

ORIGINAL RESEARCH

Enhanced transdermal delivery of lornoxicam by nanostructured lipid carrier gels modified with polyarginine peptide for treatment of carrageenan-induced rat paw edema

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School of Pharmacy, Binzhou Medical University, Yantai, People's Republic of Background: Nanostructured lipid carriers (NLCs) are emerging as attractive drug carriers in transdermal drug delivery. The surface modification of NLCs with cell-penetrating peptides (CPPs) can enhance the skin permeation of drugs.

Purpose: The objective of the current study was to evaluate the ability of the cell-penetrating peptide (CPP) polyarginine to translocate NLCs loaded with lornoxicam (LN) into the skin layers and to evaluate its anti-inflammatory effect.

Methods: The NLCs were prepared using an emulsion evaporation and low temperature solidification technique using glyceryl monostearates, triglycerides, DOGS-NTA-Ni lipids and surfactants, and then six histidine-tagged polyarginine containing 11 arginine (R11) peptides was modified on the surface of NLCs.

Results: The developed NLCs formulated with LN and R11 (LN-NLC-R11) were incorporated into 2% HPMC gels. NLCs were prepared with a particle size of (121.81±3.61)–(145.72±4.78) nm, and the zeta potential decreased from (-30.30±2.07) to (-14.66±0.74) mV after the modification of R11 peptides. The encapsulation efficiency and drug loading were (74.61 ±1.13) % and (7.92±0.33) %, respectively, regardless of the surface modification. Cellular uptake assays using HaCaT cells suggested that the NLC modified with R11 (0.02%, w/w) significantly enhanced the cell internalization of nanoparticles relative to unmodified NLCs (P<0.05 or P<0.01). An in vitro skin permeation study showed better permeation-enhancing ability of R11 (0.02%, w/w) than that of other content (0.01% or 0.04%). In carrageenan-induced rat paw edema models, LN-NLC-R11 gels inhibited rat paw edema and the production of inflammatory cytokines compared with LN-NLC gels and LN gels (P<0.01).

Conclusion: In our investigation, it was strongly demonstrated that the surface modification of NLC with R11 enhanced the translocation of LN across the skin, thereby alleviating inflammation. Keywords: lornoxicam, nanostructured lipid carriers, cell penetrating peptides, transdermal drug delivery, anti-inflammatory effect

Introduction

As a nonsteroidal anti-inflammatory drug (NSAID), lornoxicam (LN) has been used to treat various painful and inflammatory conditions and is thought to display effective analgesic and anti-inflammatory effects by selectively inhibiting cyclooxygenase-2 (COX-2).

The oral tablet and injection of LN are currently used widely to treat pain, rheumatoid diseases and inflammation. 1,2 After oral administration, LN demonstrates

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poor solubility in the acidic conditions of the stomach because of its weak acidity, causing sustained contact with the stomach wall and dyspepsia, ulceration, and anorexia. Additionally, other side effects such as anemia, hepatitis and the first-pass effect may occur upon oral administration of LN.³ Moreover, injection administration is inapplicable to chronic conditions because of its poor safety and compliance for patients.⁴

An alternative route to overcome the side effects may be the transdermal delivery route, which has been developed in the past few years. 5,6 The transdermal application of drugs has been proven to contribute to avoiding first-pass metabolism and preventing gastrointestinal damage, further improving the safety profile and patient compliance.^{5–9} In transdermal drug delivery systems, the stratum corneum (SC), with the largest barrier function, decreases the absorptive amount of active drugs in the skin layers. 10 It is important to ensure that a sufficient amount of active drugs enter the skin or remain in the skin. Additionally, the high active drug levels at specific sites of the skin is beneficial to improve the clinical efficacy in the treatment of inflammation. Therefore, to weaken the skin barrier and further enhance the skin permeation of drugs, several approaches have been developed in recent years, such as chemical 11,12 and physical enhancement methods containing magnetophoresis, 13 iontophoresis, 14,15 electroporation, 16 and microneedles, 17 but their applications are limited because of therapeutic feasibility and toxicity.

An alternative method to enhance drug penetration through the skin is the application of nanoparticles, including solid lipid nanoparticles (SLNs), nanostructured lipid carriers (NLCs), liposomes and nanoemulsions. 18,19 Among these, NLCs containing solid and liquid lipids have been developed as the second generation of lipid nanoparticles superior to SLNs.²⁰ NLCs have been reported to load more drug and prevent it from being expulsed from the carrier because of the generally high solubility of drugs in liquid lipids. NLCs have been reported to demonstrate improved stability, controlled release mode of drugs and good biocompatibility. 21,22 Additionally, NLCs can reduce the skin barrier and improve the lipid solubility of drugs, thus translating the drugs across the skin effectively, as well as protecting the active drugs from degradation.²³ To our knowledge, LN can be incorporated in some carriers, such as transdermal patches,⁵ films,⁶ gels^{9,24,25} and NLCs⁸ for transdermal drug delivery. To further enhance the delivery of NLCs into deep skin layers, cell-penetrating peptides (CPPs) have been investigated for their ability to translocate nanoparticles across the cellular membrane when modified on the surface of NLCs.²⁶ To improve the skin permeability of drugs, various CPPs, such as polyarginine (R8, R9 and R11) peptides, the transactivating transcriptional activator (TAT) peptide, penetratin (PEN) and polylysine-9 (K9), have been used.^{27–29} Among these, polyarginine peptides are widely used in transdermal drug delivery because of the membrane translocating capability produced by the interaction between the positive charges in arginine and negative charges in phospholipids of the cell membrane. Additionally, no study has specifically investigated the potential of polyarginine peptides modified with NLCs loaded with LN.

In the present study, we developed LN-NLCs (containing DOGS-NTA-Ni spacer) modified with six histidine-tagged polyarginine peptides containing 11 arginines, abbreviated as R11. Additionally, the imidazole group in histidine can be specifically bound to the metal-ion-chelating lipid (DOGS-NTA-Ni). Furthermore, a suitable proportion of R11 was selected and assessed for cellular uptake and in vitro skin permeation experiments. Moreover, hydroxypropyl methylcellulose (HPMC) was used as matrices to prepare gel formulations for the application of LN-NLC-R11. The anti-inflammatory effect of LN-NLC-R11 gels was evaluated in the carrageenaninduced paw edema experiment accompanied with the analysis of the pro-inflammatory cytokines.

Materials and methods

Material

LN was purchased from Hubin Pharmaceutical Raw Materials Co., Ltd. (Jiangsu, China). Soybean phospholipids were obtained from Tywei Pharmaceutical Co., Ltd. (Shanghai, China). 1,2-Dioleoyl-sn-glycero-3-[(N-(5-amino-1-arboxypentyl) imidodiacetic acid) succinyl nickel salt] (DOGS-NTA-Ni) was provided by Avanti Polar lipids (AL, USA). The six histidine tagged polyarginine peptides (R11: 6 histidine tag-RRRRRRRRRR) were obtained from China Peptides Co., Ltd. (Shanghai, China). Diclofenac sodium gels were purchased from Health Pharmaceutical Co., Ltd. (Shandong, China). Distilled water was used in the experiment.

Animals

The BALB/c mice (eight-week-old) and Sprague-Dawley rats (180–200 g) provided by the Animal Center of Luye

Pharma Group (Yantai, China) were used in the experiment. The experimental animals were maintained in laboratory conditions (temperature of 20±3 °C and humidity of 60±5 %) for at least a week and were free to food and water. Animal study was performed in accordance with the guidelines of Institutional Animal Care and National Institutes of Health Guide for the Care and Use of Laboratory Animals (USA), and the protocol was approved by the Committee on the Ethics of Animal Experiments of Binzhou Medical University (Permit No. SCXK20140005).

Preparation of nanoparticles

LN-NLCs were prepared using an emulsion-evaporation and low temperature solidification method. Briefly, LN (7.5 mg), glyceryl monostearate (270mg), triglycerides (90 mg), soy lecithin (150 mg), DOGS-NTA-Ni (6 mg) and Tween 80 (720 mg) were dissolved in methanol and then were heated at 80 °C in a water bath with continuous stirring. An aqueous surfactant solution was prepared by dissolving poloxamer 188 (240 mg) in water, followed by heating at 80 °C. Next, aqueous solution was added to the hot lipid phase under a stir rate of 600 rpm to obtain the final mixture (30 mL). The obtained mixture was stirred for 40 min at 80 °C to strengthen the emulsification and evaporate the residual methanol. Thereafter, LN-NLCs were obtained by solidifying the nanoemulsions in an ice water bath for 1 h.

NLCs labeled with the fluorescent dye rhodamine B (RhB-NLCs) were prepared. RhB (0.02% w/w) was added to the lipid phase in the absence of LN. The finally obtained NLC dispersion was dialyzed against ultrapure water for 24 h using a dialysis sack (MWCO =1000) with regular water replacement to remove free RhB.

For surface modification, freshly prepared LN-NLCs were mixed with 5 mg/mL of 6 histidine-tagged R11 water solution in various ratios of DOGS-NTA-Ni and 6 histidine-tagged R11 (2:1, 1:1 and 1:2). The mixtures were incubated for 3 h at room temperature with constant stirring to obtain LN-NLC-R11.

Characterization of nanoparticles Determination of particle size and zeta potential

The particle size and polydispersity indices (PDI) of LN-NLC (with or without R11) were measured in distilled water by dynamic light scattering (DLS) using Zetasizer (Malvern, Nano-ZS, UK). The zeta potential was determined in clear disposable zeta cell.

Surface morphology

The surface morphology of LN-NLC-R11 and NLC-R11 were measured by transmission electron microscopy (TEM; JEM-1230, JEOL, Japan). The suspensions were diluted and placed on a membrane-coated carbon grid surface. After staining with 1% phosphotungstic acid for 1 min, the grid surface was air dried and examined by TEM.

Entrapment efficiency of LN-NLC samples

The encapsulation efficacy and drug-loading percentages in LN-NLC and LN-NLC-R11 were determined by ultrafiltration using vivaspin columns with a filter membrane (MWCO =10,000 Da; Millipore, USA). The LN-NLC suspensions were diluted 50 times in phosphate buffer (pH 7.40), and subsequent dilutions were added to the donor compartment of the vivaspin columns to separate the unencapsulated drug from NLC at 4 °C and 6000 r/min for 20 min using a centrifuge (LR16-A; LAB Centrifuge Co., Ltd., China). The weight of free drug (W_{free}) was obtained according to the concentration of free drug in the receiver compartment. To determine the total amount of drug present in NLCs, 1 mL of LN-NLC suspension was dissolved in 10 mL of methanol via an ultrasonic machine (KQ52003; Kunshan Shumei Ultrasonic Instrument Co., Ltd., China) for 20 min, and subsequent dilutions were made with methanol. The total weight of drug (Wtotal) in the methanol solution and weight of free drug were determined using an UV-spectrophotometer (TU-1901; Beijing Persee General Instrument Co., Ltd., China) at a wavelength of 378 nm. The entrapment efficiency (EE) the drug loading percentage (DL) were calculated as follows:

$$EE \% = (1-W_{free}/W_{total}) \times 100\%$$

DL % =
$$(W_{total} - W_{free})/W_{lipid} \times 100\%$$

W_{lipid}: the weight of total materials used in NLC.

Differential scanning calorimetry (DSC) analysis

To investigate the crystallization behavior of samples, DSC analysis was carried out using a differential scanning calorimeter (TGA/DSC 3+; METTLER TOLEDO, Germany) with a STARe Software System. An appropriate amount (approximate 5 mg) of pure LN, pure glyceryl monostearate and lyophilized LN-NLCs developed in this study were placed in an alumina crucible (METTLER TOLEDO, Germany). The temperature was then heated from 30 to 300 °C with a heating rate of 10 K·min⁻¹ and a nitrogen flow of 20 mL·min⁻¹.

Preparation of gels

The exact amount of HPMC (2 %) was dispersed in warm water with continuous stirring. Next, the LN suspensions (0.05% of LN, w/w) were added slowly into the dispersion of HPMC with continuous stirring to obtain homogeneous dispersion, and LN gels (containing 0.045% LN) were obtained after being kept in the dark overnight to allow complete swelling. The LN-NLC gels and LN-NLC-R11 gels (equivalent to 0.045% LN) were obtained by LN-NLC and LN-NLC-R11 dispersion, respectively, and not the LN suspensions. Additionally, blank NLC gels were prepared without LN.

Physicochemical and rheological characterization of gels

The developed LN-NLC-R11 gels, LN-NLC gels and LN gels were characterized by their appearance, homogeneity, pH and rheological property. The appearance and homogeneity of the gels were inspected visually.³⁰

The pH of the gels was measured using a pH meter (EL20; Mettler Toledo, USA) after diluting the gels (1 g) in 20 mL of distilled water.³¹

The viscosity of LN-NLC-R11 gels was determined using a digital viscometer (SNB-1; Shanghai Hengping Instrument Co. Ltd, Shanghai, China) at room temperature and varying the shear speeds (6–60 rpm) with a number 4 spindle. The rheological property of the gels was described by the flow curve of the viscosity versus shear speed.

Stability studies of gels

The stability test was performed to investigate the stability of LN-NLC-R11 gels. Three batches of LN-NLC-R11 gels were packed into tubes (with plugs) and stored at room temperature for one month. The samples were then observed periodically for changes in appearance, homogeneity, pH and viscosity.

In vitro cellar uptake study

Cell culture

Human hyperproliferative keratinocyte (HaCaT) cell lines were purchased from the BeNa Culture Collection (Beijing, China). The cells were cultured in Dulbecco's modified Eagle's medium (DMEM) (high glucose) with 10% (v/v) fetal bovine serum (FBS) at 37 °C and in a 5% CO_2 atmosphere.

Cytotoxicity study

The cytotoxicity of NLC, LN-NLC, NLC-R11 and LN-NLC-R11 to HaCaT cells was determined utilizing the

MTT (3-(4,5-dimethylthoazol-2vl)-2,5-diphenyl-2H-tetrazolium bromide) assay. HaCaT cells were seeded in a 96well plate at a density of 10,000 cells/well and were allowed to incubate with different formulations at various concentrations of LN for 72 h at 37 °C. Next, the cells were washed with PBS (pH 7.4) and cultured with fresh DMEM (no FBS) containing MTT for 4 h at 37 °C, and then the culture medium was removed and replaced with 200 mL of DMSO to dissolve the formazan crystals in cells. The plate was shaken for 10 min in the dark. The absorbance was measured at 490 nm using a microplate spectrophotometer (SpectraMax M2; Molecular Devices, Sunnyvale, CA, USA). To obtain a background reference, wells without seeded cells were employed, and control wells were obtained using cells treated without samples. The cell viability was calculated as follows:

Cell viability (%) =
$$(A_t - A_b)/(A_c - A_b) \times 100\%$$
.

Where A_t , A_b and A_c are the absorbance of tested samples, background reference and control wells, respectively.

In vitro cellar uptake study Flow cytometry (FCM)

The in vitro cellar uptake assay of RhB-NLC and RhB-NLC-R11 (the method of preparation was as described in Preparation of nanoparticles) was studied in HaCaT cells by flow cytometry. Cells were seeded at a density of 5×10^4 cells/well in six-well culture plates and were cultured overnight. Next, the cells were incubated with different formulations (equivalent to 6 µg/mL, 12 µg/mL and 25 µg/mL of RhB) for 1, 2 and 4 h at 37 °C. After coculture, cells were washed with phosphate-buffered saline (PBS, pH 7.4) to remove extracellular particles. The associated fluorescence in cells was determined using flow cytometry (Epics XL; Beckman Coulter, Brea, CA, USA) at an excitation wavelength of 590 nm.

Additionally, the connection of the peptide (six histidine-R11 peptide) to DOGS-NTA-Ni was determined by flow cytometry. Briefly, cells were incubated with RhB-NLC-R11 and RhB-NLC (no DOGS-NTA-Ni) mixed with R11 for 2 h. After incubation, the cells were washed with PBS (pH 7.4) and the associated fluorescence was determined using flow cytometry.

Confocal laser scanning microscopy (CLSM)

Confocal laser scanning microscopy (CLSM) was employed to observe the location of RhB-NLC and RhB-NLC-R11 in

HaCaT cells. HaCaT cells were seeded in six-well plates at a density of 5×10^4 cells/well. Next, the cells were cultured with different samples (equivalent to 6 µg/mL, 12 µg/mL and 25 µg/mL of RhB). After incubation for 1, 2, and 4 h at 37 °C, the cells were fixed with paraformaldehyde, and Hoechst 33342 (10 mg/mL) was used to stain cell nuclei. HaCaT cells were then subjected to confocal laser scanning microscopy (TCS SPE; Leica Microsystems, Wetzlar, Germany) at an excitation wavelength 530 nm and an emission wavelength of 590 nm.

Mechanism of RII as a penetration enhancer

The mechanism of the cell uptake of R11 was investigated in HaCaT cells by determining the concentration of Ca²⁺ and change in the membrane potential.

Measurements of the membrane potential in HaCaT cells. The membrane potential in HaCaT cells was monitored using the fluorescence dye bis-(1,3-dibutylbarbituric acid) trimethine oxonol [DiBAC4 (3)] (Dojindo, Kumamoto, Japan), which was sensitive to the membrane potential. Briefly, the cells were cultured with LN-NLC (12 μ g/mL) and LN-NLC-R11 (12 μ g/mL) for 24 h in an incubator at 37 °C and 5% CO₂. Next, the cells were digested with pancreatin and were washed with PBS to remove the samples. The solution of DiBAC4 (3) (5 μ g/mL) was added and incubated with the cells for 0.5 h. The mean fluorescence intensity was measured using flow cytometry (Epics XL; Beckman Coulter, Brea, CA, USA) at an excitation wavelength of 488 nm and an emission wavelength of 530 nm. Cells without treatment were used as zero adjustment.

Measurement of the Ca²⁺ concentration of HaCaT cells

The changes in the intracellular Ca²⁺ concentration were determined using the calcium-sensitive fluorescent probe derivative fluo-3 AM, which easily penetrates cell membranes. Fluo-3 AM is cleaved by esterase in cells to fluo-3, which presents bright fluorescence after combining Ca^{2+,32} The fluorescence intensity of the cells cultured with LN-NLC (12 μg/mL) or LN-NLC-R11 (12 μg/mL) was measured by flow cytometry (Epics XL; Beckman Coulter, Brea, CA, USA) at excitation wavelength of 488 nm and an emission wavelength of 525 nm, and cells without treatment were used as the blank control. Briefly, after the incubation of cells with LN-NLCs and LN-NLC-R11 for 24 h at 37 °C and 5% CO₂, the cells were washed 3 times with PBS. Next, the cells were incubated in culture medium containing Fluo-3 AM (5 μmol/L) for 0.5 h. The

intracellular fluorescence intensity was measured using flow cytometry.

In vitro skin permeation studies Preparation of the isolated skin of mice

The mice were sacrificed by cervical dislocation, and abdominal hairs were removed carefully using an electric shaver. The skin from the abdominal surface was excised, and the adherent fat and subcutaneous tissue were removed carefully. Next, the isolated skin was washed with phosphate-buffered saline (PBS, pH 7.40), and its integrity was assured during the experiment.

In vitro skin permeation studies

The percutaneous permeability of LN-NLCs and LN-NLCs loaded with various amounts of cell penetrating peptide R11 (0.01%, 0.02% and 0.04% respectively) was investigated in in vitro skin permeation studies using the Franz diffusion method.

Briefly, 1 mL (0.75 mg of LN) of LN-NLC or R11-coated LN-NLC formulations was applied onto the *stratum corneum* (SC), which faced the donor compartment of Franz diffusion cells. The receptor chamber was filled with 16 mL of PBS (pH 7.40) under the condition of 400 r/min and 37±0.5 °C to maintain the physiological activity of the skin. The occlusive condition was sustained to prevent water from evaporating throughout the experiment. To draw the percutaneous permeability curve, all receiver fluid was collected and the fresh PBS (pH 7.40) with an equal volume and temperature was added directly into the receptor compartment at predetermined time intervals. The content of LN in receiver fluid was measured using HPLC (Agilent 1260) and the cumulative amount of permeated LN was calculated as follows.

$$Q = \frac{\sum_{i=1}^{n} C_n \times V}{A}$$

Here, Q (μ g/cm²) is the cumulative penetration amount per cm², C_n (μ g/mL) is the LN concentration of the *n*th sample, V (mL) is the volume of the receiver fluid, and A (cm²) is the infiltrating area.

Drug extraction from the skin

After 24 h of skin permeation, the fluid in the receptor compartment was collected, and the surface of the mouse skin was washed carefully with PBS (pH 7.4) to remove excess formulations and sucked dry. Thereafter, the mouse skin dosing area was cut off and then cut into pieces to a mixture with 0.4 mL of PBS (pH 7.4). After

homogenization in the vortex, 0.6 mL of methanol was added to the mixture. The mixture was then disposed using an ultrasonic cell disruptor (Scientz-II D; Ningbo Scientz Biotechnology Co., Ltd., China) for 10 min to destroy skin tissue maximally. Subsequently, the mixture was centrifuged at 13,500 rpm for 10 min to collect the supernatant, and then the residues were mixed again with 0.5 mL of methanol, homogenized and subjected to centrifugation as described above. After 3 rounds of homogenization and centrifugation, the total supernatant was analyzed using HPLC to determine the drug content, and the skin deposition of LN per unit area Q_s (µg/cm²) was calculated

HPLC analysis

HPLC analysis of LN was performed using a reversed-phase C18 analytical column (5 μ m; 4.6×250 mm) and an ultraviolet (UV)-visible variable wavelength detector. The mobile phase comprised methanol, and an aqueous solution containing 0.05% trimethylamine (52:48, v/v) was used to detect LN, with a flow rate of 1.0 mL/min. The drugs in the samples were detected at a column temperature of 25 °C, an injection volume of 20 μ L and a wavelength of 378 nm.

Skin irritation test

Six New Zealand white rabbits were used to evaluate the skin irritation potential of LN-NLC-R11 gels. Hairs on the backside area of rabbits were removed by hair clippers 24 h prior to experiment. LN-NLC-R11 gels were smeared uniformly on the left side of hairless skin once a day for 7 days, and no treatment was carried out on the right side of hairless skin. The skin was observed for any response, including redness and erythema, after the application of LN-NLC-R11 gels. In the skin irritation test, we referred to the scoring criteria in the literatures. 33,34 The scores of erythema and edema were recorded (ranging from 0 to 4), and the scores were called as the irritation index. It was calculated to evaluate the degree of irritation. An irritation index below 0.4 indicates negligible irritation.

In vivo anti-inflammatory activity of the formulations of LN

Experimental design to analyze pharmacological activity

Before the implementation of the experiment, 40 healthy Sprague-Dawley rats were discretionarily sectionalized into 5 groups (n=8). These groups were administered

either blank NLC gels (A, negative control), diclofenac sodium gels (B, positive control), LN gels (C), LN-NLC gels (D) and LN-NLC-R11 gels (E). The dose of drug administered to mice was 2 mg/kg in every group.

Carrageenan-induced rat paw edema test

The study of paw edema induced by carrageenan was carried out to explore the anti-inflammatory activity of different formulations. Prior to 1 h subcutaneous injection of carrageenan (1% w/v) with 0.1 mL into the middle part of the right hind paw, the samples were smeared on the right hind paw at an established dosage every 1 h for 3 h. The paw volume at the marked position was measured using a toe volume meter (PV-200; Chengdu Techman Software Co., Ltd, China) at 0, 1, 2, 3, 4, 5, 6 and 7 h after the application of 1% carrageenan. Paw edema was presented as an average increase in the paw volume at certain positions.

Estimation of pro-inflammatory cytokines and chemokines in rat serum

At the period of severe inflammation (3rd h of the experiment), blood samples from the rats were collected and serum was obtained by centrifugation (3K15; Sigma, Germany) at 2,000 rpm for 20 min at 4 °C. The serum was stored at -80 °C before determination, and the levels of IL-6, cyclooxygenase-2 (COX-2) and inducible nitric oxide synthase (INOS) in rat serum were determined using enzyme-linked immunosorbent assay (ELISA) (Shang Hai Lengton Bioscience Co., LTD, China). 35

Statistical analysis

One-way analysis variance (ANOVA) followed by the Fisher's least significant difference (LSD) test was used to compare two different groups of samples using SPSS version 19.0 software (IBM Corp., Armonk, NY, USA). Additionally, the data were expressed as the means \pm standard deviation. A *P*-value<0.05 was considered to indicate a significant difference between groups.

Results and discussion

Characterization of nanoparticles and gels Characterization of nanoparticles

The mean particle size of LN-NLCs was found to be 121.81 ± 3.61 nm, with a polydispersity index (PDI) of 0.26 ± 0.01 . The mean particle size of LN-NLC-R11 was increased to 145.72 ± 4.78 nm (with a PDI of 0.27 ± 0.01) because of the surface

modification of LN-NLCs (Figure 1A). The zeta potential of LN-NLCs in double-distilled water was -30.30 ± 2.07 mV and further decreased to -14.66 ± 0.74 mV (Figure 1B) after surface modification with R11 because of the positive charges of the guanidine group in poly-arginines. The zeta potential is often considered to indicate whether poly-arginines have been

modified on the surface of nanoparticles because the zeta potential of the nanoparticles turns positive when poly-arginines are successfully modified.^{36,37}

The images (Figure 1C) confirmed the formation of uniformly spherical particles regardless of the surface modification.

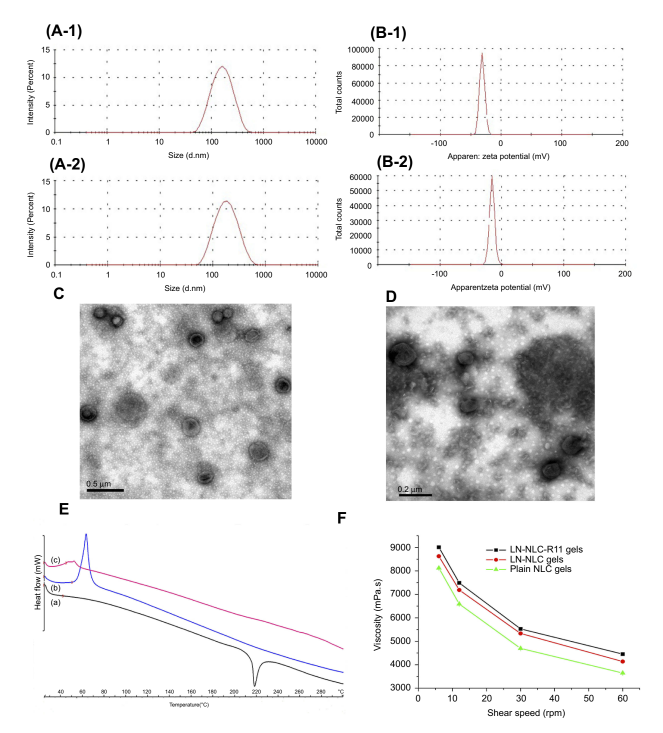


Figure 1 Pharmaceutical characterization of nanoparticles and gels.

Notes: (A-I), (A-2) Size distribution of LN-NLC and LN-NLC-RII, respectively; (C and D)

TEM imagines of LN-NLC-RII and NLC-RII; (E) DSC thermograms of LN (A), glyceryl monostearate (B) and LN-NLC (C); (F) Flow curves of formulation gels. Data presented as mean ± SD (n=3).

The entrapment efficiency of LN in LN-NLCs was (74.61 ± 1.13) %, and the drug-loading percentage was (7.92 ± 0.33) %. The entrapment efficiency and drug loading percentages of LN in LN-NLCs were unaffected by surface modification.

The thermodynamic properties of LN, glyceryl monostearate and LN-NLCs were analyzed by DSC, and the DSC thermograms of different formulations are shown in Figure 1E. A single endothermic peak at 219 °C was observed in the DSC curve of LN, which agrees with its melting point, and the crystalline nature of LN was confirmed. The exothermal peak of glyceryl monostearate was found at 63 °C. Moreover, the melting endotherm of LN does not occur in the DSC thermogram of LN-NLC, indicating that the drugs were encapsulated completely inside the lipid matrix of NLCs in the amorphous form.

Pharmaceutical characterization of gels

The formulation gels prepared in this study presented slightly opaque and good homogeneity.

The pH values of LN formulation gels were in the range of 7.0–7.2, which would likely not produce irritation. Thus, they were considered suitable for skin application.³⁸

The rheological properties of gels are closely related to the ease of application to the skin by affecting the adhesion and retention for topical application. The viscosities of LN-NLC-R11 gels (4452±12 mPa·s) and LN-NLC gels (4136±8 mPa·s) were higher than those of LN gels (3645±8 mPa·s) at a constant shear speed of 60 rpm. The cause might be that the incorporation of nanoparticles into the gel base increased the viscosity of gels.³⁹ The flow curves shown in Figure 1F indicated that the gels revealed a nonnewtonian, pseudo plastic flow pattern (increasing shear speed with a decrease in viscosity), maximizing the area coverage on application. Thus, it is an important characteristic for topical formulations.^{40,41}

Stability studies of gels

As shown in Table 1, the LN-NLC-R11 gels were found to be stable at room temperature for one month, no stratification was observed, and the appearance of LN-NLC-R11 gels was

still slightly opaque with good homogeneity. No significant change was observed in the pH and viscosity of gels, indicating that the LN-NLC-R11 gels developed in this study had good storage stability at room temperature for one month.

Cytotoxicity of nanoparticles

The cytotoxicity of NLC, LN-NLC, NLC-R11 and LN-NLC-R11 on HaCaT cells was evaluated by MTT assays. As demonstrated in Figure 2, the blank NLC formulations resulted in cell viabilities greater than 90%, indicating that the blank NLC had good biocompatibility to HaCaT cells before and after R11 peptide coating. Upon LN loading, the cell viability decreased as the LN concentration increased, and LN-NLCs coated with R11 peptide presented lower cell viability than uncoated LN-NLCs because of the enhanced cell permeability of the R11 peptide.

In vitro cellar uptake study Flow cytometry

Flow cytometry experiments were carried out to quantify the intracellular accumulation of samples. Figure 3 shows the accumulation of RhB-NLC-R11 in cells in the presence of various amounts of R11. When the cells were exposed to RhB-NLC-R11, the fluorescence intensity in cells was higher than that in cells exposed to RhB-NLC, indicating that the DOGS-NTA-Ni spacer (0.02% in NLC, w/w) is on the surface of NLCs, resulting in the increased R11 peptide binding affinity to NLCs. Moreover, when the ratio of R11 to DOGS-NTA-Ni was 1:1 (w/w), the fluorescence intensity was higher than that of the other two ratios (1:2 or 2:1). The results indicated that a further increase in the R11 content did not induce a further increase in the fluorescence intensity in cells. Moreover, no obvious increase in the fluorescence intensity in cells was observed after the coincubation of cells with RhB-NLCs mixed with R11 without the DOGS-NTA-Ni spacer, indicating no specific interaction between NLCs and the R11 peptide (Figure 3A–C). Therefore, the ratio (1:1 w/ w) of DOGS-NTA-Ni to the R11 peptide was adopted in further studies.

Table I Stability studies of LN-NLC-R11 gels at room temperature for one month

Parameter	0 d	10 d	20 d	30 d
Appearance	Slightly opaque	Slightly opaque	Slightly opaque	Slightly opaque
Homogeneity	Good	Good	Good	Good
pН	7.09±0.04	7.08±0.02	7.10±0.04	7.11±0.03
Viscosity (mPa 's)	4452±12	4450±11	4447±11	4443±10

Notes: Data presented as mean \pm SD (n=3).

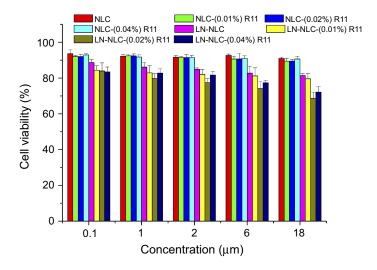


Figure 2 Cell viability of HaCaT cells incubated with different formulations with different concentration of LN for 72 h.

Notes: The concentration of NLC without LN was calculated corresponding with LN-NLC. Data presented as mean ± SD (n=3).

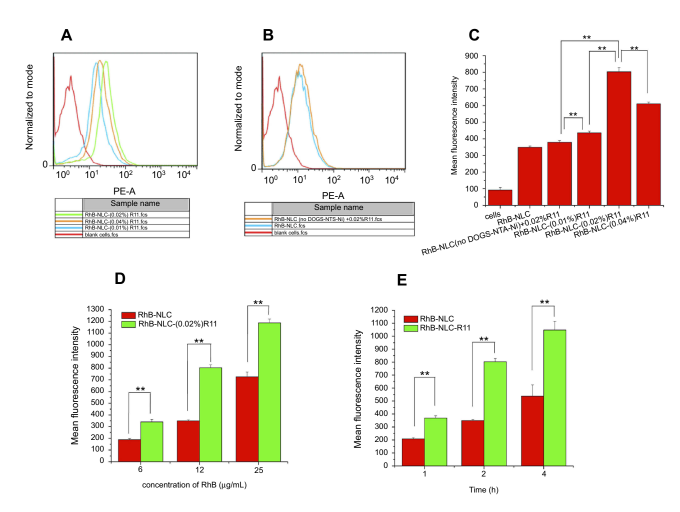


Figure 3 Cell uptake of different formulations in HaCaT cells analyzed by flow cytometry.

Notes: Flow-cytometry analysis (A and B) mean fluorescence intensities (C) of different formulations in HaCaT cells. (D and E) Mean fluorescence intensities of RhB-NLC and RhB-NLC-(0.02%) R11 incubated with HaCaT cells in various concentrations and time. **P<0.01. Data presented as mean ± SD (n=3).

As shown in Figure 3D and E, the mean fluorescence intensity (MFI) inside HaCaT cells increased with a longer time and higher concentrations, demonstrating that HaCaT cells showed time-dependent and concentration-dependent uptake of RhB-NLCs with or without R11. Additionally, the results in Figure 3 D and E) showed that the MFI of HaCaT cells treated with RhB-NLC-R11 was much higher than that of cells treated with RhB-NLCs at each time or concentration point.

Confocal laser scanning microscopy imaging

The uptake effect of RhB-NLC-R11 in HaCaT cells was visualized using CLSM. As shown in Figure 4, Hoechst 33342-stained nuclei appeared blue, and RhB labeled NLC (R11) appeared red. Higher rhodamine fluorescence intensity was observed at longer incubation times (1 h to 4 h) and higher concentrations of rhodamine (6 μg/mL to 25 μg/mL) (Figure 4). As shown in Figure 4, the redder fluorescence was observed in the cytoplasm of HaCaT cells incubated with RhB-NLC-R11 at 2 h than in the cytoplasm of RhB-NLC. The HaCaT cells incubated with RhB-NLC-(0.02%) R11 showed brighter florescence (red) than with other formulations, indicating that NLC modified with R11 at the

ratio (1:1 w/w) of DOGS-NTA-Ni to R11 peptide showed enhanced cellular permeability. The results were consistent with the quantitative uptake measurements of FCM. Therefore, NLC-R11 is expected to be a suitable system to deliver LN to the post stratum corneum with enhanced cellular uptake.

Study of the mechanism of RII as a penetration enhancer

Effects of LN-NLC and LN-NLC-R11 on the membrane potential of HaCaT cells

The HaCaT cell line derived from the human abdominal epidermis is an important model to study the human skin and is used widely as a substitute for normal human epidermal keratinocytes. The membrane potential plays a key role in the biophysical characteristics of the cell membrane, maintaining the balance of the ion concentration between the inside and outside of the cells. In general, the membrane potential changed accompanied by the change in the membrane permeability and structure of the membrane lipid and protein.42-44

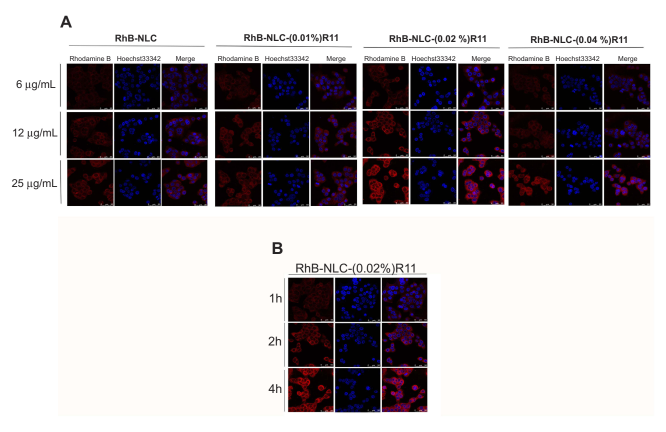


Figure 4 Confocal laser scanning microscopy images of HaCaT cells incubated with different formulations with various concentrations (A) and incubated with RhB-NLC-RII in different time (B).

As shown in Figure 5, the mean fluorescence intensity of cells treated with LN-NLC-R11 was higher than that in cells treated with LN-NLCs, indicating that LN-NLC-R11 significantly decreased the membrane potential of HaCaT cells compared with LN-NLC. The negative charge inside the cell membrane and positive charge outside the cell membrane are maintained in the resting state. Nanoparticles could induce depolarization of the membrane when acting on the cell membrane. The more fluorescent probe DiBAC4 (3) with a negative charge could be transported across the membrane with increased depolarization of the membrane, leading to a decreased membrane potential. Furthermore, the decrease in the membrane potential is beneficial for nanoparticles to penetrate the cell membrane.

Change in the intracellular ca²⁺ concentration induced by RTI

Figure 5 shows significant enhancement of the intracellular fluo-3 fluorescence in cells incubated with LN-NLC-R11, reflecting the corresponding increased Ca²⁺ concentration compared with cells cultured with LN-NLCs, indicating that R11 may enhance the permeability of cell membranes via changing the degree of intercellular Ca²⁺ concentration. Particularly, intercellular Ca²⁺ is a crucial second messenger that regulates signal transduction and other intracellular processes. Changes in the intracellular Ca²⁺ concentration could induce changes in the potential, fluidity and degree of tight junctions of the cell membrane, further influencing some cellular processes, such as endocytosis and micropinocytosis.⁴⁵

In vitro skin permeation of nanoparticles

The results in Figure 6 show the effect of the surface modification of NLC on the skin permeation of LN. LN-NLC-R11 significantly increased the cumulative penetration amount and skin deposition of LN (P<0.01 or P<0.05) compared with LN-NLC alone. The cause may be due to the interaction of the positive charge in R11 with the negative charge in lipids and residues of proteins of SC, further inducing the destabilization of the lipid bilayer and membrane and leading to enhanced permeation of NLCs. 46 Furthermore, the negatively charged NLCs obtained in this study showed less affinity to the skin. After modified with polyarginine peptides, the zeta potential was reduced to -14 mV from -30 mV, which induced an increased affinity of the nanoparticles to the skin. Additionally, similar results were concluded in other studies. 47,48 It was also found that CPP-conjugated nanoparticles penetrate through the skin by some appendages such as hair follicles and sebaceous glands. 49 Although the enhanced permeation ability of CPPs is achieved by various pathways, such as diffusion through intercellular lipids and skin appendages, the amount of CPPs modified on NLCs could influence the permeating efficiency. As shown in Figure 6A and B, especially with LN-NLC modified with 0.02% (w/w) R11, the cumulative penetration amount and retention in skin of LN was approximately 2.0 and 1.7 times higher than that with LN-NLC alone, respectively, indicating the greater enhanced permeation effect than with LN-NLC modified with 0.01% (w/w) R11 and 0.04% (w/w) R11. The cause may be that the redundant R11 had no

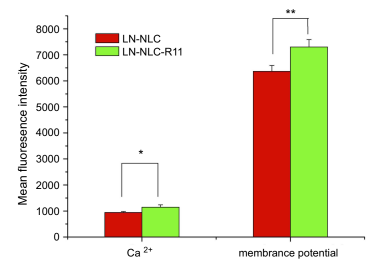


Figure 5 The results of Ca^{2+} concentration and membrane potential in HaCaT cells. **Notes:** Data presented as mean \pm SD (n=3). *P<0.05, **P<0.01.

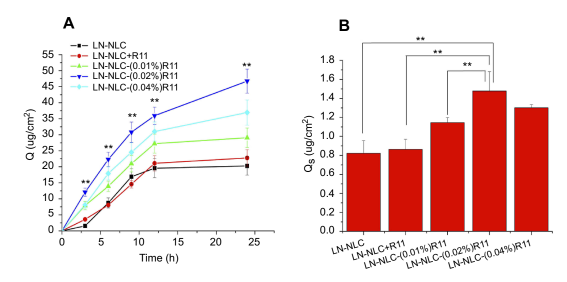


Figure 6 In vitro percutaneous permeability and retention in skin of LN.

Notes: (A) In vitro cumulative penetration amounts of lornoxicam per cm² (Q) versus time profiles in different formulations, **P<0.01, compared LN-NLC; (B) The skin drug deposition (Q_s) of different formulations in transdermal permeation, **P<0.01. Data presented as mean \pm SD (n=3).

binding affinity to the surface of NLCs, limiting the space in which the nanoparticles penetrate the skin. Furthermore, NLCs prepared without DOGS-NTA-Ni in the presence of six histidine-tagged polyarginine peptides did not significantly increase the amount of LN in or through skin, indicating that no nonspecific interaction occurred between the nanoparticles and six histidine-tagged polyarginine peptides. Thus, NLCs prepared with 0.02% (w/w) R11 and DOGS-NTA-Ni (R11:DOGS-NTA-Ni=1:1, w/w) allowed NLCs to penetrate the skin, leading to the maximum skin permeation of LN.

Skin irritation evaluation

In the skin irritation test, no erythema and edema were observed in rabbits, the mean irritation index was 0, and no difference was found between the left side and right side of hairless skin. Thus, LN-NLC-R11 gels exhibited no irritation for topical application and will be safe for patients.

Anti-inflammatory effect of the formulations of LN

Carrageenan-induced rat paw edema

To evaluate the anti-inflammation effect of various formulations of LN, the carrageenan-induced paw edema method was adopted, and the results are presented in Figure 7A. As shown in Figure 7 (A), carrageenan-induced edema was alleviated significantly after the administration of LN-NLC gels (D) and LN-NLC-R11 gels (E) at the dosage of 2 mg/kg compared with blank NLC gels (A) and LN gels (C), respectively

(*P*<0.05 or *P*<0.01), demonstrating the good activity of NLC as the LN carrier in inhibiting acute inflammation.

Moreover, LN-NLC-R11 (E) gels (2 mg/kg of LN) significantly inhibited the paw edema of rats compared with LN-NLC gels (D) (2 mg/kg of LN) at various time points (P<0.01), indicating that the cell-penetrating peptide R11 on the surface of LN-NLCs increased the local drug concentration by enhancing the percutaneous permeability of LN-NLCs, leading to an obvious anti-inflammatory effect. Additionally, inhibition of the paw edema of LN-NLC-R11 (E) gels was higher than that of diclofenac sodium gels (B) during severe inflammation (3 h-5 h, P<0.05) and was similar to that during other periods, indicating the excellent anti-inflammatory effects of LN-NLC-R11 (E) gels. Moreover, with the extension of time, especially 5 to 7 h after inflammation, LN-NLC-R11 gels (E) manifested sustained inhibition of paw edema, showing that LN-NLC-R11 gels not only promoted drugs through the skin but also extended the action time of drugs to achieve an anti-inflammatory effect in 7 h. The edema degree, edema percentage and inhibitory rate of edema were calculated as follows:

Edema degree =
$$V_{right} - V_{left}$$

Edema percentage =
$$(V_{right}-V_{left})/V_{left} \times 100\%$$

Inhibitory rate of edema =
$$(E_A - E_{test})/E_A \times 100\%$$

Where V_{right} is the mean volume of the right hind paw, V_{left} is the mean volume of left hind paw, E_{A} is the mean

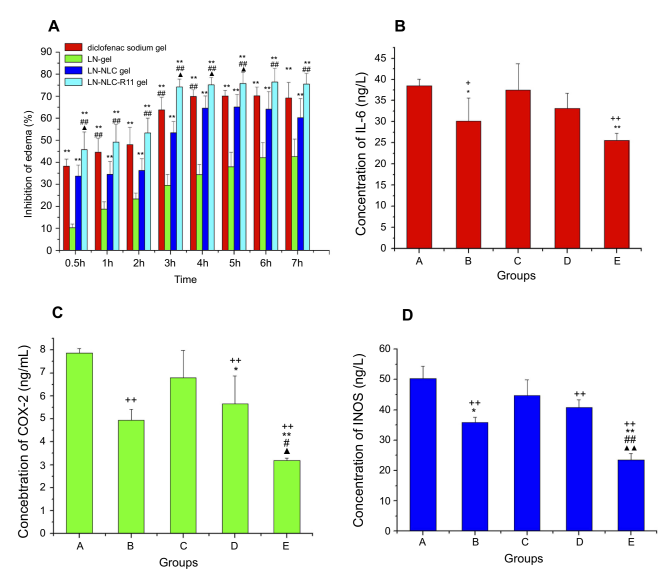


Figure 7 Anti-inflammatory activity tested by carrageenan-induced paw edema.

Notes: (A) The inhibition of various formulations on carrageenan-induced paw dema in rats. (**P<0.01, as compared to LN-gel; **#P<0.01, as compared to LN-gel; **#P<0.01, as compared to LN-gel; **P<0.05, as compared to diclofenac sodium gel). (B-D) The inhibition of various formulations on pro-inflammatory cytokine and chemokines (IL-6, COX-2, INOS) content in rats. (*P<0.05, **P<0.01, compared to blank NLC gels; *P<0.05, **P<0.01, compared to LN-NLC gels; *P<0.05, **P<0.01, compared to LN-NLC gels; *P<0.05, **P<0.01, compared to LN-NLC gels; *P<0.05, **P<0.01, compared to Blank NLC gels; *P<0.05, **P<0.01, compared to LN-NLC gels; *P<0.01, compared to LN-NLC gels; *P<0.05, **P<0.01, compared to LN-NLC gels; *P<0.01, compared to LN-NLC gels; *

edema percentage of negative control (group A) and E_{test} is the mean edema percentage of test groups (B, C, D and E).

Estimation of pro-inflammatory cytokines and chemokines in rat serum

As a nonsteroidal anti-inflammatory drug (NSAID), LN reduces swelling in inflamed tissues.⁵⁰ The principal mechanism of its action is via the inhibition of prostaglandin synthesis, which is mediated by the enzyme COX in the arachidonic acid pathway.^{51,52} The COX isoform COX-2 is produced during inflammation, resulting in the formation of excessive prostaglandins. Moreover, other

inflammatory mediators such as reactive oxygen products and cytokines have been found to considerably induce inflammation. ^{53,54} The overexpression of inducible nitric oxide synthase (iNOS) results in the increase of nitric oxide (NO) in inflamed tissues, further inducing edema in inflammatory sites. ⁵⁵ Among the pro-inflammatory cytokines, interleukin (IL)-6, which is relevant to the formation of inflammatory mediators, could be effectively reduced by LN. ⁵⁰ As the well-established inflammatory cytokine and enzymes, serum IL-6, COX-2 and INOS play crucial roles in the pathology of inflammation and are determined to assess the degree of inflammation. ^{56,57}

Therefore, the effect of the developed different formulations in reducing inflammatory symptoms was investigated by determining markers such as IL-6, COX-2 and INOS in this study.

Figure 7B-D shows the levels of IL-6, COX-2 and INOS in the rat serum of different formulations, respectively. Significant decreases in the serum levels IL-6, COX-2 and INOS in rat models were observed after the administration of LN-NLC-R11 gels (E) compared with those in the negative control (P<0.01) and LN gels (P<0.01) in Figure 7B-D, indicating that LN-NLC-R11 gels could effectively alleviate the severity of inflammation by reducing the serum content of inflammatory cytokines and relevant enzymes—ie, IL-6, COX-2 and INOS. The serum COX-2 and INOS levels of LN-NLC gels were significantly different from those of the negative control (P<0.01), although no significant differences were found in the serum IL-6 level among the negative control, LN gels and LN-NLC gels. LN-NLC-R11 gels showed no significant difference from the marketed product (diclofenac sodium gels) at the serum IL-6 level but showed a higher inhibition effect than the marketed product at the serum COX-2 and INOS levels (P<0.05 or P<0.01), indicating their excellent effect treat inflammation.

Conclusion

The present study suggested that LN-NLCs were prepared successfully by an emulsion-evaporation and low temperature solidification method and were characterized by an appropriate size, high entrapment efficiency and drug loading. To enhance the transdermal permeability of LN, R11 was used in the formulation to prepare LN-NLC-R11. FCM and CLSM studies suggested that the NLC modified with R11 (0.02%, w/ w) had enhanced cell internalization of nanoparticles. The results of the in vitro permeation study demonstrated that LN-NLCs modified with 0.02% R11 exhibited favorable skin permeation and deposition in the skin by increasing the amount of LN penetrated through the skin and retained in skin layers, a condition that was favorable for topical application. Additionally, HPMC was used to prepare LN-NLC-R11 gels for the in vivo anti-inflammatory study. In the carrageenaninduced rat paw edema model, LN-NLC-R11 gels inhibited obviously rat paw edema and produced inflammatory cytokines compared with LN-NLC gels and LN gels. Thus, LN-NLC gels containing the cell-penetrating peptide R11 developed in our study enhanced the translocation of LN across the skin, thereby alleviating the severity of inflammation and making it a promising preferred formulation of LN.

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Disclosure

The authors report no conflicts of interest in this work.

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