Polymer-Coated Hydroxyapatite Nanoparticles for the Delivery of Statins

Thomas L Moore1, Ann-Sofie Schreurs2, Rachel A Morrison1, Erika K Jelen1, Joachim Loo2, Ruth K Globus3* and Frank Alexis1,4*
1Department of Bioengineering, Clemson University, Clemson, SC 29634, USA
2Space Biosciences Division, NASA Ames Research Center, Moffett Field, CA 94035, USA
3School of Materials Science and Engineering, Nanyang Technological University, Singapore 639798, Singapore
4Institute of Biological Interfaces of Engineering, Department of Bioengineering, Clemson University, Clemson, SC 29634, USA

Abstract
Statin drugs have been previously reported to increase the activity of osteoblasts, and are being investigated to treat low bone density pathologies. A controlled release nanoparticle drug delivery system may be used to prolong the exposure of osteoblasts to statin. Furthermore, nano-hydroxyapatite (nHA) has known osteoconductive and osteoinductive properties. The success of nanoparticle drug delivery systems is associated with safety, stability, biodegradability, and pharmacological parameters. nHA have been used as gene and drug delivery vehicles, however in vivo studies have shown that nHA aggregate, accumulate in the lungs, and impede the function of pulmonary surfactant. To address some of these challenges, we present a nHA coated with the biodegradable co-polymer poly(glycolide)-poly(ethylene glycol) (PGA-PEG). PGA coating thickness could be precisely tuned by surface initiated ring opening polymerization to control the release of lovastatin. PGA-PEG coated nHA were shown to be internalized in human osteoblasts, and were also shown to maintain 70% cell viability in human umbilical vein endothelial cells at doses up to 500 µg mL−1 in vitro. Preliminary in vivo maximum tolerated dose studies showed that PGA-PEG coated nHA were less toxic compared to uncoated nHA. This hybrid polymer-nHA nanoparticle drug delivery system has potential as a controlled release system for the delivery of statins to treat low bone density pathologies.

Keywords: Polymer-coated hydroxyapatite nanoparticles; Strains; Statin drugs; Polymerization

Introduction
Hydroxyapatite nanoparticles (nHA) have been investigated for numerous biomedical applications including implant coatings, tissue engineering scaffolds, in addition to gene and drug delivery systems [1-4]. In drug delivery, nHA are appealing due to drug loading and release through adsorption/electrostatic interactions, ability to tune nHA morphology, and ability to facilitate endosomal escape [4-6]. However, systemically delivered nHA has limited utility due to aggregation, and particle heterogeneity. Previously, nHA were shown to aggregate in vivo and be rapidly cleared by the liver [7,8]. nHA also shown to block pulmonary capillaries, or inhibit the function of pulmonary surfactant [9,10]. Another challenge with nHA for drug delivery is that drug loading on nHA is generally limited to loading via adsorption or electrostatic interactions [4,11-14]. Here we developed a nHA drug delivery system with a multifunctional polymer coating to sequester and control the release of drugs (Figure 1).

Previous studies have investigated nHA for the delivery of anti-cancer drugs, antibiotics, genes, peptides, and bone growth drugs [4,7,13-16]. nHA can be loaded with charged molecules via adsorption or electrostatic interaction, and Kundu et al. [14] showed that doxorubicin, an anticancer drug, could load 30-50% onto the surface of nHA. Furthermore, Palazzo et al. [12] loaded alendronate, cisplatin, or di(ethyleneediamineplatinum) medronate onto nHA via adsorption at 92%, 82%, and 52% of drug adsorbed, respectively. Drugs or molecules could also be loaded via encapsulation into the calcium phosphate matrix. Bhisi et al. [4] encapsulated plasmid DNA (pDNA) within calcium phosphate nanoparticles (NP) at 5.88 µg per mg NP for intracellular pDNA delivery. Calcium phosphate NP have also been loaded with organic dyes via encapsulation, as a proof-of-concept for loading small, organic molecules [17]. Loading drug by adsorption to nHA surface may however result in potentially undesirable burst release of drug [18]. This could also result in changes to the physico-chemical properties of the NP, which may in turn affect the NP physiological behavior [19].

Use of a polymer coating may mitigate some of the challenges associated with nHA-mediated drug delivery, and polymeric nanoparticles have been previously investigated to treat bone pathologies [15,20,21]. Polymer coatings are able to encapsulate or bind drugs, prolong and tune release profiles, and mediate the interaction of the NP with the biological environment [15,21-24]. Shi et al. [22] reported microspheres of poly(lactide-co-glycolide) (PLGA) which encapsulated nHA and alendronate, a bisphosphonate drug used to treat bone diseases such as osteoporosis and Paget’s disease. These microspheres, between 77 and 137 µm in diameter, could sustain the release of alendronate over a 30-day period. Another study showed that while hydroxyapatite microspheres loaded with doxorubicin via adsorption exhibited burst release, PLGA-coated HA microspheres were able to sustain release for up to 7-days [23]. We report nHA coated with a poly(glycolide)-poly(ethylene glycol) (PGA-PEG) copolymer for the controlled release ofLovastatin. As a proof of concept for using this PGA-PEG coated nHA drug delivery system (nHA-PGA-PEG), lovastatin was chosen as a potential therapeutic for the anabolic treatment of bone density-loss diseases. With nHA-PGA-PEG, the polymeric coating is able to encapsulate hydrophobic lovastatin within the PGA layer via hydrophobic-hydrophobic interactions. This loading approach could also be utilized to load other hydrophobic drugs.

*Corresponding authors: Frank Alexis, Department of Bioengineering, Clemson University, Clemson, SC 29634, USA, Tel: 864-656-3051; Fax: 864-656-4468; E-mail: falexis@clemson.edu
Ruth Globus, Space Biosciences Division, NASA Ames Research Center, Moffett Field, CA 94035, USA, Tel: 864-656-3051; Fax: 864-656-4468; E-mail: ruth.k.globus@nasa.gov

Received September 01, 2014; Accepted October 15, 2014; Published October 24, 2014

Citation: Moore TL, Schreurs AS, Morrison RA, Jelen EK, Loo J, et al. (2014) Polymer-Coated Hydroxyapatite Nanoparticles for the Delivery of Statins. J Nanomed Nanotechnol 5: 237. doi: 10.4172/2157-7439.1000237

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Furthermore, encapsulation also protects the drug from premature degradation, and controls release.

Statins are one class of drug shown to stimulate bone formation by enhancing the expression of bone morphogenic protein-2 (BMP-2) in osteoblasts [25,26]. In a culture of murine (2T3) or human (MG-63) bone cells, statins enhanced the expression of BMP-2 mRNA, increased the expression of type I collagen mRNA, lowered the expression of collagenase in MC3T3-E1 mouse osteoblast cells, and increase the expression of alkaline phosphatase [27]. Local controlled delivery of lovastatin at a fracture site using lovastatin microparticles has been shown to significantly improve fracture healing in mice [28], and controlled release of statins also improved osteogenic responses [21,29,30]. Thus, release kinetics and sustained exposure to lovastatin are important factors for the use of statins to treat bone pathologies.

PLGA nanoparticles have previously been used to deliver lovastatin, and Ho et al. [21] reported PLGA particles that could deliver lovastatin over a 15-day period. Tai et al. [15] utilized PLGA nanoparticles co-loaded with simvastatin and nHA, and the nHA could tune the simvastatin release kinetics and adjust the pH of the microenvironment around the drug. Simvastatin could be released over a 14-day period.

Our study also utilized an outer PEG layer to stabilize the nanoparticle in an aqueous environment, as PEG has been shown to stabilize NP, reduce protein adsorption, and prolong the circulation time of nanocarriers in vivo [31,32]. Finally nHA, a primary component of bone tissue, has been shown to be osteoinductive and osteoconductive [33-38]. Thus, the nHA-PEG drug delivery system is designed such that the outer PEG shell stabilizes the NP in solution, the PGA coating sequesters hydrophobic lovastatin and controls its release, and the nHA core may further enhance osteoblast activity.

**Materials and Methods**

**Chemicals and reagents**

Nano-hydroxyapatite was purchased from SkySpring Nanomaterials, Inc. (Houston, TX, USA). Methoxy-poly(ethylene glycol)-isocyanate (mPEG-isc), MW 5000, was purchased from Nanocs, Inc. (New York, NY, USA). Glycolide, phosphazene base P$_2$-t-Bu at 2 M in THF, and all solvents were purchased from Sigma-Aldrich. Lovastatin was purchased from TCI America (Philadelphia, PA, USA). All materials were used as provided.

**PGA-PEG coating of nHA**

In a typical reaction, 40 mg of nHA and 464 mg (4 mmol) of glycolide were dried overnight under 32 in Hg vacuum. Glycolide and nHA were dissolved in 3 mL anhydrous dimethylformamide (DMF), and P$_2$-t-Bu was added. The reaction vessel was purged under N$_2$, and the reaction continued for 24 hr. PGA coating thickness was controlled by varying the ratio of nHA and glycolide. Next, 100 mg (0.02 mmol) of mPEG-isc was added directly into the reaction. This reaction continued under N$_2$ for an additional 6 hr. The final product was washed twice with DMF and possible free monomer/polymer, and once with HyPure H$_2$O via centrifugation at 5,000 rpm for 5 min. The pellet was then lyophilized and stored at -20°C under N$_2$, until used.

**Nanomaterial characterization**

Thermogravimetric analysis (TGA) was recorded on a TA Instruments Hi-Res TGA 2950 thermogravimetric analyser under N$_2$, from 25°C to 600°C at 20°C min$^{-1}$. Transmission electron microscopy (TEM) was taken on a Hitachi H9500 high-resolution TEM at 300 kV. Fourier transform infrared spectroscopy (FTIR) was recorded on a Thermo-Nicolet Magna 550 equipped with a Thermo-SpectraTech
Foundation series Endurance Diamond ATR. IR: 2881 cm⁻¹ (-CH₂-
CH₂-O-), 1745 cm⁻¹ (C=O).

**In vitro release of lovastatin**

Release of lovastatin from nHA-PGA-PEG was determined using high performance liquid chromatography (HPLC). nHA nanoparticles coated with varying thicknesses of PGA were loaded with lovastatin using a solvent evaporation method. Briefly, particles were dissolved at 5 mg mL⁻¹ in a solution of lovastatin in acetonitrile (1 mg mL⁻¹). This NP solution was then dropped into HyPure H₂O at a 1:2 ratio and stirred for 2 hr. Lovastatin/NP was then washed three times in HyPure H₂O via centrifugation in 100 kD MWCO centrifugal filter units at 3500 rpm for 5 minutes. 1 mg of Lovastatin/NP was then loaded into the top of 3.5 kD MWCO Slide-A-Lyzer MINI dialysis units. Dialysis units were stored in microcentrifuge tubes containing HyPure H₂O at 37°C. At each time point, the dialysate was collected and replaced with fresh H₂O. Dialysis was lyophilized and analysed for lovastatin content. HPLC was performed using a Waters 1525 Binary HPLC pump with a 2998 photodiode array detector. An Alltima C18 column (4.6x25 mm, 5 μm) was used with a mobile phase of 71% acetonitrile (ACN) and 29% 0.5 M ammonium acetate in H₂O at pH 4.0, and a flow rate of 1 mL min⁻¹. Lovastatin was detected at a wavelength of 238 nm.

**Cell culture**

Human umbilical vein endothelial cells (HUVEC) and endothelial growth media (EGM) were purchased from Lonza (Walkersville, MD, USA). Fetal bovine serum (FBS) was purchased from Atlanta Biologicals (Lawrenceville, GA, USA). HUVEC were grown on collagen-1 coated growth media (EGM) were purchased from Lonza (Walkersville, MD, USA). HUVEC were seeded at 10,000 cells well⁻¹ in a 96-well plate with five repeats per test concentration. Uncoated nHA and nHA-PGA-PEG were prepared by first dissolving at 5 mg mL⁻¹ in acetonitrile (ACN). Next, the nHA or nHA-PGA-PEG solution was dropped at a 1:2 ratio into sterile HyPure H₂O in an aseptic hood and stirred for 2 hr. NP were washed twice with HyPure H₂O, and once with sterile filtered phosphate buffered saline (PBS) via centrifugation in a 100 kD MWCO centrifugal filter unit. These separate solutions were then dispensed in EGM at varying concentrations and added to cells. HUVEC were incubated with nanomaterials for 24 hr at 37°C and 5% CO₂. Media was removed, and cells were gently washed twice with sterile PBS.

A PrestoBlue cell viability assay (Life Technologies) was performed by mixing PrestoBlue reagent at a 1:9 ratio with EGM. 100 μL of working solution was added to cells, and these were incubated protected from light for 45 min at 37°C and 5% CO₂. Fluorescent intensity was measured with a BioTek Synergy 4 plate reader at λ₅60 of 560 nm, and λ₆0 of 590 nm. Cell viability was determined by normalizing the fluorescent intensity of each test well to the average intensity of control cells treated with only EGM.

**In vitro cellular uptake**

Bone marrow cells grown under osteoblastogenic conditions were approximately 80% confluent after 10 days in culture. Alexa Fluor 532 was loaded into nHA-PGA-PEG via encapsulation following the lovastatin loading procedure. Multiple controls were included to ensure the specificity of nHA uptake and dye delivery as follows. The cells were imaged without any addition of the nHA or dye, or with the dye added alone or with nHA added alone. Cells were exposed to media only, media with fluorescent dye, or dye-loaded nHA-PGA-PEG. Fluorescent images were taken from 5 min to 72 hr following exposure. Fluorescent microscopy was recorded on a Zeiss LSM 510 microscope.

**In vivo maximum tolerated dose**

All in vivo work was done in accordance with Clemson Institutional Animal Care and Use (IACUC) approved protocols. Animals were housed at Clemson University’s Godley-Snell Research Center. BALB/c mice were treated with escalating doses of either uncoated nHA or nGP. Nanoparticles were prepared similar to the in vitro cytotoxicity studies, and systemically administered via tail vein injection. Mice were observed for behavioral signs of significant toxicity. Non-responsive behavior, or a mass loss of 15% of initial body weight was classified as toxic [40,41]. Mice were observed for 7 days following injection. Following the completion of the maximum tolerated dose study, mice were euthanized as per IACUC protocol requirements. Organs were fixed in 4% formaldehyde overnight and processed for histological analysis. Briefly, organs were embedded in paraffin, sectioned at 5 μm, and stained with hematoxylin and eosin (H&E), or Alizarin red.

**Results and Discussion**

Bone density loss can be attributed to a number of pathological abnormalities. For example, microgravity-induced bone density loss is caused by the lack of mechanical stresses in space [42,43]. As previously stated, nHA has been investigated for drug delivery applications. However, loading is often limited to drug adsorption or electrostatic interactions. Also, nHA use as a systemic therapy is limited due to instability and aggregation. For bone applications though nHA is promising due to its osteoconductive and osteoinductive capabilities, and because it is a naturally occurring, primary component of bone [33-36,38,44]. Thus, this nHA-PGA-PEG delivery system is able to overcome the challenge of drug loading by encapsulating drug within the PGA, increasing stability with a PEG coating, and retaining the nHA core. The multifunctional nanoparticle could prolong the release of an osteoblast-stimulating drug (lovastatin), and enable long-term space flight or potentially treat other pathologies [15,21,44].

**Nanoparticle synthesis and characterization**

nHA was coated by directly functionalizing the surface with poly(glycolide)-poly(ethylene glycol) (PGA-PEG). Hydroxyl groups within the crystal structure of nHA were utilized to initiate the ring-opening polymerization of PGA directly on the nHA surface (nHA-PG) [16,36,45,46]. By controlling the ratio of the monomer glycolide (M) and the catalyst P2-C3 (C), it was possible to change the thickness of the PGA coating. PEG was then coupled with the terminal hydroxyl groups on PGA chain ends by directly introducing isocyanate-functionalized PEG into the reaction mixture. Figure 2 shows the chemical characterization of nHA coated with PGA-PEG (nHA-PG-PEG). Figure 2A shows thermogravimetric analysis (TGA) of nHA, nHA-PG, and nHA-PG-PEG. There was no mass loss for nHA over a temperature range from 25°C to 600°C, indicating no presence of organic polymer (Figure 2A). However, by varying the monomer to catalyst ratio (M/C) at 200, 100, and 50, it was possible to achieve PGA coatings of 20, 40, and 60 wt%, respectively. Moreover, by introducing PEG-isocyanate directly in a one-pot reaction it was possible to further functionalize PGA coated nHA with a layer of PEG (Figure 2B).

Fourier transform infrared (FTIR) spectroscopy of nHA, nHA-
PGA, and nHA-PGA-PEG further verifies the presence of a polymer coating on nHA (Figure 2C). The carbonyl stretch at 1745 cm⁻¹ is present in both nHA-PGA and nHA-PGA-PEG, and corresponds to the ester bond in PGA. The carbonyl group also results in CH₂-C=O bending at 1425 cm⁻¹. In the nHA-PGA-PEG sample, there is a slight CH stretching vibration at 2880 cm⁻¹ which corresponds to the ether bond in PEG (Figure 2C, inset) [47].

Transmission electron microscope images show morphological differences between the uncoated nHA and nHA-PGA-PEG (Figure 3). Uncoated nHA were rod-shaped, and nHA-PGA-PEG showed a contrast between the nHA core and a polymeric shell. These data show that nHA was successfully coated with PGA-PEG, and that the thickness of the PGA layer could be tuned by varying the M/C ratio.
Lovastatin loading and release

The rate ofLovastatin release was controlled by varying the polymer coating thickness. Release from polymeric systems is controlled by a number of factors including polymer-drug interactions, polymer degradation and swelling, drug diffusivity through the polymer matrix, and polymer thickness [48-50]. Release from nHA-PGA was inversely related to polymer thickness: as polymer coating thickness increased, Lovastatin release slowed. As drug diffusivity is controlled by factors such as polymer crystallinity, swelling, degradation kinetics, and thickness, it is expected that thicker polymers coatings will result in slower release kinetics. Figure 4 shows that release from 20 and 40 wt% PGA coated nHA was instantaneous, while there was an initial lag period of 24 hr for Lovastatin release from 60 wt% PGA. Moreover, 20 wt% nHA-PGA showed a 1.32-fold increase in release compared to 60 wt% nHA-PGA after 14 days. While Lovastatin is generally administered as an oral therapeutic, it however exhibits low oral bioavailability [51]. Several studies have investigated Lovastatin for local injection to treat bone pathologies [21,28-30], however by encapsulating Lovastatin within a NP drug delivery system it may be possible to improve the treatment of bone pathologies systemically by delivering the drug to multiple sites (Figure 5).

**In vitro toxicity and cellular uptake**

While all components of nHA-PGA-PEG are independently used clinically for biomedical applications, nanoparticle drug delivery systems must be robustly tested for safety of use. Here, we measured cytotoxicity in HUVEC after treating with increasing doses of either uncoated nHA, or nHA-PGA-PEG for 24 hr. Five repeats were used for each test, and data were analyzed using a Tukey’s honestly significant difference test. Compared to the control, all formulations showed a slight decrease in cell viability. However, there was no significant difference in cytotoxicity from either uncoated or coated NP, and HUVEC maintained 70% viability at doses as high as 500 µg mL⁻¹. Nanomaterial cytotoxicity is a factor of NP stability, size, shape, surface functionalization, method of synthesis, uptake, degradation, and material. These cytotoxicity data appear similar to other cell types in vitro, where human monocyte-derived macrophages showed 70% viability after treatment with rod-shaped nHA at up to 500 µg mL⁻¹ [52]. Zhao et al. [53] tested the cytotoxicity of rod-shaped nHA in BEAS-2B human bronchial epithelial cells, RAW264.7 mouse monocyte/macrophage cells, and HepG2 human hepatocellular liver carcinoma cells. Cell viability was >90% following exposure up to concentrations of 300 µg mL⁻¹.

Primary bone marrow cells from adult mice were flushed and cultured in medium containing factors that promote osteoblastogenesis, and uptake studies were conducted to verify cellular internalization of nHA-PGA-PEG loaded with Alexa Fluor 532 (Figure 6). Some extracellular aggregation of nHA-PGA-PEG was observed, evident as intense fluorescence. These decreased over time and correlated with an increase of fluorescence localization inside the cells. The peak of intracellular fluorescence was reached in approximately 30-60 mins. Figure 7 shows a higher magnification to image the localization of the nHA-PGA-PEG within the cells. The fluorescence was clearly localized inside the cells, and seemed to concentrate into some organelles. These images were collected 24 hr after addition of the NP and after the media
was changed, thus to remove any remaining nHA-PGA-PEG. The fluorescence observed is specific to the cells.

### In vivo toxicity

One limitation for the use of nHA as a systemic delivery system is instability that can lead to toxicity. Previous studies have shown that nHA can aggregate in blood and become localized in the lungs. nHA also has been shown to adversely affect pulmonary surfactant in the lungs [9,10]. We hypothesized that the PGA-PEG coating therefore mediates the interaction of the NP with cells. Both PGA and PEG are FDA-approved materials frequently used in biomedical applications. PEG has also been shown to improve the aqueous stability of nanoparticles [32,54,55]. Thus, the polymer coating may reduce inherent cytotoxicity associated with nHA.

Preliminary maximum tolerated dose studies were performed in BALB/c mice by injecting escalating doses of uncoated nHA or nHA-PGA-PEG. Mice were monitored for signs of morbidity (lack of mobility, >15% loss of initial body mass), and were observed over 168 hr. Table 1 shows the survival time of mice injected with the two different formulations. Mice were able to tolerate higher doses of nHA-PGA-PEG compared to nHA. This is potentially due to the polymer coating, particularly the PEG improving the dispersion of nHA in blood. Aoki et al. [9] found the medium lethal dose of intravenously administered nHA in Wistar rats to be 160 mg kg⁻¹. The cause of morbidity in the rats was attributed to nHA aggregation and blockage of pulmonary capillaries. Our histological examination of mice treated with nHA at high doses show that this may also have been the cause of death for BALB/c mice treated with nHA (Figure 8). Tissues were stained with haematoxylin and eosin, or Alizarin red to show tissue structure or the presence of calcium, respectively. Morphologically, the saline injected lung tissue appeared healthy. H&E micrographs clearly show the clean structure of healthy alveolar sacs. The tissue from the nHA injection however appeared severely inflamed with a high presence of immune cells. Furthermore there is significant aberration in the alveolar structure in the nHA lung tissue compared to the saline injection. Alizarin red staining for calcium showed healthy lung tissue (pink color) following the saline injection, while the nHA injection showed large calcium deposits, shown as red aggregates. Small deposits of nHA were also found in the brain, liver, and heart tissues. These data supports results by Aoki et al. [9] that the cause of morbidity from systemically administered nHA was aggregation and blockage of pulmonary capillaries.

| Dose (mg kg⁻¹) | Survival Time (hr) | nHA-PGA-PEG |
|---------------|-------------------|-------------|
| 160           | 168.000           | 168.000     |
| 200           | 0.030             | 168.000     |
| 240           | 0.008             | 24.000      |
| 280           | 0.008             | 24.000      |

Table 1: Survival time of mice treated with uncoated or coated nHA.
Conclusions

This work shows a new approach for the functionalization of nHA with PGA-PEG to deliver lovastatin. PGA was grafted directly onto the surface of nHA, and PEG was attached to the chain-ends of PGA. The relatively hydrophobic PGA was able to encapsulate and control the release of lovastatin over a 2-week period. Moreover, release of lovastatin could be controlled by tailoring the thickness of the PGA coating. While nHA and nHA-PGA-PEG showed some toxicity in vitro, HUVECs showed 70% viability at doses as high as 300 µg mL⁻¹. nHA-PGA-PEG uptake was shown in mouse osteoblasts using fluorescent and differential interference microscopy. Finally, the PGA-PEG coating raised the maximum tolerated dose in BALB/c mice when compared to uncoated nHA. This work shows that the nHA system could be used to deliver lovastatin for treatment of low-bone density pathologies.

Acknowledgement

This work was supported by SC EPSCOR-NASA 2098490, and the SC EPSCOR-NASA South Carolina Space Undergraduate Research Grant. Facilities used were supported by the South Carolina COBRE Center of Biomaterials for Tissue Regeneration (CBTR) funded under NIH Grants 5P20RR021949-04 and 8P20GM103444-04. The authors would like to thank Kim Ivey for use of the TGA and FTIR, and Dr. John Parrish and Travis Pruitt for help with the in vivo

![Figure 8: Tissues of BALB/c mice injected with either saline or nHA at 280 mg kg⁻¹ were stained with haematoxylin and eosin (left) for tissue structure or Alizarin red (right) for calcium. Scale bars represent 100 μm.](image)
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