporate drugs into polymeric matrices. Chlorosilanes have been used on multiwall carbon nanotubes to functionalize their surface. Gousse et al. modified cellulose whisker dispersed in organic solvents and concluded that it is possible to partially silylate the whiskers where the surface of the whiskers is silylated but the core is kept almost intact. Silylether chemistry has also been used for incorporating drugs into the matrix of particles as a pro-drug strategy with hydroxyl ethyl acrylate. Figure 2A compares the spectra of dialyzed BSA post-functionalization (12−24 h) with intact material. After 12 hr of functionalization, mass over charge (m/z) peaks for BSA are...
shifted toward higher values, which serves as the evidence of functionalization with DIDCS (Figure 2B). Exposure of BSA to DIDCS for 24 h causes formation of visible precipitates. However, due to low signal intensity after 24 h of silylation, no conclusions could be drawn regarding this sample. Once the spectra for intact and 12 hr modified samples are deconvoluted, an increase in the molecular weight of the BSA due to silylation is observed as illustrated in Figure 2C.

Liquid chromatography-tandem mass spectrometry (LC-MS/MS) evaluation of the structure of trypsin digested 12 hr functionalized BSA molecules did not reveal any detectable modification of the recovered peptidic sequence. This can be attributed to the susceptibility of the bonds formed to hydrolysis during the digestion period.

Glycerol is capable of forming a variety of combinations with DIDCS under the reaction conditions (Figure S2, Supporting Information). The proposed structures are shown in Figure 3 for compounds A through H. The most abundant component is compound E formed from the reaction of two equivalents of DIDCS with one equivalent of glycerol. Although less abundant compared to compound E, multiple glycerol molecules can also be linked together due to the multifunctionality of both DIDCS (2 × chlorine [Cl]) and glycerol (3 × hydroxyl [OH]) as confirmed by the presence of compounds D, G, and H. Furthermore, the reaction conditions are conducive to the formation of larger fragments. Nevertheless, assigning structures to these peaks is not straightforward due to a multitude of combinations. Errors involved between the observed and theoretical masses of the identified peaks are presented in Table S2, Supporting Information.

No evidence of reactivity of lactose with DIDCS could be detected under the reaction conditions. Although lactose bears multiple hydroxyl groups, we believe that insolubility of the material makes it immune to solid state functionalization carried in extra-dry ACN. Lactose is not soluble in acetonitrile—water mixtures containing less than 15% water.

Particle Ensemble Analysis. XPS is a powerful technique for detecting compositional variations on the surface that are induced by silanization. The amount of silicon on the surface of the particles was measured with XPS, and relative intensities of the peaks are plotted in Figure 4A. The sulfur (S 2p) peak serves as a protein marker (disulfide bond-cysteine) and therefore was used as the appropriate reference element for comparing and quantifying the relative intensities of each peak.
Both silicon (Si 2s) and (Si 2p) peaks increase with respect to sulfur (S 2p) as a function of reaction time. However, the data for Si 2s is not shown for brevity. High resolution peaks of silicon (Si 2s, Si 2p) and sulfur (S 2p) for unmodified and reacted samples along with the survey scans are compared in Figure S3, Supporting Information. High resolution scans of oxygen (O 1s), carbon (C 1s), and nitrogen (N 1s) peaks were performed to detect any chemical changes associated with functionalization. Figure 4B,C compares oxygen (O 1s) scans for unmodified particles and the sample silylated for 24 hr. The shifted values of the binding energies for the deconvoluted components are listed in Table S3, Supporting Information. Lopez and co-workers55 reported the value of 532.8 eV for the oxygen (O 1s) peak of 1,10-decanediol and 532.7 eV for poly(vinyl alcohol) and poly(propylene glycol) with reference to an aliphatic carbon peak at 285.0 eV. Therefore, it can be concluded that the reduction in the intensity of the oxygen (O 1s) peak (labeled B) is consistent with consumption of hydroxyl groups on the surface due to the functionalization after 24 hr. The oxygen (O 1s) peak (labeled A) of 531.4 eV is very close to peptidic oxygen of BSA as reported by Iucci et al.56 at 531.6 eV. Carbon (C 1s) and nitrogen (N 1s) scans for the same samples are also presented in Figure S4, Supporting Information.

**Individual Particle Analysis.** A representative particle prior to exposure to the electron beam of EDS is shown in Figure 5A. Post-EDS analysis, the particle retains its shape, although there is minor structural damage due to the electron beam (Figure 5B). Elemental mapping at 20.0 kV over the top surface of the particle qualitatively confirmed the presence of silicon (Si) in the particle as illustrated in Figure 5C. Elemental maps of oxygen (O), sulfur (S), chlorine (Cl), nitrogen (N), and carbon (C) are also shown in Figure 5D−H for comparison. The corresponding spectrum obtained from elemental mapping of the particle in Figure 5 is shown in Figure S5, Supporting Information.

**Cross Sectional Analysis.** Figure 6C,D shows the top and side views of a particle after being cross sectioned with the FIB, respectively. The distribution of the elements along the yellow line depicted in Figure 6D is plotted in Figure 6B as a function...
of transverse distance. The sum spectrum of the linescan is also compared with the spectrum of the background to confirm that no silicon signal is detected on the carbon background (Figure 6A). No variation could be observed in the silicon content along the scanned line of Figure 6D, which suggests that DIDCS penetrates all the way through the particle and the reaction is not limited only to the surface of the particle.

**Mechanism and Dissolution Profile.** To study the dissolution of particles, silylated particles were exposed to pH 7.4 buffer and the fractional release of Alexa Fluor 555 conjugated BSA was measured (Figure 7A, symbols). The data demonstrates that the longer the treatment of particles with DIDCS, the slower is the dissolution rate of the particles. This is consistent with the higher silicon content of the particles as a function of reaction time, obtained from EDS and XPS experiments.

Numerous mathematical models exist in the literature for describing the dissolution behavior. Weibull and Korsmeyer-Peppas models are fitted to the experimental data (Figure 7A, lines), and corresponding model parameters are listed in Table S4, Supporting Information. The diffusion exponent in the Korsmeyer-Peppas model is indicative of a release mechanism (mode of transport of solute) and is shape dependent.

To visualize the dissolution process for particles post-functionalization, the behavior of a single 12 hr silylated particle is tracked in pH 7.4 buffer as a function of time. As observed in Figure 7B, the intensity of Alexa Fluor 555 conjugated BSA in the particle gradually diminishes due to dissolution of the fluorophore from the particle into the surrounding environment. Effects of both acidic (0.1 v/v% TFA, approximate pH = 2.0) and basic (pH 10.0 buffer) conditions on the size of particles are illustrated in Figure 7C,E for particles functionalized for 24 hr, respectively. At these pH extremes, particles significantly swell to 4–5 times their original size of Figure 7D.

Shirai et al. have studied the effect of different pHs on the hydrolytic removal of silyl groups from trialkyl silyl ethers. Their results indicate that there exists a pH where the observed hydrolysis rate constants are minimized. They demonstrated that this rate of hydrolysis is also affected by the electronegativity and steric size of the substituents on both silicon and oxygen atoms. The extreme swelling of these particles in both acidic and basic conditions suggests that the prevailing mechanism of dissolution is desilylation (removal of silyl groups).

The extreme swelling of the particles in acidic pH as observed in Figure 7C leaves the protein content of the particles prone to degradation in the harsh acidic environment of the gastrointestinal tract, and therefore, an oral delivery route for these suspensions is not envisioned. A measure for the stability of the suspension of 24 hr functionalized particles in pH 7.4 buffer is presented in Figure S6, Supporting Information.

**Structural Evaluation.** CD signal can determine structural changes associated with processing or functionalization of the protein in the particles. An examination of the melting behavior of intact BSA was done to evaluate the structural changes associated with heat denaturation of the molecule, which will
enable us to compare the structures of processed BSA to that of the intact or denatured molecule (Figure 8C). It can be seen that the intensity of the peaks at 209 and 222 nm is greatly reduced upon heating the sample. Figure 8C merely serves as a reference on how the structure of thermally denatured BSA evolves and illustrates the significant reduction in the helical structure upon thermal denaturation.

CD data for unmodified and modified particles are presented in Figure 8A and compared to intact and heat denatured BSA. Houen et al.63 showed that denatured BSA has limited solubility in water, and thus, structural changes postdenaturation are expected to be detectable with CD. Figure 8B shows the corresponding structural components obtained from deconvoluting/fitting the CD spectra while accuracy of the fits to the experimental data is examined in Figure 8A. The data in Figure 8B suggests that the silylation process used herein did not perturb the secondary structure of the fraction of BSA molecules that were dissolved in the buffer through the course of the dissolution experiment. Moreover, no evidence of thermal denaturation post particle fabrication could be obtained, primarily due to the fact that the denaturation mechanism is a time dependent process.64 This method of analysis had been used by Harn et al.2 to analyze the structure of recombinant humanized IgG1κ and also by Estey et al.65 to study the degradation of BSA under acidic conditions.

Aggregates of denatured protein can induce antibody, but they are not expected to induce neutralizing antibody due to lost protein conformation on denaturation. Formation of natively conformed aggregates on the other hand can be detrimental for therapeutic proteins due to the generation of neutralizing antibody response.66 Although subvisible particles less than 10 μm are not being evaluated in the USP (788) test, they can potentially impact clinical performance.67 Nevertheless, Singh et al.68 emphasize the uncertainty around the potential immunogenicity risk of subvisible particles in pharmaceutical formulations and conclude that the direct link between aggregation and clinical immunogenicity is not well established. They also assert that preclinical studies are not yet capable of evaluating the immunogenicity potential of product attributes such as aggregates and particles smaller than 10 μm. Experiments of van Beers et al.69 with different structures of recombinant human interferon Beta-1a proved that large unreacted aggregates (3.2 μm) and guanidine treated aggregated molecules (0.35 μm) are not necessarily immunogenic in transgenic immune-tolerant mice. The relation between structure and immunogenicity of recombinant human interferon Alpha-2b was investigated in wild-type and transgenic mice by Hermeling et al.,70 and no antibody responses for GA cross-linked molecule were detected. Moreover, boiled/denatured molecules (1−3 μm) proved to be poor immunogens in transgenic mice.

CONCLUSIONS

We have demonstrated that the combination of PRINT technology and silylation provides a valuable tool for tuning the solubility of the protein particles while controlling their size and shape. EDS and XPS confirm the modification at the particle and ensemble levels, respectively. The silylation process...
renders the solubility of the particles to be tunable, eventually dissolving in aqueous media while not perturbing the secondary structure of the fraction of BSA released from the particles. The rate of dissolution can be controlled by the time particles spend in the functionalization media consisting of extra-dry acetonitrile and diisopropylchlorosilane. Extreme swelling of the particles in low and high pH confirms that desilylation is the mechanism of particle dissolution. To further understand the nature of functionalization, EDS cross sectional analysis confirmed that the chlorosilane diffuses all the way through the particles and is not just limited to the surface.

Lowering the temperature used in the filling/harvesting stages of the particle fabrication enables us to better protect the sensitive contents of the particles against various modes of instabilities. Fabricating particles from homologous albumin instead of a heterologous one is recommended as it allows us to prepare particles that are made from nonimmunogenic ingredients. Immunogenicity of our subvisible functionalized particles loaded with a therapeutic protein will be thoroughly investigated in future, to elucidate the role of functionalization and fabrication parameters. In vivo behavior of these functionalized particles and their potential in extending protein retention is a subject of a separate study.

**ASSOCIATED CONTENT**

1. Supporting Information
2. HPLC analysis of the particles postfabrication. (2) Schematic of particle silylation. (3) FT-MS analysis of reaction products of glycerol and DIDCS. (4) Estimated error involved in the structure assignment for the reaction products of glycerol and DIDCS. (5) High resolution XPS scans of silicon and sulfur. (6) High resolution XPS scans and peak locations. (7) EDS elemental mapping performed on an individual particle. (8) Parameters for Korsmeyer-Peppas and Weibull models. (9) Optical density measurement for the suspension of functionalized particles. This material is available free of charge via the Internet at http://pubs.acs.org.

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**Notes**
The authors declare the following competing financial interest(s): Joseph M. DeSimone is a founder and maintains a financial interest in Liquidia Technologies.

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