Delivery system for DNAzymes using arginine-modified hydroxyapatite nanoparticles for therapeutic application in a nasopharyngeal carcinoma model

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Abstract: DNAzymes are synthetic, single-stranded, catalytic nucleic acids that bind and cleave target mRNA in a sequence-specific manner, and have been explored for genotherapeutics. One bottleneck restricting their application is the lack of an efficient delivery system. As an inorganic nanomaterial with potentially wide application, nano-hydroxyapatite particles (nHAP) have attracted increasing attention as new candidates for nonviral vectors. In this study, we developed an nHAP-based delivery system and explored its cellular uptake mechanisms, intracellular localization, and biological effects. Absorption of arginine-modified nanohydroxyapatite particles (Arg-nHAP) and DZ1 (latent membrane protein 1 [LMP1]-targeted) reached nearly 100% efficiency under in vitro conditions. Using specific inhibitors, cellular uptake of the Arg-nHAP/DZ1 complex was shown to be mediated by the energy-dependent endocytosis pathway. Further, effective intracellular delivery and nuclear localization of the complex was confirmed by confocal microscopy. Biologically, the complex successfully downregulated the expression of LMP1 in nasopharyngeal carcinoma cells. In a mouse tumor xenograft model, the complex was shown to be delivered efficiently to tumor tissue, downregulating expression of LMP1 and suppressing tumor growth. These results suggest that Arg-nHAP may be an efficient vector for nucleic acid-based drugs with potential clinical application.

Keywords: hydroxyapatite nanoparticles, DNAzymes, latent membrane protein 1, transfection efficiency, cellular uptake

Introduction

Epstein–Barr virus (EBV) has been closely linked with the oncogenesis of many cancers, including nasopharyngeal carcinoma. EBV-encoded latent membrane protein 1 (LMP1) is considered to be one of the key oncoproteins contributing to EBV-mediated tumorigenesis. Genetic manipulation of LMP1 expression has been suggested as a novel strategy for the treatment of EBV-associated carcinomas.¹,²

DNAzymes are synthetic, single-stranded DNA catalysts that bind to their complementary sequence in a target mRNA by Watson–Crick base pairing and subsequently cleave the mRNA at predetermined phosphodiester linkages.³–⁵ Many studies have demonstrated that delivery of specific DNAzymes targeting tumor-associated genes exerts anticaner effects, supporting the efficiency of specific DNAzymes in downregulation of appropriate genes and highlighting DNAzymes as potential genotherapeutic molecules.⁶–¹² In our previous studies, we discovered that EBV-LMP1-targeted DNAzymes could downregulate LMP1 expression, resulting in inhibition of cell
proliferation, promotion of apoptosis, and enhancement of radiosensitivity both in vitro and in vivo.\textsuperscript{13-16}

Active DNAzymes must penetrate the cell membrane, be appropriately trafficked in cells, enter the nucleus, and locate their target mRNA before achieving their biological function.\textsuperscript{17,18} However, due to their poor pharmacokinetic properties and biological permeability restrictions, therapeutic use of DNAzymes is dependent on the capability of delivery systems. In recent years, much effort has been made to develop highly effective, efficient, and safe vectors that provide a stable complex and protect payloads from degradation, thereby ensuring delivery of DNAzymes to target cells.\textsuperscript{19}

The rise of nanotechnology has provided new methods and tools for the study of gene carriers.\textsuperscript{20} Among these, calcium phosphate nanoparticles have emerged as a potential vector for nonviral gene delivery.\textsuperscript{21-23} Hydroxyapatite [Ca\textsubscript{10}(OH)\textsubscript{2}(PO\textsubscript{4})\textsubscript{6}, HA], a ceramic compound with a composition similar to that of mammalian bone and the dentin mineral compartment, has the advantages of absorbability and a high binding affinity with a variety of molecules, excellent biocompatibility, low cytotoxicity, nonimmunogenicity, lack of oncogenicity, and relatively high loading capacity and transfection efficiency.\textsuperscript{24,25} Nanohydroxyapatite particles (nHAP) incorporating pEGFP-N1 are reportedly able to deliver DNA into gastric cancer cells without any significant cytotoxicity.\textsuperscript{26} Tan et al\textsuperscript{27} discovered that nHAP modified with protamine could improve the efficiency of gene transfection. Sun et al\textsuperscript{28,29} successfully used nHAP to deliver the NT-3 gene into the cochlear neurons of guinea pigs both in vitro and in vivo, and further demonstrated that surface modification of nHAP with polyethylenimine carrying specific genetic materials could pass through the intact round window membrane of the chinchilla with high transfection efficiency and low toxicity. Yan-Zhong et al\textsuperscript{30} used arginine-modified nanohydroxyapatite to change the surface charge of nHAP, thereby improving adsorption capacity in human epithelial cells. These studies demonstrate that nHAP may be a potentially effective and safe gene vector with possible clinical application.

In this study, we designed and prepared arginine-modified hydroxyapatite nanoparticles (Arg-nHAP) and examined the absorption efficiency of Arg-nHAP and DZ1 in vitro. We demonstrated that Arg-nHAP can efficiently deliver DNAzyme into cells, release it, and have biological functions both in vitro and in vivo. We further elucidated the mechanisms of cellular uptake and intracellular trafficking of the Arg-nHAP/DZ1 complex as an energy-dependent endocytotic process.

### Materials and methods

#### Materials

The chemicals, inhibitors, transfection reagents, and cell culture media used in these experiments were sourced as follows: fluorescein isothiocyanate (FITC)-labeled DZ1 (FITC-DZ1) and control DNAzyme (CON) were synthesized by Oligos Etc Inc (Portland, OR, USA); Lipofectamine\textsuperscript{TM} 2000, ProLong® gold antifade reagent with DAPI (4′,6-diamidino-2-phenylindole), and trypsin-EDTA were from Invitrogen Life Technologies (Grand Island, NY, USA); high-performance liquid chromatography grade filipin III (>85%), phenylarsine oxide (≥97%), MTS (3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium), and sodium azide were from Sigma-Aldrich (St Louis, MO, USA); 2-deoxy-D-glucose was from Tokyo Chemical Industry (Tokyo, Japan); Fugene HD was from Roche (Basel, Switzerland); and fetal bovine serum was from Gibco (Grand Island, NY, USA).

#### Design and synthesis of DZ1

DZ1 was designed to be effective in suppressing expression of the target protein LMP1.\textsuperscript{13} CON was designed based on the sequence of DZ1 by introducing two mutations in the catalytic core at positions 6 and 7 (5′–3′). To determine localization of DZ1 in vitro and in vivo, DZ1 was labeled with FITC at the 5′ end.

#### Preparation and characterization of Arg-nHAP

Arg-nHAP was synthesized at the State Key Laboratory for Powder Metallurgy of Central South University by a chemical coprecipitation hydrothermal technique. First, 0.2 mol/L Ca(NO\textsubscript{3})\textsubscript{2} and 0.2 mol/L (NH\textsubscript{4})\textsubscript{2}HPO\textsubscript{4} solutions were mixed at a ratio of 5 to 3 (v/v, Ca to P mole ratio of 1.67) with arginine (4%) preadded to the phosphate solution. The reaction temperature was 60°C, and the pH of the mixture was adjusted to 10–11 by ammonia water. After stirring for 30 minutes, the mixed solution was poured into a Teflon\textsuperscript{®}-lined stainless-steel autoclave and underwent hydrothermal treatment at 170°C for 5 hours. After filtration and drying, crystalline Arg-nHAP was obtained.\textsuperscript{26,29}

Crystalline Arg-nHAP was diluted to 5 mg/mL using ultrasonic dispersion for 60 minutes (ultrasonic homogenizer, VC500/750, Sonics & Materials, Inc., Newtown, CT, USA) and was observed for 2 hours until it appeared layered and milky. Finally, the Arg-nHAP suspension was stored at 4°C after autoclaving. The particle size was measured using a
transmission electron microscope (JEM-2100F, JEOL, Tokyo, Japan). The zeta potential was measured by a Zetasizer Nano ZS (Malvern Instrument Company, Malvern, UK). Arg-nHAP phase analysis was measured using an X-ray diffractometer (D-Max/2550VB+; Rigaku, Tokyo, Japan) with Cu Kα radiation (λ = 1.54178 Å, 40 kV, 30 mA). The scanning angle/speed was 25°–55°/2.4° per minute and 5°–75°/5° per minute.

Absorption efficiency of DZ1 and Arg-nHAP

The absorption efficiency of Arg-nHAP and DZ1 was determined by centrifugation assay. The Arg-nHAP/DZ1 complex was prepared by mixing 100–750 µg Arg-nHAP solution with 5–60 µg DZ1 followed by incubation on a thermostat shaker (500 rpm, 21°C) for one hour. The mixtures were centrifuged (13,000 g, 4°C) for 10 minutes. The nonadsorbed DNAzyme was quantitated in the supernatant by ultraviolet spectrophotometry at 260 nm using a nucleic acid analyzer (Beckman Coulter Inc, Fullerton, CA, USA).

DZ1 release in vitro

DZ1 release from Arg-nHAP/DZ1 was determined using a centrifugation assay. DZ1 was labeled with FITC at the 5′ end. The Arg-nHAP/DZ1 complex was prepared in Roswell Park Memorial Institute (RPMI) 1640 medium or in cytoplasm, plated in a 96-well Nunc plate (Sigma-Aldrich), and incubated at 37°C for 0, 8, 12, 24, and 48 hours, respectively. The mixtures were centrifuged (20,000 g, 4°C) for 15 minutes. The DZ1 released was quantitated in the supernatant using a multifunctional microplate reader (Varioskan Flash, 3001, Thermo Fisher Scientific, Waltham, MA, USA) at 488 nm/525 nm.

Cell culture

CNE1 is a poorly differentiated EBV-negative nasopharyngeal squamous carcinoma cell line and CNE1-LMP1 is a stable LMP1-integrated nasopharyngeal carcinoma cell line. All cells were cultured in RPMI 1640 medium with 10% fetal bovine serum. All media were supplemented with 100 IU/mL penicillin, 100 µg/mL streptomycin, and 2 mmol/L L-glutamine. All cultures were incubated in a humidified atmosphere containing 5% CO₂ at 37°C.

Arg-nHAP/DZ1 transfection of CNE1-LMP1 cells

Prior to transfection, 2.0 × 10⁵ CNE1-LMP1 cells per well were seeded in a six-well plate in RPMI 1640 medium with 10% fetal bovine serum and incubated at 37°C until 60%–70% confluence at the time of transfection. The Arg-nHAP/DZ1 complex (40 µg DZ1 to 400 µg Arg-nHAP) was prepared as described above. Lipofectamine 2000 or Fugene HD complexes containing the same amount of FITC-DZ1 were prepared as a comparison following the manufacturer’s protocol. The complexes were then added to the cells and incubated at 37°C for 4 hours, followed by addition of complete medium to the wells for further experiments.

FACS analysis

The transfected CNE1-LMP1 cells were incubated at 37°C for 4 hours. The cells were washed with phosphate-buffered saline, harvested by treatment with trypsin-EDTA, resuspended in ice-cold phosphate-buffered saline (pH 7.4), and examined immediately using a FACSort flow cytometer (FACS, Becton Dickinson, Franklin Lakes, NJ, USA). After 20,000 total events were acquired per sample, cells were defined by assessing cell size and granularity in the forward scatter and side scatter channels. FITC fluorescence in the cell population was expressed as the geometric mean of positive events after subtraction of background fluorescence. The percentage of FITC-positive cells was determined using the CellQuest software program (Becton Dickinson).

Cell viability assay

The MTS assay was used to measure cell cytotoxicity. CNE1-LMP1 cells were plated at a density of 1.0 × 10⁵ cells/well in 96-well plates containing RPMI 1640 medium with 10% fetal bovine serum and incubated at 37°C until 60%–70% confluence at the time of transfection. Arg-nHAP/DZ1, Lipofectamine 2000/DZ1, or Fugene HD/DZ1 complexes were prepared as described above (final mass of DZ1, 40 µg). Twenty-four hours later, the complexes were added to the cells and incubated at 37°C for 24 and 48 hours, respectively. Next 20 µL of MTS was added to each well and incubated at 37°C. The absorbance of each well was read on a microplate reader at 490 nm.

Mechanism studies

Cells were seeded in a 12-well plate at a density of 6.0 × 10⁴ cells/well one day prior to transfection. Twenty-four hours after seeding, the cells were pretreated with inhibitors. The concentrations of the inhibitors were chosen based on previous studies, i.e., 10 mM sodium azide and 50 mM 2-deoxy-D-glucose for one hour; 0.15 µM phe nylarsine oxide for 10 minutes; and 1.25 µg/mL of filipin for one hour. The effect of low temperature on the cellular uptake of the complex was investigated at 4°C.
pretreatment, the cells were transfected with the Arg-nHAP/ FITC-DZ1 complex and analyzed by FACSsort flow cytometry as described above.

Confocal microscopic analysis
Cells were plated in four-chamber polystyrene vessels on glass slides (Falcon™, BD Biosciences, Bedford, MA, USA) at a density of 4.0 \( \times \) 10^4 cells per chamber one day prior to transfection. Twenty-four hours after seeding, the cells were transfected with the Arg-nHAP/DZ1 complex. After incubation at 37°C for 4 hours, the cells were fixed with 4% paraformaldehyde in phosphate-buffered saline at 4°C for 30 minutes. The cells were then washed with cold phosphate-buffered saline before their nuclei were stained with DAPI. Each slide was then sealed with a cover slip (Fisher Scientific, Pittsburgh, PA, USA). All slides were protected from light and stored at 4°C for further analysis using confocal microscopy (Leica TCS SP5, Nussloch, Germany).

Western blot analysis
The cells were transfected as described above and incubated for 48 hours at 37°C, then lysed in lysis buffer. After protein quantification using the bicinchoninic acid assay (Pierce Chemicals, Rockford, IL, USA), total protein was resolved on 10% polyacrylamide sodium dodecyl sulfate gel and transferred onto a nitrocellulose membrane by electroblotting. The membrane was incubated in blocking buffer for 2 hours, followed by incubation with a primary antibody against LMP1 (Dako, Carpinteria, CA, USA) (1:250) or β-actin (1:1000) (Santa Cruz Biotechnology, San Diego, CA, USA) as an internal control overnight at 4°C. The membrane was washed in Tris-buffered saline Tween® 20 and further incubated with a secondary antibody (1:2000, anti-LMP1, Dako, M0897). Protein expression was determined using a supersignal chemiluminescence system (Pierce Chemicals) followed by exposure to autoradiographic film.

Xenograft model of human nasopharyngeal carcinoma in nude mice
Athymic BALB/C nude mice (aged 4–6 weeks, female) were injected subcutaneously with 5 \( \times \) 10^6 CNE1-LMP1 cells in 0.2 mL of RPMI 1640 medium. Tumor volumes were determined according to the formula of length \( \times \) width \( \times \) height \( \times \) (\( \pi/6 \)) using a digital caliper. When the tumor volume reached 60–110 mm\(^3\), the animals were randomly divided into five groups (n = 5 each) to receive an intratumoral injection of 20 µL saline containing 100 µg of DZ1 or CON adsorbed by 3 µL of Fugene HD or 20 µL of Arg-nHAP, respectively, twice a week (six times in total). Tumor growth was monitored every 2 days. The mice were sacrificed at the end of the experiments, and tumor tissues were weighed and stored in 10% neutral-buffered formalin for histochemical examination.

Immunohistochemistry
Paraffin-embedded tissues were sectioned and stained for LMP1 using a streptavidin-biotin antibody coupled to peroxidase. Immunohistochemical staining of LMP1 was performed according to standard procedures. The immune complex was visualized with DAB solution and counterstained with hematoxylin (Sigma-Aldrich). Slides were scored for positivity (brown-stained cytoplasm) and negativity (blue nuclei) using a light microscope.

Statistical analyses
All statistical analyses were performed using Statistical Package for the Social Sciences version 11 software (SPSS Inc, Chicago, IL, USA). If not otherwise stated, the means of at least three independent experiments \( \pm \) standard deviation are shown. All data were calculated using the Student’s t-test. A difference with \( P < 0.05 \) was considered to be statistically significant, and \( P < 0.01 \) was considered to be very significant.

Results
Preparation and characterization of Arg-nHAP
Crystalline Arg-nHAP was synthesized by a chemical coprecipitation hydrothermal technique, and its X-ray diffraction graph is shown in Figure 1B. The diffraction peaks of each sample are in agreement with the hexagonal nHAP in the P6\(_3\) m space group (JCPDS 09-0432). This X-ray diffraction pattern is consistent with that of standard hydroxyapatite. No other phases could be detected, indicating that the sample was pure nHAP crystal. The size and morphology of the nHAP crystal was identified in transmission electron micrographs (Figure 1A and C), and it could be seen that the Arg-nHAP crystal had a rod-like shape with a diameter of 20–50 nm and length of 50–200 nm. Imaging also showed that DZ1 adsorbed with Arg-nHAP had no significant effect on morphology and size. At pH 7.4, the zeta potential of Arg-nHAP, DZ1, and Arg-nHAP/DZ1 was 25.47 ± 2.9, −16.46 ± 2.62, and 18.06 ± 1.67 mV, respectively (Figure 1D).
Absorption efficiency of DZ1 and Arg-nHAP

The efficiency of DZ1 adsorbed to Arg-nHAP was assessed by centrifugation assay. DZ1 was added to different concentrations of Arg-nHAP. The complex was centrifuged, and the absorbance of the supernatant was measured by ultraviolet spectrophotometry at 260 nm. The absorption efficiency of Arg-nHAP and DZ1 was high, reaching almost 100% under optimal conditions, suggesting that DZ1 could be effectively adsorbed on Arg-nHAP (Table 1). Increasing Arg-nHAP was likely to increase the total capacity of DZ1 loading. As the amount of Arg-nHAP increased to 600 μg, the amount of DZ1 adsorbed decreased substantially, suggesting that exclusion of DZ1 occurred due to saturation of absorption.
In our preliminary study, we found that cellular uptake efficiency was optimal when 40 µg of DZ1 was transfected into CNE1-LMP1 cells. Therefore, we used conditions (40 µg DZ1 to 400 µg Arg-nHAP) for subsequent experiments where the absorption efficiency was 76.48% ± 1.38%.

**DZ1 release in vitro**

To determine whether DZ1 could be released from the Arg-nHAP/DZ1 complex, we performed the release assay both in medium and in cytoplasm, which was prepared from CNE1-LMP1 cells. As showed in Figure 2, DZ1 release from Arg-nHAP/DZ1 was increased in a time-dependent manner, both in RPMI 1640 medium and in cytoplasm. Interestingly, DZ1 release from the Arg-nHAP/DZ1 in cytoplasm reached 80% at 48 hours, which was significantly higher than that in RPMI 1640 (55%). This suggests that the cellular environment could facilitate release of DZ1 from the complex.

**Cellular uptake efficiency of Arg-nHAP/DZ1 complex in CNE1-LMP1 cells**

To determine cellular uptake efficiency and cellular toxicity, DZ1 was labeled with FITC at the 5’ end. CNE1-LMP1 cells were transfected with the complex, and the cellular uptake efficiency was analyzed by FACS. The results demonstrated that only a slight increase in fluorescence was observed in the cells treated with free FITC-DZ1. In marked contrast, approximately 74.2% of cells were FITC-positive for uptake of the Arg-nHAP/DZ1 complex. As control systems, the Fugene HD/DZ1 and Lipofectamine 2000/DZ1 complexes had a cellular uptake efficiency of 89.8% ± 9.42% and 93.8% ± 7.94%, respectively (Figure 3A). When the transfected cells were subjected to fluorescent microscopy, no adverse effects on cell morphology were observed in cells treated with the Arg-nHAP/DZ1 complex (Figure 2B). A cytotoxicity study was carried out using MTS to quantify the survival rate of CNE1-LMP1 cells with that of control groups with the same level of DZ1. As shown in Figure 3C, the viability of Arg-nHAP was slightly better (approximately 80%) than the commercially available transfection reagents (the cell viabilities for Fugene HD and Lipofectamine 2000 were approximately 65% and 50%, respectively), suggesting that Arg-nHAP was not cytotoxic at the concentration range used in this study.

**Mechanisms for cellular uptake of the Arg-nHAP/DZ1 complex**

To determine if cellular uptake and intracellular trafficking of the Arg-nHAP/DZ1 complex was an active process or adenosine triphosphate-dependent, the transfected cells were treated with a mixture of metabolic inhibitors, ie, sodium azide and 2-deoxy-D-glucose. The effect of low temperature on cellular uptake of the complex was evaluated by incubation at 4°C. The results show that treatment with the mixture of sodium azide and 2-deoxy-D-glucose significantly reduced the cellular uptake efficiency to 31.9% as compared with uptake in the absence of these reagents. As shown in Figure 4, a significant reduction in cellular uptake efficiency was also observed at 4°C. These results suggest that the complex was likely to be taken up by cells through an energy-dependent endocytotic pathway.

To confirm that endocytosis was relevant to cellular uptake of the Arg-nHAP/DZ1 complex, the cells were pre-treated with the endocytosis inhibitors, filipin or phenylarsine oxide, prior to transfection. Phenylarsine oxide is a potent...
Figure 3 Cellular uptake efficiency of the Arg-nHAP/DZ1 complex, cell morphology, and viability. (A) Transfected cells were harvested and analyzed using FACS (DZ1, FITC-labeled). FITC fluorescence in the cell population was expressed as the geometric mean of positive events after subtraction of background fluorescence (nontransfected cells). The percentage of FITC-positive cells was determined using the CellQuest software program. All values are the mean of three measurements and are shown with error bars. (B) Transfected cells were subjected to fluorescent microscopy. (200×). (C) MTS assay was used to assess cellular viability of CNE1-LMP1 cells 24 and 48 hours following treatment with DZ1, Arg-nHAP, Lipofectamine® 2000, Fugene, Arg-nHAP/DZ1, Lipofectamine 2000/DZ1 and Fugene/DZ1. The absorbance of each well was read on a microplate reader at 490 nm, and the results indicating the cell viability were plotted as the percentage over controls (MOCK cells).

**Abbreviations:** FITC, fluorescein isothiocyanate; Arg-nHAP, arginine-modified nano-hydroxyapatite particles; DZ1, DNAzyme 1; FACS, fluorescence activated cell sorting; MTS, 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium; LF2000, lipofectamine 2000.

Inhibitor of clathrin-mediated endocytosis, whereas filipin inhibits the raft/caveolae pathway. Preincubation of cells with phenylarsine oxide 0.15 μM resulted in significantly reduced cellular uptake of the Arg-nHAP/DZ1 complex (36.7%). However, phenylarsine oxide did not completely inhibit active uptake of the complex when compared with cellular uptake efficiency at 4°C (Figure 4). Filipin 1.25 μg/mL also had a distinct effect on cellular uptake efficiency, with a reduction to approximately 50%. Using phenylarsine oxide and filipin, we demonstrated that endocytosis of the Arg-nHAP/DZ1 complex is both clathrin-dependent and caveolae-dependent.
Intracellular localization of the Arg-nHAP/DZ1 complex

To visualize the intracellular localization of the complex in CNE1-LMP1 cells, we utilized confocal microscopy to examine the transfected cells. Consistent with the results of FACS analysis, free FITC-CON added to the culture media was poorly internalized by the cells, and confocal images revealed very limited localization in punctate cytoplasmic regions. When the cells were treated with the Arg-nHAP/FITC-DZ1 complex for 24 hours, the bright, punctate fluorescence of FITC was distributed in the cytoplasm and nuclear region, as shown in the merged picture and the FITC image (Figure 5), and the same intracellular localization was found in the Lipofectamine 2000/DZ1 and Fugene HD/DZ1 controls, suggesting successful trafficking of DZ1 to the nucleus.

Downregulation of LMP1 expression in CNE1-LMP1 cells

We had previously demonstrated that DZ1 transfection of LMP1-expressing cells resulted in downregulation of expression of the target gene, LMP1.13,15 To examine whether Arg-nHAP could deliver a biologically active payload to cells, we transfected the Arg-nHAP/DZ1 complex into the cells and assayed DZ1-mediated inhibition of LMP1 expression. The data in Figure 6 show that DZ1 efficiently inhibited LMP1 expression when compared with CON-transfected and nontreated cells (P < 0.01), suggesting that DZ1 was successfully taken up into the cells, released into the cytoplasm, and suppressed LMP1 expression in cells.

Antitumor effects and downregulation of LMP1 expression in a CNE1-LMP1 xenograft model in nude mice

To determine whether the Arg-nHAP/DZ1 complex could downregulate LMP1 expression in vivo, we pre-established a CNE1-LMP1 xenograft model in nude mice to explore the effects of the complex on tumor growth. Fugene HD was used as the commercial control in vivo experiments due to its low cytotoxicity. As shown in Figure 7 and Table 2, Arg-nHAP/DZ1 significantly inhibited tumor growth compared with Arg-nHAP/CON and saline (P < 0.05). In addition, Fugene HD/DZ1 and CON had only a marginal effect on tumor growth (P > 0.5). These data suggest that Arg-nHAP may be used as a delivery vehicle for nucleic acid-based agents. To demonstrate that LMP1 expression was indeed inhibited in the tumor tissues, immunohistochemical staining was done on cross-sections of paraffin-embedded formalin-fixed tissues from the different treatment groups. The results show that LMP1 expression in tumor tissues treated with the Arg-nHAP/DZ1 complex was significantly reduced compared with CON (Figure 8), indicating that Arg-nHAP was delivered.
successfully to the tumor cells and that DZ1 could locate its target mRNA and exert its biological effects in vivo.

**Discussion**

The characteristics of DNAzymes, including their remarkable catalytic efficiency and specificity, make them an attractive approach to gene therapy. One of the problems for potential therapeutic use of DNAzymes is the lack of efficient delivery systems. Such systems should provide efficient delivery of DNAzymes into target cells, with protection from nucleases, promotion of endosomal release of DNAzymes into the cytoplasm, and location of the target mRNA.

Commercially available liposomes have demonstrated high cellular uptake efficiency and capacity in vitro tests. However, in vivo applications, they are usually accompanied by unwanted regional and systemic toxicity.

Calcium phosphate nanoparticles represent a unique class of nonviral vectors that have attracted increasing attention as new candidates for nonviral vectors in gene therapy.

![Figure 6](image_url)  
**Figure 6** Inhibition of LMP1 protein expression in CNE1-LMP1 cells using DZ1. Cells were transfected with DZ1 or control (40 µg). All values are the mean of three measurements and are shown with error bars.  
**Abbreviations:** Arg-nHAP, arginine-modified nanohydroxyapatite particles; GAPDH, glyceraldehyde 3-phosphate dehydrogenase; LMP1, latent membrane protein 1; DZ1, DNAzyme 1; LF2000, lipofectamine 2000.

![Figure 7](image_url)  
**Figure 7** Tumor growth curve of each group in a nasopharyngeal carcinoma xenograft model. When the tumor volume reached 60–110 mm³, the animals were injected intratumorally with 100 µg of DZ1 or CON adsorbed by 3 µL of Fugene HD or 20 µL of Arg-nHAP (20 µg per injection) twice a week (six times total). The tumor volume was measured twice a week. Tumor tissues were harvested and weighed at the end of the experiment.  
**Abbreviations:** Arg-nHAP, arginine-modified nanohydroxyapatite particles; CON, control DNAzyme; DZ1, DNAzyme 1.

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**Table 2** Statistical analysis of mean tumor volume

<table>
<thead>
<tr>
<th>Group</th>
<th>Cases</th>
<th>Mean ± SD</th>
<th>P value</th>
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<tr>
<td>Fugene HD/control</td>
<td>5</td>
<td>464.74 ± 164.78</td>
<td>0.730^a</td>
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<tr>
<td>Fugene HD/DZ1</td>
<td>5</td>
<td>353.82 ± 107.58</td>
<td>0.309^b</td>
</tr>
<tr>
<td>Arg-nHAP/control</td>
<td>5</td>
<td>505.82 ± 102.82</td>
<td>0.940^c</td>
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<tr>
<td>Arg-nHAP/DZ1</td>
<td>5</td>
<td>290.86 ± 85.40</td>
<td>0.037^d</td>
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<tr>
<td>Saline</td>
<td>5</td>
<td>498.83 ± 188.82</td>
<td></td>
</tr>
</tbody>
</table>

**Notes:**  
^a Fugene HD/control compared with saline;  
^b Fugene HD/DZ1 compared with saline;  
^c Arg-nHAP/control compared with saline;  
^d Arg-nHAP/DZ1 compared with saline;  
^e Fugene HD/control compared with Fugene HD/DZ1;  
^f Arg-nHAP/control compared with Arg-nHAP/DZ1.  
^Statistically significant (P < 0.05) versus control.  
**Abbreviations:** Arg-nHAP, arginine-modified nanohydroxyapatite particles; SD, standard deviation; DZ1, DNAzyme 1.
In the present study, we established an effective Arg-nHAP/DNAzyme delivery system in which nHAP was modified with arginine conjugated to a guanidyl group \((-\text{CH}_2)_3\text{NHC(NH}_3^+)\) on the surface of Arg-nHAP to prevent crystal growth. This system was shown to facilitate absorption of DNAzyme effectively, with cellular uptake of covalently attached particles to deliver active DNAzyme into cells and animals. The observed biological activity both in vitro and in vivo indicates that DZ1 delivered by nHAP should have been released from the complex in the cytoplasm and located its target mRNA to exert its biological effects.

In comparison with Lipofectamine 2000 and Fugene HD, which have been used widely in vitro settings, Arg-nHAP showed similar cellular uptake efficiency both in test tubes and in cell cultures. However, Arg-nHAP/DNAzyme showed much more efficient delivery in vivo than did Fugene, as evidenced by the strong biological activity of Arg-nHAP/DZ1 in suppressing nasopharyngeal tumor growth (Figure 7). This discrepancy between the in vitro and in vivo activity of Arg-nHAP and that of the commercially available transfection reagents may be due to the unique properties of the Arg-nHAP nanoparticles, including loading capacity, low toxicity, and biocompatibility.44,45

In this study, we explored the mechanisms of cellular uptake and intracellular trafficking of the Arg-nHAP/DZ1 complex. By inhibition studies using specific metabolic inhibitors, i.e., sodium azide and 2-deoxy-D-glucose, which block production of adenosine triphosphate by interfering with glycolytic and oxidative metabolic pathways in cells, the complex was determined to be probably taken up by cells through an active process.44,45 Further, a significant reduction in transfection efficiency of the complex was observed at 4°C, confirming that cellular uptake of the complex relied on an energy-dependent pathway.

Endocytosis is known to be a prevalent and major process for cellular uptake of nanomaterials.46,47 Studies have suggested the involvement of both clathrin-mediated and caveolae-mediated pathways in lipid-based and polymer-based gene delivery systems.48,49 For calcium phosphate-based gene delivery systems, both clathrin-dependent and caveolae-dependent endocytosis was suggested to be involved.50 To examine if endocytosis was relevant to cellular uptake of the Arg-nHAP/DZ1 complex, the cells were pretreated with endocytosis inhibitors, such as filipin or phenylarsine oxide, prior to transfection. Phenylarsine oxide inhibits clathrin-mediated endocytosis by reacting with vicinal sulfhydryls to form stable ring structures, whereas filipin interacts with 3-b-hydroxysterols in the plasma membrane to form filipin-sterol complexes and subsequently induces the filamentous caveolin-1 coat to disassemble rapidly, leading to inhibition of caveolae-mediated endocytosis. We demonstrated that treatment of cells with these inhibitors significantly reduced cellular uptake of Arg-nHAP/DZ1, suggesting that endocytosis of the Arg-nHAP/DZ1 complex was clathrin-dependent and caveolae-dependent.

The features of Arg-nHAP indicate that it may be an efficient vector for delivery of biologically active molecules. Further studies are needed to address issues such as the stability, pharmacokinetics, and toxicity of these nanoparticles before realization of their clinical potential.

**Conclusion**

Data from this study indicate that DZ1 was successfully encapsulated with Arg-nHAP and transfected into cells, where it was released, entered the nucleus, located the target mRNA, and inhibited LMP1 protein expression in vitro and in vivo. Arg-nHAP nanoparticles represent a promising nonviral carrier for gene therapy.

**Acknowledgment**

This work was supported by the Key Projects in the National Science and Technology Pillar Program during the Twelfth Five-year Plan Period (2013BAI01B07) and the National Nature Science Foundation of China (81172188; 81072220; 91129709; and 51102285).
Disclosure

The authors report no conflicts of interest in this work.

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