ORIGINAL RESEARCH

Atrial natriuretic peptide modified oleate adenosine prodrug lipid nanocarriers for the treatment of myocardial infarction: in vitro and in vivo evaluation

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Purpose: Myocardial infarction is a major cause of mortality and heart failure worldwide. One of the most effective methods of this injury is direct delivery of cardioprotective drugs to ischemia–reperfusion (IR) myocardium. The aim of the present study was to fabricate an adenosine (Ade) prodrug-based, atrial natriuretic peptide (ANP)-modified nanosystem for the treatment of myocardial infarction.

Materials and methods: Oleate adenosine prodrug (Ade-OA) and ANP-distearoylphosphatidylethanolamine-polyethylene glycol were synthesized. ANP-modified Ade-loaded lipid nanocarriers (ANP Ade/LNCs) were then self-assembled by using solvent evaporation method. In vitro drug release in the presence of plasma was evaluated. In vivo inhibition effect on infarct size, tissue distribution, and pharmacokinetics were investigated in rats with ischemic myocardium after intravenous injection.

Results: In vivo inhibition effect on infarct size, tissue distribution, and pharmacokinetics studies in acute myocardial infarction (AMI) rats showed that ANP Ade/LNCs exhibited better efficiency than non-modified Ade/LNCs and free Ade in all respects.

Conclusion: These results indicated that the ANP Ade/LNCs can be used as a promising system for the treatment of cardiovascular diseases.

Keywords: myocardial infarction, atrial natriuretic peptide, lipid nanoparticles, adenosine, prodrug

Introduction

Myocardial infarction is a major cause of mortality and heart failure worldwide.¹ Compared with the United States or Europe, the incidence of acute myocardial infarction (AMI) increases dramatically in China, which is ~2.5 million AMI patients at present and estimated to be 23 million in 2030.² Therefore, significant clinical effort has been devoted to preventing AMI and reduce the infarction size. Currently, early reperfusion therapies, including thrombolysis and percutaneous coronary intervention, are standard strategies in patients with ST-segment elevation AMI for the reduction of infarction size.^{3,4} However, it is well recognized that reperfusion therapy would induce myocardial ischemia–reperfusion (IR) injury, which further induces calcium overload, impaired glycemic control, increased oxidative stress, and inflammatory response.^{5,6} One of the most effective methods of this injury is direct delivery of cardioprotective drugs to IR myocardium.

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© 2018 ¹ ue tal. This work is published and licensed by Dove Medical Press Limited. The full terms of this license are available at https://www.dovepress.com/terms.php hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission from Dove Medical Press Limited, Previded the work is properly attributed. For permission for commercial use of this work, please see paragraph 4.2 and 5 of our Terms (http://www.dovepress.com/terms.php). Adenosine (Ade) is a ubiquitous endogenous nucleoside that regulates many physiological functions particularly in the heart and brain.⁷ It has most of the IR injury mechanisms, restoring calcium homeostasis, reducing reactive oxygen species (ROS) synthesis, inhibiting the inflammatory response, promoting the preservation of microvascular flow and stabilizing cellular membranes, reducing free radical formation, and promoting angiogenesis following vascular injury.^{6,8–11} However, its systemic administration would bring side effects and limited pharmacological efficiency, and it's extremely short plasma half-life has hampered its clinical application.^{12–14}

Lipid prodrug-based nanosystems are receiving considerable attentions.¹⁵ This strategy provides important benefits: 1) an increase of drug chemical stability in vivo; 2) a sustained drug release; and 3) a reduced toxicity before the metabolization occurs.¹⁶ Lipid covalently conjugated to the drugs contains fatty acids, cholesterol derivatives, phospholipids, and so on.¹⁷ In this study, oleic acid (OA) was chosen as the lipid to conjugate to Ade, so as to achieve sustained release. By combining the lipid prodrug strategy with the use of lipid nanoparticulate systems as drug carriers, oleate adenosine prodrug (Ade-OA)-loaded lipid nanocarriers (LNCs) were developed.

Lipid nanoparticulate systems can be harnessed to deliver therapeutic agents precisely to the infarcted heart by passive targeting.^{18–20} Compared with passive targeting, active targeting due to the specific interaction of the tissuerecognition ligand and receptor may deliver more drugs to the injured tissue than the intact tissue.²¹ Atrial natriuretic peptide (ANP), a member of the natriuretic peptide family, exerts its cardioprotective functions not only as a circulating hormone but also as a local autocrine and/or paracrine factor.²² Research studies^{22,23} have shown that ANP inhibits IR injury and reduces infarct size. Moreover, ANP could be bound to specific natriuretic peptide receptors (NPRs), which are expressed predominantly in the endocardium of the ischemic heart.²³

In this study, ANP-modified and Ade-OA (oleic acid) prodrug-loaded LNCs were fabricated. Ade and OA were conjugated to obtain an amphiphilic prodrug (Ade-OA). ANP was conjugated to distearoylphosphatidyletha-nolamine-polyethylene glycol (DSPE-PEG) by a coupling reaction to get ANP-PEG-DSPE. LNCs were then self-assembled by using the solvent evaporation method. Particle size and morphological analyses revealed uniform spherical shape of NLC. In vitro drug release in the presence of plasma was evaluated. In vivo inhibition effect on

infarct size, tissue distribution, and pharmacokinetics were investigated in rats with ischemic myocardium after intravenous injection.

Materials and methods Materials

Ade (\geq 99%, A9251), ANP rat (amino acid sequence: Ser-Leu-Arg-Arg-Ser-Ser-Cys-Phe-Gly-Gly-Arg-Ile-Asp-Arg-Ile-Gly-Ala-Gln-Ser-Gly-Leu-Gly-Cys-Asn-Ser-Phe-Arg-Tyr, ≥97%, HPLC, A1663), OA (≥99%, GC, O1008), 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC, 39391), dimethyl sulfoxide (DMSO, D 2650), Dulbecco's Modified Eagle's Medium (DMEM, D5796), fetal bovine serum (FBS, F9665), and 3-[4,5-dimethylthiazol-2yl]-2,5diphenyltetrazolium (MTT, M2128) were purchased from Sigma-Aldrich Co. (St. Louis, MO, USA). DSPE-PEG (MW~5kDa)-COOH (DSPE-PEG₅₀₀₀-COOH, PS2-ODE-5K) and DSPE-PEG-FITC (PS2-DEFT-5K) were obtained from Shanghai Pengsheng Biological Co., Ltd. (Shanghai, China). Injectable soyabean lecithin (ISL, TW-SL-IJ01) was provided by Shanghai Taiwei Pharmaceutical Co., Ltd. (Shanghai, China).

Synthesis and characterization of Ade-OA prodrug and ANP-PEG-DSPE ligand

Ade-OA prodrug was synthesized by forming an amide linkage between Ade-NH₂ and OA-COOH (Figure 1A).²³ OA-COOH was dissolved in DMSO and activated by adding EDC·HCl (1.5 equivalents) and NHS (1.5 equivalents) for 2 h at room temperature with stirring (400 rpm). One equivalent of Ade-NH₂ was added to OA-COOH and stirred (400 rpm) for 24 h in the dark at room temperature. Then it was dialyzed against water for 24 h and lyophilized to obtain Ade-OA prodrug.

ANP-PEG-DSPE ligand was synthesized by conjugating APN-NH₂ to DSPE-PEG₅₀₀₀-COOH (DSPE-PEG) by amide bond (Figure 2A).²² The free carboxyl groups of DSPE-PEG were again activated by EDC·HCl (1.5 equivalents) and NHS (1.5 equivalents) for 2 h at room temperature with stirring (400 rpm). ANP was dissolved in phosphate-buffered saline (PBS, pH <7.2) and incubated with DSPE-PEG for 24 h in the dark with stirring (400 rpm) at room temperature, and then it was dialyzed against water for 24 h and lyophilized to obtain ANP-PEG-DSPE.

The chemical structure of Ade-OA and ANP-PEG-DSPE was determined by using ¹H nuclear magnetic resonance (¹H-NMR) analysis by using DMSO-d6 as solvent.



Figure I Synthesis scheme (**A**) and 'H-NMR spectroscopy (**B**) of Ade-OA prodrug. **Note:** Ade-OA was synthesized by forming an amide linkage between Ade-NH₂ and OA-COOH. **Abbreviations:** 'H-NMR, 'H nuclear magnetic resonance; Ade, adenosine; OA, oleic acid.

Preparation and characterization of LNCs

LNCs were then self-assembled by using the solvent evaporation method.²⁴ Briefly, Ade-OA (50 mg) and ISL; (200 mg) were dissolved in acetone (5 mL) (1). ANP-PEG-DSPE (100 mg) was dissolved in acetone–water solution (5 mL, 1:1, v/v) (2). 0.05% poloxamer 188 was dissolved in Milli-Q[®] water (10 mL) (3). Then solutions (1) and (2) were simultaneously and separately added dropwise into solution (3). The mixture was stirred at 600 rpm at room temperature until complete evaporation of the acetone. Redundant stabilizers were removed by centrifugation at $1,000 \times g$ and 4°C for 30 min. The pellet was resuspended in Milli-Q water, vortexed and washed three times, filtered through a 0.45 µm membrane, and adjusted to pH 7.4. The obtained ANP-PEG-DSPE-modified Ade-OA-loaded LNCs (ANP Ade/LNCs, Figure 3) were stored at 2–8°C. ANP-PEG-DSPE-modified, no Ade-loaded



Figure 2 Synthesis scheme (A) and ¹H-NMR spectroscopy (B) of ANP-PEG-DSPE ligand.

Notes: ANP-PEG-DSPE was synthesized by conjugating ANP-NH₂ to DSPE-PEG₅₀₀₀ COOH (DSPE-PEG) by amide bond.

Abbreviations: 'H-NMR, 'H nuclear magnetic resonance; ANP, atrial natriuretic peptide; PEG, polyethylene glycol; DSPE, distearoylphosphatidylethanolamine.



Figure 3 Scheme graph of ANP Ade/LNCs.

Note: ANP Ade/LNCs were self-assembled by using solvent evaporation method. **Abbreviations:** ANP, atrial natriuretic peptide; Ade, adenosine; LNC, lipid nanocarrier; OA, oleic acid; ISL, injectable soyabean lecithin.

blank LNCs (ANP LNCs) were prepared by the same method, using OA instead of Ade-OA. Ade-loaded LNCs without ANP modification (Ade/LNCs) were prepared by the same method, using PEG-DSPE instead of ANP-PEG-DSPE.

Particle size, polydispersity index (PDI), and ζ -potential of each sample were measured at room temperature by Zeta Sizer Nano ZS apparatus (Malvern Instruments, Malvern, UK).²⁵ Samples were prepared in disposable capillary cells without dilution. The measurements were performed under conditions of low ionic strength where the surface charge of the particles can be measured accurately. The average particle size was reflected in volume mean diameter.

The Ade encapsulated in LNCs was separated from the LNCs by dissolving in aqueous HCl (2 M), sonicated for 30 min and stirred for 3 h.⁷ The solution was then centrifuged at 2,000× g for 30 min, the supernatant was collected, and the concentration of Ade was measured at 280 nm by using UV-vis spectrophotometer (UV-1700; SHIMADZU, Kyoto, Japan). The entrapment efficiency (EE) was calculated as (encapsulated Ade in LNCs/mass of the total Ade added) × 100%. The loading capacity (LC) was evaluated by the formula (encapsulated Ade in LNCs/mass of LNCs) × 100%.

In vitro Ade release of LNCs

In vitro release of Ade from the LNCs was studied by dialysis method in a pH 7.4 medium containing 10% FBS.²⁶ Briefly, suspensions of Ade/LNCs and ANP Ade/LNCs were placed

inside a dialysis bag (cutoff 12,000 Da) and stirred at 37°C for 72 h. At predetermined time points, 100 μ L samples were withdrawn and replaced with fresh medium intervals. The Ade concentration in samples was analyzed by the same method mentioned in the "Preparation and characterization of LNCs" section.

Cells

The H9c2 cells (rat cardiomyoblasts) were obtained from American Type Culture Collection (ATCC, Manassas, VA, USA). Cells were cultured in DMEM (Sigma-Aldrich Co., USA) supplemented with 10% fetal FBS (Fisher Chemicals, Fairlawn, NJ, USA) and maintained in a humidified incubator at 37° C and 5% CO₂.

Cellular uptake of LNCs

Fluorescently tagged LNCs were prepared by using DSPE-PEG-FITC instead of DSPE-PEG. H9c2 cells were replaced in 96-well plates with fresh media after 24 h of incubation and treated with fluorescently tagged Ade/LNCs, ANP Ade/ LNCs, and Ade/LNCs. After 4, 24, and 48 h of incubation, the cells were harvested and washed in cold PBS for three times and determined the fluorescence intensity of the cells by flow cytometer (BD Biosciences, San Jose, CA, USA) equipped with a 488 nm argon laser for excitation.²⁷

Cytotoxicity of LNCs

Cytotoxicity of LNCs was determined by using the MTT assay.²⁸ Briefly, H9c2 cells were seeded in 96-well plates at a density of 1×10^4 cells per well with fresh media and incubated for 24 h prior to drug treatment. Subsequently, cells were treated separately with Ade/LNCs, ANP Ade/LNCs, ANP LNCs, free Ade, along with 0.9% saline control and incubated for 72 h. 20 µL PBS containing 5 mg/mL MTT reagent was then added to each well, incubated for an additional 4 h at 37°C. 200 µL of DMSO was used to dissolve the formed formazan crystals, and absorbance was read at 570 nm. Cell viability and half-maximal inhibitory concentration (IC₅₀) was then calculated for each sample.

Animals and AMI model induction

Sprague-Dawley rats (220–240 g) were purchased from Beijing Vital River Laboratory Animal Technology Co., Ltd. (Beijing, China) and fed regular chow, and water was available ad libitum. All experiments were performed in accordance with the Guide for the Care and Use of Laboratory Animals and approved the Medical Ethics Committee of Jining Medical University (no. JNMC201712.2-001). AMI rats were induced as follows:²⁹ rats were anesthetized with a combination of ketamine (40 mg/kg) and xylazine (10 mg/kg), incubated, and mechanically ventilated. The chest was opened by left thoracotomy, the left anterior descending was ligated for 45 min with an intramural 5-0 silk suture, followed by persistent reperfusion. Then chest was closed, and rats could recover.

In vivo effects of LNCs on infarct size

AMI rats were randomly divided into six groups (n=8) that received an intravenous injection of 1 mL of Ade/LNCs (1), ANP Ade/LNCs (2), ANP LNCs (3), free Ade (4), respectively, along with AMI group (AMI rats received physiological saline (5), and sham-operated group (identical surgery except for the coronary artery ligation and injected with physiological saline (6). The hearts were excised and cut into 2 mm thick slices parallel to the atrioventricular groove.³⁰ The slices were immersed in a 1% solution of 2,3,5triphenyltetrazolium chloride at 37°C (pH 7.4) for 15 min and photographed again for the identification of infarct area. The area of AMI was calculated by using ImageJ (National Institutes of Health, Bethesda, MD, USA) software for analyzing the stained or unstained areas of the heart tissue. The size of the infarct area was evaluated as a percentage of whole size of the left ventricle.

In vivo tissue distribution of LNCs

After intravenous injection, rats of Ade/LNCs (1), ANP Ade/ LNCs (2), and free Ade (4) were sacrificed at predefined times (1 and 24 h postadministration).³¹ Various tissues (heart, kidney, liver, brain, and lung) were immediately harvested, kept in saline solution to remove the blood and contents, blotted on filter paper, and then weighed for wet weight. After homogenization using a threefold volume of 0.1 M phosphate buffer (pH 7.4), tissue samples were stored at -20° C for further analysis. Concentrations of Ade in rat plasma were analyzed by the same method mentioned in the "Preparation and characterization of LNCs" section.

In vivo pharmacokinetics of LNCs

After intravenous injection, blood samples of Ade/LNCs (1), ANP Ade/LNCs (2), and free Ade (4) were collected into heparinized centrifuge tubes just before intravenous administration (0 h) and after at 0.25, 0.5, 1, 2, 4, 6, 8, 12, 16, 24, 36, and 48 h.³² We then centrifuged the blood sample at $13,000 \times g$ for 3 min and isolated the plasma. All samples were stored at -20° C until later analysis. Concentrations of Ade in

rat plasma were analyzed by the same method mentioned in the "Preparation and characterization of LNCs" section.

Statistical analysis

The statistical analyses were performed with computer software SPSS 20.0 (IBM Corp, Armonk, NY, USA). Data distribution was tested with Kolmogorov-Smirnov test. Unpaired *t*-test was used to compare 2 independent groups and statistical significances in pharmacokinetics were evaluated using one-way analysis of variance (ANOVA) followed by Tukey's test. The levels of the significant differences were set at probabilities of *P<0.05, **P<0.01, and ***P<0.001.

Results

Characterization of Ade-OA and ANP-PEG-DSPE

¹H-NMR spectrograms of Ade-OA and ANP-PEG-DSPE in DMSO-d6 are presented with point to point of each proton peak according to the structure. Ade-OA (Figure 1B): δ 0.97 (a, -CH₃, OA); 1.35 (b, -CH₂-, OA); 2.06 (c, -OH, Ade); 2.31 (d, -CH₂-CO-N-, amide linkage); 3.51–3.93 (e, f, g, -CH-CH₂-, Ade); 5.45 (h, -CH=CH-, OA); 6.04 (i, -N-CH-O-, Ade); 8.17 (j, -CO-NH-, amide linkage); 8.79 (k, -N-CH=N-, Ade). ANP-PEG-DSPE (Figure 2B): δ 2.49 (a, -CO-O-CH-, DSPE); 2.86 (b, -P-O-CH₂-, DSPE); 3.44 (c, -CH₂-CH₂-O-, PEG); 3.52 (d, -CO-NH-, amide linkage between PEG and DSPE); 6.03 (h, -CO-NH-, amide linkage between ANP and PEG); 4.37, 6.53–8.39 (e, i–n, chemical shifts belong to ANP). The chemical shifts of the amide linkages, Ade, OA, PEG, and DSPE could determine the formation of Ade-OA and ANP-PEG-DSPE.

Characterization of LNCs

The particle size, PDI, ζ -potential, EE, and LC of LNCs were analyzed and are summarized in Table 1. The size of the ANP Ade/LNCs and Ade/LNCs was 168 and 122 nm, respectively. The PDI of the LNCs was <0.2. The ζ -potential of ANP Ade/LNCs was 30 mV; however, the

Table I	Characterization	of LNCs	(mean ± standard	deviation,	n=3)
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LNCs	Particle	PDI	ζ-Potential	EE (%)	LC (%)
	size (nm)		(mV)		
ANP LNCs	166.4±4.2	0.16±0.02	29.5±2.8	N/A	N/A
Ade/LNCs	121.5±3.9	0.13±0.01	15.6±2.1	83.5±3.2	10.2±0.9
ANP Ade/	167.9±4.6	0.17±0.02	30.4±3.1	82.2±3.5	7.1±0.8
LNCs					

 $\label{eq:abbreviations: Ade, adenosine; LNC, lipid nanocarrier; ANP, atrial natriuretic peptide; EE, entrapment efficiency; PDI, polydispersity index; LC, loading capacity; N/A, not applicable.$

 ζ -potential of Ade/LNCs was 16 mV. The EE of Ade-loaded LNCs was >80%. The LC of Ade/LNCs and ANP Ade/LNCs was 10.2 and 7.1%, respectively.

In vitro Ade release of LNCs

The in vitro Ade release profiles (Figure 4) demonstrate that both ANP Ade/LNCs and Ade/LNCs show sustained release behavior. The results illustrate that the release of Ade from ANP Ade/LNCs was slower than Ade/LNCs. Over 80% of drug release was achieved at 24 and 48 h, respectively.

Cellular uptake

Flow cytometry analysis (Figure 5) showed that ANP Ade/LNCs had higher cell uptake compared with Ade/LNCs at 24 (P=0.0309) and 48 h (P<0.01). ANP LNCs also had higher cell uptake compared with Ade/LNCs at 24 (P=0.0293) and 48 h (P<0.01). Higher cellular uptake efficiency was observed in the studied groups along with the time (from 4 to 48 h). There are no significant differences in cell uptake efficiency between ANP Ade/LNCs and ANP LNCs group at all the time points tested (P>0.05).

In vitro cytotoxicity

Cell viability (Figure 6) of the LNCs groups over the concentration range from 1 to 200 μ M was over 80% compared with the controls. These results mean than neither Ade nor LNCs would introduce toxicity to the H9c2 cells when used at concentrations below 200 μ M.





Note: In vitro release of Ade from the LNCs was studied in an aqueous medium containing potassium hydrogen phosphate (pH 7.4, 0.01 M) and 10% FBS. **Abbreviations:** Ade, adenosine; LNC, lipid nanocarrier; FBS, fetal bovine serum; ANP, atrial natriuretic peptide.



Figure 5 Cellular uptake of LNCs determined the fluorescence intensity of the cells by flow cytometer.

Notes: *P<0.05, **P<0.01. Data are presented as mean ± standard deviation, n=6. Abbreviations: Ade, adenosine; LNC, lipid nanocarrier; ANP, atrial natriuretic peptide.

In vivo effects on infarct size

In vivo experiment (Figure 7) showed that the infarct size was significantly inhibited in all the Ade-containing groups than the AMI group (58%). Compared with AMI group, the infarct size was obviously decreased in the free Ade group (47%) (P=0.0426), Ade/LNCs group (39%) (P<0.01), and ANP Ade/LNCs group (27%) (P<0.001). ANP Ade/LNCs group exhibited the strongest efficiency in the inhibition of infarct size.



Figure 6 H9c2 cells cytotoxicity of LNCs determined using MTT assay. **Note:** Data are presented as mean ± standard deviation, n=6. **Abbreviations:** Ade, adenosine; LNC, lipid nanocarrier; ANP, atrial natriuretic peptide; MTT, 3-[4,5-dimethylthiazol-2yl]-2,5diphenyltetrazolium.



Figure 7 In vivo effects of Ade/LNCs, ANP Ade/LNCs, ANP LNCs, and free Ade on infarct size.

Notes: *P<0.05, **P<0.01, and ***P<0.001 compared with AMI group. Data are presented as mean \pm standard deviation, n=8. No data in Sham group.

Abbreviations: Ade, adenosine; LNC, lipid nanocarrier; ANP, atrial natriuretic peptide; AMI, acute myocardial infarction.

In vivo tissue distribution

Tissue distribution of Ade/LNCs, ANP Ade/LNCs, and free Ade are different at 1 (Figure 8A) and 24 h (Figure 8B) post administrations in vivo. At 1 h, Ade/LNC (P=0.0227) and ANP Ade/LNC (P=0.0246) distributions in the heart were higher than free Ade. At 24 h, ANP Ade/LNC distribution in the heart was significantly higher compared with free Ade (P<0.001). Ade/LNCs also showed more obvious heart distribution at 24 h in comparison with free Ade (P<0.01).

In vivo pharmacokinetics

The plasma drug concentration-time profile (Figure 9) illustrated that the free Ade was rapidly cleared from the circulation in the first few hours, while Ade/LNCs and ANP Ade/LNCs exhibited a longer plasma circulation time. The half-lives of Ade/LNCs (2.0 ± 0.4 h) and ANP Ade/LNCs (3.4 ± 0.7 h) were longer than that of the free Ade (0.8 ± 0.1 h) (Table 2). ANP Ade/LNCs exhibited the maximum area under the curve (AUC) of 138.5 mg/L·h. The plasma clearance (CL) and volume of distribution (V) of ANP Ade/LNCs were significantly higher than those of Ade/LNCs and free Ade.

Discussion

Lipid nanoparticulate systems can be harnessed to deliver therapeutic agents precisely to the infarcted heart. In this study, we would like to make a comparison of the lipid prodrug to







Figure 9 In vivo plasma drug concentration-time profile of Ade/LNCs. **Notes:** ANP Ade/LNCs, and free Ade at 0, 0.25, 0.5, 1, 2, 4, 6, 8, 12, 16, 24, 36, and 48 h. Data are presented as mean ± standard deviation, n=8. **Abbreviations:** Ade, adenosine; LNC, lipid nanocarrier; ANP, atrial natriuretic peptide.

cell-membrane-based nanoparticles (NPs). Cell-membranecoated nanocarriers have been explored for their beneficial biocompatibility and modifications through lipid-insertion or chemical conjugation.^{33,34} Cell-membrane-coated surface engineering approach provides a platform to functionalize nanocarriers without sacrificing the natural function of cell membranes. In contrast, lipid prodrug-based nanocarriers provide an increase of drug chemical stability in vivo and a reduced toxicity before the metabolization occurs. The aim of the present study was to fabricate an Ade prodrug-based, ANP-modified nanosystem for the treatment of myocardial infarction. First, Ade-OA prodrug and ANP-PEG-DSPE were synthesized. ANP-PEG-DSPE is an amphiphilic compound constructed by a lipophilic DSPE head and a hydrophilic ANP-PEG tail. The amphiphilic nature of ANP-PEG-DSPE makes it possible to self-assemble to LNCs with a lipid inner

Table 2 Pharmacokinetic parameters after administration in rats(mean \pm standard deviation, n=8)

Parameters	Unit	Ade/LNCs	ANP Ade/LNCs	Free Ade
AUC	mg/L∙h	67.4±2.8**	I 38.5±3.7***,∆	7.8±0.6
AUC 📖	mg/L∙h	68.3±2.6**	I 39.7±4. I ***.∆	8.2±0.7
$T_{1/2}\alpha$	h	0.9±0.2**	I.3±0.3***,∆	0.4±0.1
$T_{1/2}\beta$	h	2.0±0.4**	3.4±0.7***,∆	0.8±0.1
V	L/kg	44.5±2.5**	96.8±2.9***,∆	5.6±0.5
CL	L/h/kg	12.4±1.2**	26.9±2.2***.	1.6±0.3

Notes: **P<0.01, ***P<0.001 compared with free Ade; ^P<0.05 compared with Ade/LNCs.

Abbreviations: Ade, adenosine; LNC, lipid nanocarrier; ANP, atrial natriuretic peptide; $AUC_{0,a}$, total area under the curve; $AUC_{0,a}$ area under the curve until the last time point; $T_{1/2}$, terminal phase half-life; CL, clearance; V, volume of distribution.

core and a hydrophilic shell. The hydrophobic Ade-OA could also be assembled to the lipid core during the preparation process. Larger size of ANP Ade/LNCs than that of Ade/LNCs could be explained by ANP modification on the LNC surface that increased the diameter of the nanocarriers. This could also be taken as an evidence that ANP was successfully modified on the outer layer of the system.³⁵ Particle size is a key effect that can influence the in vivo distribution of carriers. The great advantages of nano-sized carriers include prolonged blood circulation time and improved bioavailability.³⁶ PDI exhibits the size distribution of NPs. PDI <0.2 could be considered as a uniform system. The positive charge of LNCs could exploit the negative charge present at the cell surface, increase the residence time of the carriers, and promote cell penetration, thus the internalization process would be facilitated.³⁷

The in vitro Ade release from ANP Ade/LNCs was slower than Ade/LNCs. This difference may be due to the ANP modification on the surface of LNCs hindered the drug release.³⁸ The reason behind this sustained release behavior is attributed to slow degradation of the nanomaterials, and drugs were diffused from the matrix in a sustained manner. Moreover, the PEG shell on the outside of carriers also protects the drugs and let them have longer release time. In addition, in the presence of serum, the release of drugs was not affected, indicating the stability of LNCs.

Cellular uptake research could provide some circumstantial evidence to display the advantages of the nanocarriers to enter the cells.³⁹ Cell uptake efficiency of ANP-modified LNCs was significantly higher compared with Ade/LNCs. This could be attributed to the ability of ANP modification that promotes the LNCs pass through the cells and accumulate in the cells, and also the PEG could prolong blood circulation time, enhance the solubility of hydrophobic drugs and improve cellular uptake.

The cytotoxicity of any nanoparticulate systems is dependent on several material-related and biological factors, such as cell type and exposure conditions.⁴⁰ For example, the particle size, hydrophobicity, and surface charge can be crucial for the production of ROS and create potential sites of interaction with receptors, leading to variable degrees of cytotoxicity. In the present study, we have evaluated the cytocompatibility of cell viability of the LNCs over the concentration range from 1 to 200 μ M on H9c2 cells that were to elucidate the best non-toxic concentration to be used later in the in vivo studies. In all cases, it was possible to observe that there is no obvious cytotoxicity of the LNCs in the studied concentrations. So, the concentration below 200 μ M can be the safe range for further studies.

In vivo infarct therapy effect was evaluated by measuring the inhibition of the infarct size. Infarct size is conceived as one of critical indices for evaluating the cardiac damage in the generation of ischemic heart disease.⁴¹ To establish the AMI rat model for the evaluation of in vivo infarct therapy effect, coronary artery ligation and isoprenaline injection are the two methods that are commonly used.⁴² Between these two methods, coronary artery ligation is more clinically relevant for it can be used to imitate the clinical status of patients. In this study, compared with the AMI group, the infarct size was obviously decreased in the free Ade group, Ade/LNC group, and ANP Ade/LNC group. ANP Ade/LNC group exhibited the strongest efficiency in the inhibition of infarct size. This may attribute to ANP that could bind to specific NPRs, which are expressed predominantly in the endocardium of the ischemic heart. Moreover, ANP itself could inhibit IR injury and reduces infarct size. In vivo blood circulation half-lives were calculated based on twocompartment model of pharmacokinetics via PKSolver.43,44 In vivo tissue distribution and pharmacokinetics data showed long-circulating characteristics of LNCs than that of free drug, which proved that LNCs elongate the circulation time of drugs in serum and have great potential to accumulate in the myocardial infarct area. ANP-modified LNCs are candidates to target the infarct myocardium in a receptordependent manner.⁴⁵ The aim of the ANP modification is to deliver more ingredient to the infarct zone. ANP Ade/LNCs exhibited higher heart concentration in comparison with Ade/ LNCs in AMI rats which could be due to the targeted ability of ANP used for the modification. ANP Ade/LNCs exhibited higher AUC in comparison with Ade/LNCs and free Ade. This phenomenon could be due to the sustained release of Ade due to the long circulation effect of the PEG.⁴⁶ It was reported that PEG could improve the surface hydrophilicity of nanocarriers and prevented the absorption of lipoproteins and opsonins effectively. Therefore, the conformational clouds of PEG over the nanocarriers could avoid the recognition of the reticuloendothelial system and prolong LNC circulation time in vivo.

Conclusion

In this research, Ade-OA prodrug and ANP-PEG-DSPE ligand were synthesized. ANP-PEG-DSPE-modified Adeloaded LNCs were prepared as a novel drug delivery carrier due to the targeting ability and therapeutic efficiency to AMI. In vivo inhibition effect on infarct size, tissue distribution, and pharmacokinetics studies in AMI rats showed that ANP Ade/LNCs exhibited better efficiency than non-modified Ade/LNCs and free Ade in all respects. These results indicated that the ANP Ade/LNCs can be used as a promising system for the treatment of cardiovascular diseases.

Disclosure

The authors report no conflicts of interest in this work.

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