Novel lipophilic SN38 prodrug forming stable liposomes for colorectal carcinoma therapy

Jing Xing1,2,*
Xiquan Zhang3,*
Zhe Wang4
Huanqing Zhang3
Peng Chen1,2
Gaoxin Zhou5
Chunlong Sun6
Ning Gu1,2
Min Ji1,2

1School of Biological Science and Medical Engineering, Southeast University, Nanjing 210096, People’s Republic of China; 2Department of Biological Science and Medical Engineering and Collaborative Innovation Center of Suzhou Nano Science and Technology, Suzhou University, Suzhou 215123, People’s Republic of China; 3Nanjing Institute of Pharmaceutical Research and Development, Chia-Tai Tianqing Pharmaceutical Group Co. Ltd, Nanjing 210023, People’s Republic of China; 4Emergency Department, The Second Affiliated Hospital of Nanjing Medical University, Nanjing 210029, People’s Republic of China; 5School of Biomedical Engineering, Shenzhen University, Shenzhen 518071, People’s Republic of China; 6College of Biological and Environmental Engineering, Binzhou University, Binzhou 256603, People’s Republic of China

*These authors contributed equally to this work

Background: SN38 (7-ethyl-10-hydroxy camptothecin), as a potent metabolite of irinotecan, is highly efficacious in cancer treatment. However, the clinical utility of SN38 has been greatly limited due to its undesirable properties, such as poor solubility and low stability. Therefore, the development of an acceptable SN38 prodrug is highly anticipated. In order to overcome these weaknesses, moeizitecan, a lipophilic SN38 prodrug containing a SN-38, a trolox, a succinic acid linker, and a hexadecanol chain, was loaded into liposomal nanoparticles by ethanol injection method.

Materials and methods: In order to overcome these weaknesses, moeizitecan, a lipophilic SN38 prodrug containing a SN-38, a trolox, a succinic acid linker, and a hexadecanol chain, was loaded into liposomal nanoparticles by ethanol injection method.

Results: Experiments showed that the moeizitecan-loaded liposomal nanoparticles (MLP) with a diameter of 105.10±1.49 nm have a satisfactory drug loading rate (90.54±0.41%), high solubility and stability, and showed sustained release of SN38. Notably, MLP exhibited better antitumor activity against human colon adenocarcinoma cells than irinotecan, a FDA-approved drug for the treatment of advanced colorectal cancer. Furthermore, xenograft model results showed that MLP outperformed irinotecan in terms of pharmacokinetics, in vivo therapeutic efficacy and safety. Finally, we used molecular dynamic simulations to explore the association between the structure of MLP and the physical and functional properties of MLP. Mooritecan molecules in MLP folded themselves inside the hydrocarbon chain of the lipid bilayer, which led an increased acyl chain order of the lipid bilayer, and therefore enhanced the lactone ring stability protecting it from hydrolysis.

Conclusion: Our MLP construction strategy by liposome engineering technology may serve as a promising universal approach for the effective and safe delivery of lipophilic prodrug.

Keywords: SN38, lipophilic prodrug, liposomes, molecular dynamic simulations, cancer therapy

Introduction

SN38 (7-ethyl-10-hydroxy camptothecin) is a potent antineoplastic agent. As a member of the camptothecin family, SN38 inhibits DNA topoisomerase I, prevents DNA religation and replication via irreversibly binding to topo-I and DNA, and ultimately causes apoptosis in tumor cells.1,2 The high potency of SN38 makes it a great candidate against many malignancies, such as colorectal carcinoma, lung tumors, gastric, cervical and ovarian cancers.3 However, the clinical application of SN38 is limited due to its poor solubility in most pharmaceutically acceptable solvents and side effects caused by uncontrolled conversion from active lactone ring to inactive carboxylate form at pH>6.4–6 Then, a water-soluble prodrug of SN38, irinotecan (CPT-11) was developed and has been widely used as a clinically first- or second-line drug to treat the advanced colorectal cancer.7–9 However, only 2–8% of CPT-11 administrated transforms into the active metabolite.10,11 Moreover, the active lactone ring of CPT-11 is readily converted...
to the inactive carboxylate form after infusion in humans. Thus, unpredictable therapeutic side effects and dose-limiting toxicity usually happen when treating patients with CPT-11.

To overcome these problems, numerous drug delivery systems, such as prodrugs, polymeric micelles, and liposome-based formulations, have been extensively developed. The macromolecular prodrugs and small-molecule prodrugs with amphiphilic characteristic can self-assemble into nanoscale particle or be wrapped with nanoscale vehicles. Although the prodrugs-assembled nanoparticles were effective in high drug loading, enhancing drug delivery to tumor tissue and suppressing tumor growth, their inherent shortcomings must be considered, especially inadequate pharmacokinetics. Employing liposomes as the drug vector of SN38 and its derivatives is an effective strategy to prolong the systemic circulation of their in the bloodstream and optimal concentration in tumor tissues. Liposomes are lipid bilayer vesicles. And as one of the best drug delivery systems, they were investigated for delivery of both lipophilic and hydrophilic drugs. Especially the nanoliposomes have some advantages, such as increased solubility and stability, enhanced cellular uptake, improved bioavailability, and specific targeting of organs or tissues for the controlled release of drugs. Due to the robust biocompatibility, several liposomal formulations are available for chemotherapeutic application in clinic. Encapsulation of drugs within liposomes often confers the drugs with high stability to avoid chemical degradation. However, it is a challenge to employ liposomes to deliver SN38 because of its unfavorable physicochemical properties and low drug entrapment efficiency. Increasing the lipophilicity through lipophilic conjugate of SN-38 has multiple benefits, including enhancement of drug loading in lipid-based formulations, possible stabilization of the active lactone form, and improvement of the permeability of the drug through cell membranes. Indeed, many research have focused on the biological activities of the liposomal formulations of lipophilic prodrug of SN38. A few studies concern on the effect of lipophilic prodrug on the membrane of liposome formulation. However, there have been no reports on the influence of encapsulation of the lipophilic prodrug of SN38 on the structure and the stability of the lipid bilayer.

Motivated by this rationale, we have recently developed a liposome-based delivery system for encapsulation of a lipophilic SN38 prodrug (designated as moeixitecan) to improve its stability and biological activities. In our previous study, we already synthesized moeixitecan, which contained a SN-38 component, a trolox component, a succinic acid linker, and a hexadecanol chain, with excellent properties in vivo and in the clinic. The physiochemical properties of moeixitecan-loaded nanoliposomes were characterized and the antitumor efficacy were evaluated in colorectal tumor models in vitro and in vivo, respectively. Meanwhile, molecular dynamics (MD) simulations were performed to expound the underlying molecular mechanisms of the influence of moeixitecan molecules on the structure and the stability of the phospholipid bilayer. Finally, the in vivo biodistribution of moeixitecan-loaded nanoliposomes and a safety evaluation compared to CPT11 were examined in mouse.

Materials and methods
Materials
2-(hexadecyloxy carbonyl) -2, 5, 7, 8-tetramethylchrooman-6-yl 7-ethyl-camptothecin-10-y1 succinate (moeixitecan) obtained internally by Chia-tai Tianqing Pharmaceutical (Nanjing, People's Republic of China). 7-Ethyl-10-hydroxycamptothecin (SN38) was provided by Sigma-Aldrich (Shanghai). Dipalmitoyl phosphatidylcholine (DPPC), hydrogenated soybean phospholipids (HSPC), and distearoyl phosphoethanolamine-PEG2000 (DSPE-PEG2000) were purchased from Southeast Pharmaceuticals Co. (Soochow, People's Republic of China). Chloroform and methanol were purchased from Shanghai Chemical Reagent Company.

Preparation of moeixitecan (SN38 prodrug)-loaded liposomal nanoparticles (MLP)
Liposomes loaded with moeixitecan were prepared on the basis of ethanol injection as described previously. Briefly, DPPC, HSPC, and DSPE-PEG2000 were first dissolved in ethanol at a molar ratio of 75:20:5. The moeixitecan was added to the above ethanolic solution with the ratio of lipids to moeixitecan fixed at 20:1 (w/w) and incubated at 60°C. The mixture (1 mL) was then rapidly injected into 5% dextrose solution (9 mL) at 60°C. Finally, the resulting vesicles suspension was extruded through polycarbonate filters (Avanti Polar Lipids, Alabaster, AL, USA) of 200 and 100 nm (five times each) for MLP.

Characterization of MLP
The particle size and surface potential of MLP were measured at 25±0.1°C by Nano-ZS 90 Nanosizer (Malvern, UK). The morphology and structure were observed by transmission electron microscope (JEOL, Japan) with negative stain method. Before analysis, the samples were prepared by
placing a drip onto 300-mesh copper-grid, and a drip of 3 wt% phosphotungstic acid was dropped on the samples, then kept the sample at room temperature overnight for drying.

The encapsulation efficiencies (EE) of moeixitecan in liposomes were determined. Firstly, the nanoparticles were extracting 1 mL of MLP with 10 mL acetonitrile for 1 hr while stirring. Then, the samples were then filtered and analyzed by HPLC. A C18 ODS reverse-phase column (5 mm, 250 mm×4.6 mm, Shimadzu, Japan) was used for HPLC analysis at room temperature. The mobile phase consisted of methanol and 1% formic acid aqueous solution (80:20, v/v) and the flow rate was maintained at 1 mL/min. The injection volume was 20 μL and the determined wavelength was 220 nm. The EE of moeixitecan were calculated according to the following formula EE (%)=(Moeixitecan\text{total}−Moeixitecan\text{free})/Moeixitecan\text{total}×100%.

**In vitro drug release from liposomes**

The drug release behavior from free moeixitecan and MLP was evaluated using the dialysis method against PBS (pH 7.4) containing 0.2% Tween 80. Free moeixitecan solutions and MLP solutions were infused into dialysis bags (1 mL solution per bag at an SN38 equivalent concentration of 0.5 mg/mL) and submerged into 50 mL of PBS containing 0.2% Tween 80 (w/v) and maintained at 37°C under 100 rpm shaking for 96 hrs. At pre-determined time intervals, 1 mL of the release media was collected and equal volumes of fresh media were supplemented. After freeze-drying and redissolution with 1.5 mL methanol, the concentration of moeixitecan and SN38 was analyzed by HPLC. Finally, the cumulative amount of drug released into the media at each time point was calculated as the percentage of total released drug to the initial amount of the drug.

**Physical stability of MLP**

The physical stability of MLP was estimated by measuring the size change of liposomes in the rat plasma as well as PBS buffer (pH 7.4) with or without 20% FBS at 37°C. Lastly, the MLP prepared with the highest physical stability was further evaluated for any size change during storage at 4°C for 3 months.

**MD simulations of the effect of moeixitecan on the structure of lipid membranes**

We performed atomistic MD simulations for six model systems (Table 1) containing lipid bilayers with 0–25 mol% of moeixitecan. Each system was added with 4000 water molecules and Na\textsuperscript{+} or Cl\textsuperscript{−} ions were added in order to achieve physiological concentration.

Simulations were carried out using the GROMACS 4.6 software package. The all-atom OPLS force field was employed with a recent extension for lipids to parameterize all lipid molecules. The TIP3P model, compatible with the OPLS-AA force field, was used for water. The time step was set to 2 fs, and the simulations were carried out at 1 bar and 300 K. The Nosé-Hoover method was used to couple the temperature with separate heat baths for the membrane and the rest of the system with time constants of 0.4 ps. The linear constraint solver algorithm was used to minimize the energy of the initial configurations.

**Cell lines and cell culture**

Human colon adenocarcinoma cells (HT-29) were used in this study provided by Cell Bank of Shanghai Institute of Cell Biology, Chinese Academy of Sciences. HT29 cells were maintained in McCoy’s 5A (modified) medium, supplemented with 10% FBS and 1% penicillin/streptomycin at 37°C.

**In vitro cytotoxicity**

The cytotoxicity of MLP was assessed against HT-29 cells by the CCK-8 assays. Briefly, HT29 cells were seeded in

| System | Dipalmitoyl phosphatidylcholine | Moeixitecan | Water | Simulation length (ns) |
|--------|---------------------------------|-------------|-------|-----------------------|
| 1 (0 mol%) | 200 | 0 | 4000 | 2×500 |
| 2 (5 mol%) | 190 | 10 | 4000 | 2×500 |
| 3 (10 mol%) | 180 | 20 | 4000 | 2×500 |
| 4 (15 mol%) | 170 | 30 | 4000 | 2×500 |
| 5 (20 mol%) | 160 | 40 | 4000 | 2×500 |
| 6 (25 mol%) | 150 | 50 | 4000 | 2×500 |
96-well plates at a density of 5000 cells per well and incubated for 24 hrs. Then, MLP at various concentrations was added to the culture media. Moeixitecan and CPT-11 were used as controls. After incubation for 24 and 48 hrs, the cell viability was determined using CCK8 assays. At the end of the incubation period, 10 μL of CCK-8 reagent (Beyotime, Shanghai, People’s Republic of China) was added to the cells and further incubated for 2 hrs. Afterward, the 96-well plate was analyzed at a 450 nm wavelength in an Infinite 200 PRO plate reader (Tecan, Switzerland) to determine cell viability.

**Apoptosis analysis using flow cytometry**

The apoptosis of HT29 cells was measured by Annexin-V-FITC/PI double staining method. HT29 cells were seeded in 6-well plates at a density of 2×10^5 cells per well and incubated overnight. Then, moeixitecan, CPT-11, and MLP (with the equal SN38 dose of 2 μg/mL) were added to the culture media. Cells that were untreated were used as controls. After 24 or 48 hrs of treatment, the HT29 cells were collected and detected on FACS CantoII flow cytometry (BD, MA) by using an Annexin-V-FITC Apoptosis Detection Kit (Beyotime).

**Animals and tumor model**

The animal experiments were carried out according to the protocol approved by the Ministry of Health in People’s Republic of China (document no. 55, 2001) and the guidelines for the care and use of Laboratory Animals of Southeast University. Female BALB/C nude mice (20±2 g, 6–8 weeks old) were purchased from Nanjing model animal research institute (Nanjing, People’s Republic of China) and received care under standard housing conditions. HT-29 cells (6×10^5/200 μL PBS) were administrated by subcutaneous injection into the flank region of the mice. Two weeks after post-inoculation, the xenograft tumor model was built. Tumor volume was calculated as (tumor length)×(tumor width)^2/2.

**In vivo pharmacokinetic studies**

In vivo pharmacokinetic study was performed according to the previous report. Briefly, the Sprague-Dawley rats (250 ±20 g) were injected with a single dose of 2.5 mg/kg (SN38 equivalent) of CPT11, free moeixitecan, MLP via tail vein (n=3), respectively. Blood samples were collected into heparinized tubes after 0.5, 1, 3, 6, 12, and 24 hrs. Subsequently, whole blood samples were centrifuged at 3000 rpm for 10 mins at 4°C to separate the plasma. The plasma was stored at 80°C prior to analysis by LC/MS/MS. An aliquot 5 μL of the processed samples were injected to LC/MS/MS system and the sample was separated on an Agilent Eclipse Plus C18 (4.6×100 mm, 3.5 μm, Agilent Technologies, USA), which was kept at room temperature (25±2°C). In the current study, SN38 were separated on a gradient elution, which was performed by varying the proportion of mixture A (5 mM ammonium acetate with 0.2% formic acid) and mixture B (acetonitrile) as follows: 0 min (75% A–25% B), at 1.0 min (50% A–50% B), at 1.5 mins (75% A–25% B), at 1.6 mins (25% A–75% B), and at 2.2 mins (25% A–75% B) was used and delivered at a rate of 0.60 mL min^-1 into the mass spectrometer’s electrospray ionization chamber. SN38 were detected by triple-quadrupole LC/tandem mass spectrometric detection (A Shimadzu-LC-30AD from Shimadzu, Tokyo, Japan equipped with MS/MS; AB Sciex triple quad 6500, from Sciex, Framingham, MA) with electrospray ionization interface running in positive ionization mode. After extracting, samples were analyzed using a LC/MS/MS analysis.

**In vivo distribution by near-infrared (NIR) imaging**

A NIR region probe (iDSPE)-embeded liposomes were used to evaluate the in vivo distribution of MLP in HT29 tumor-bearing mice. The NIR probe, iDSPE was co-encapsulated with moeixitecan in iDSPE-embeded MLP using the same method. The tumor-bearing mice were intravenously injected 0.2 mL of ir623 and iDSPE-embeded MLP, respectively. The equivalent ir623 dose was kept at 0.5 mg/kg. The NIR images of mice were obtained at 0.5, 2, 4, 8, 24, and 48 hrs postinjection using the ex/in vivo imaging system (Caliper Life Science, USA) with a 630 nm excitation wavelength and 710 nm filter. After imaging, the mice were immediately sacrificed and organs were harvested for the ex vivo imaging under the same conditions as described above.

**In vivo antitumor efficiency**

The antitumor activity was conducted using HT29 tumor-bearing mice. When the tumor volume reached about 100 mm³, the mice were randomly divided into four groups (n=5 per group) and treated with PBS, CPT11, free moeixitecan, MLP (with the equal SN38 dose of 2.5 mg/kg). Drugs were administered intravenously via the tail vein at designated times. The tumor sizes were measured with a...
caliper, and the tumor volumes were calculated according to the formula (length×width²)×1/2. The body weight of mice in each group was also monitored. At the end of the “antitumor activity study”, the mice were sacrificed for humane reasons. Meanwhile, the tumors and major organs were excised and fixed with 4% formalin. Then, the organs sections were mounted on glass slides and stained with H&E and observed using a digital microscope.

Safety evaluation
The blood collected from the HT29 tumor-bearing mice were used to perform hematological tests using a blood counter (Sinnowa, HBVet-5 model) with adapted dilutions. Hematological analysis involves assessment of white blood cell count, red blood cell (RBC) count, hematocrit, % neutrophils, % lymphocyte, % monocyte, % eosinophil, and % basophile.

Results and discussion
Preparation and characterization of MLP
The structure of moeixitecan is 2-(hexadecyloxycarbonyl)-2,5,7,8-tetramethylchroman-6-yl 7-ethyl-camptothecin-10-yl succinate, containing an SN-38 component, a trolox component, a succinic acid linker, and a hexadecanol chain, as shown in Figure 1A. MLP were prepared by ethanol injection method, the lipids were composed of DPPC, HSPC, and DSPE-PEG2000, as depicted in Figure 1A. Then, the physiochemical characters of MLP were systematically evaluated and the results are presented in Table 2, including mean diameter, polydispersity index (PDI) value, zeta potentials, and EE. The average hydrodynamic diameter of MLP was 105.10±1.49 nm, which was a little lower than the blank liposomes (117±2.49 nm), as shown in Figure 1C. Compared with the blank liposomes, MLP had a much lower surface charge, indicating a good stability of the formulations against vesicle aggregation and fusion. As expected, when moeixitecan was added, the negative zeta potential increased significantly, which would maintain the stability of the liposome solution by electrostatic repulsion. In addition, the EE of MLP was 90.54±0.41%, which was higher than that (34.62–60.11%) of CPT11-loaded nanoliposomes reported in the literature.

The morphology of the MLP was observed by transmission electron microscopy. As shown in Figure 1B, the image obviously verified the formation of liposomes with uniform spherical nanostructures. With the addition of moeixitecan, MLP became more stable and maintained a consistent size distribution without significant variation during storage at 4°C for 3 months compared with the blank liposomes (Figure 1D). In order to provide further evidence for in vivo applications, the particle stability in plasma of the liposomes for 7 days was evaluated. MLP showed no significant change in sizes within rat plasma (~150 nm) in 7 days (Figure 1E). In contrast, without moeixitecan, the mixture of lipid could not form stable bilayer liposomes, and formed large aggregates in rat plasma (data were not shown). All of these results proved that moeixitecan entrapped in liposomes formed a stable nanoparticle, and MLP exhibited great size stability either in storage condition (4°C) or physiological environment (in plasma).

In vitro drug release
The in vitro release properties of MLP were investigated in PBS containing 0.2% Tween 80 via dialysis followed by HPLC. Figure 2 presents in vitro release of free SN38 from liposomal formulation and its solution as control. As shown in Figure 2, MLP exhibited a slower and sustained release behavior in comparison with its solution. Cumulative release of SN38 after 72 hrs was 63.55±6.81% from the MLP, whereas 80.57±5.37% for moeixitecan solution within 48 hrs. There was no burst release of SN38 detected from the liposomes, which strongly suggested that moeixitecan was located in lipid bilayers, and thereby enhance drug retention within the liposomes but SN38 can be slowly released. The sustained release of SN38 could be beneficial for cancer treatment as a result of the maintenance of constant SN38 concentration in tumor tissue.36

MD simulations of moeixitecan-lipid membranes interaction
To clarify the interaction between moeixitecan and simplified lipid bilayer in more detail, we carried out atomistic MD simulations. We performed all-atom MD simulations of a DPPC bilayer before and after adding 5 mol%, 10 mol%, 15 mol%, 20 mol%, and 25 mol% moeixitecan. The MD simulations can provide a detailed description of the changes in bilayer structure incurred by the incorporation of moeixitecan into a lipid bilayer. Figure 3A and B illustrates the location (depth) and orientation of moeixitecan molecules inside membranes at the end of the simulations. The moeixitecan molecular may integrate itself into the hydrocarbon chain region of the lipid bilayer. To estimate the position of moeixitecan with respect to the lipid molecules, we calculated their mass density profiles across the bilayer. Figure 3C...
shows the density profiles along the membrane normal for the moeixitecan. At the end of the modeling, the distance between the moeixitecan molecule and the center of membrane was 4 nm. Therefore, these figures show that moeixitecan folding is preferably located inside the fatty acid chains of the phospholipid. The thickness of the membrane increased with increasing concentration of moeixitecan (Figure 3D). Meanwhile, the areas per molecule of lipid

**Figure 1** The characterization and physical stability of moeixitecan-loaded liposomal nanoparticle (MLP). (A) Schematic illustration of the SN38 prodrug, MLP. (B) Morphological characteristics of MLP. (C) Particle size distribution of MLP. (D) Size changes of MLP were examined as a function of time at 4°C for 3 months. (E) Size changes of MLP in different medium were examined for 7 days.

| MLP | Composition with the molar ratio | Size (nm) (PDI) | Zeta potential (mV) | EE (%) |
|-----|---------------------------------|-----------------|---------------------|--------|
| DPPC:HSPC:PEG-DSPE=75:20:5 | 105.31±0.43 (0.152±0.02) | 20.31±6.24 | 90.54±0.41 |

**Abbreviations:** DPPC, dipalmitoyl phosphatidylcholine; HSPC, 1-palmitoyl-2-stearoyl-sn-glycero-3-phosphocholine; PEG-DSPE, 1,2-distearoyl-sn-glycero-3-phosphoethanolamine-N-[methoxy(polyethylene glycol)-2000]; PDI: polydispersity index.
bilayer decreased with increasing concentration of moxizitecan (Figure 3E). Lipid membrane thickness and the area per molecule are structural parameters that are needed to accurately determine other bilayer structural parameters and are directly related to lipid–lipid, and lipid–protein interactions in biomembranes. Our MD simulations results confirmed the experimental results and provided an accurate picture of moxizitecan’s location and effect for lipid bilayers: moxizitecan molecules fold themselves located inside the hydrocarbon chain of the lipid bilayer inducing an increased acyl chain order of the lipid bilayers and lactone ring stability. Both physical stability of MLP and MD simulations data consistent with each other and showed that moxizitecan has an ordering effect, and increases the stability of bilayer.

In vitro cytotoxicity
The in vitro cytotoxicity was evaluated by CCK8 assay on human colon adenocarcinoma cells (HT-29). The MLP, moxizitecan, and CPT-11 were incubated with HT29 cells for 24 and 48 hrs to compare their cytotoxicity in vitro (Figure 4). The IC₅₀ (concentration inhibiting cell growth to 50% of control) values of CPT-11, MLP, and moxizitecan at 48 hrs were 79,024, 0.8461, and 3.139 μg/mL, respectively. More specifically, the MLP achieved an outstanding cytotoxic effect compared to the clinical anticancer drug CPT-11. Two possible reasons were thought to induce the increased cytotoxicity of moxizitecan and MLP. First, the combination of moxizitecan and phospholipids was more effective for cellular uptake, which could contribute to the enhanced cytotoxicity. Second, once be uptaken by cancer cells, the moxizitecan can sufficiently convert to active SN38 molecules by hydrolysis, and reached a relatively high drug concentration. Hence, MLP exhibited potent antitumor activity comparable or even slightly superior in comparison
with moeixitecan, both of which were significantly better than CPT-11 in vitro.

**In vitro cell apoptosis induced by released SN38 molecules**
The apoptosis assay was used to elucidate the death mechanism of HT-29 cells incubated with various drug formulations. After treatment for 24 and 48 hrs, the cells were measured using Annexin-V-FITC/PI double staining analyzed by flow cytometry. The motion of phosphatidylserine to the exterior of the cell membrane is an essential event in apoptosis cells which can be specifically detected by fluorescently labeled Annexin V. Indeed, a high level of early and late apoptosis after 24 or 48 hrs of treatment was induced by MLP and moeixitecan, which was comparable to the level induced by CPT-11. After 24 hrs, the percentage of apoptotic HT-29 cells treated with CPT-11, moeixitecan, and MLP were 16.6%, 34%, and 31.4%, respectively (Figure 4C). However, after 48 hrs of incubation, the apoptosis proportion of HT-29 cells incubated with CPT-11, moeixitecan, and MLP increased to 29.4%, 41.6%, and 46.4%, respectively (Figure 4C). Hence, increased cellular apoptosis was observed because of extending treatment time. In summary, the cell apoptosis against cancer cells was consistent with the results of in vitro cytotoxicity assays, suggesting that moeixitecan-loaded NPs can perform better anti-proliferative potency than CPT-11.

**In vivo pharmacokinetic studies**
To evaluate the MLP in the blood circulation system, the pharmacokinetic study was assessed in Sprague-Dawley rats after intravenous injection of the CPT-11, moeixitecan, and MLP solutions (at the equivalent SN38 dose of 2.5 mg/kg).
The concentration-time profiles of CPT-11, moeixitecan, and MLP are shown in Figure 5 and the main pharmacokinetic parameters are provided in Table 3. The maximum SN38 concentrations (C_max) for MLP were evidently higher when moeixitecan was encapsulated in the liposomal formulations. The C_max of the rats treated with CPT-11, moeixitecan, and MLP were 56, 137, and 616 ng/mL, respectively. This was further reflected in differences in t_{1/2} and the values of mean residence time (MRT): t_{1/2} with CPT-11 at 1.73 hrs, moeixitecan at 2.42 hrs, and MLP at 5.23 hrs; MRT with CPT-11 at 2.44 hrs, moeixitecan at 4.64 hrs, and MLP at 6.70 hrs. Moreover, the AUC of SN38 is higher upon administration of both moeixitecan (774 ng/L h) and MLP (3454 ng/L h) compared to CPT-11 (118 ng/L h). In particular, the AUC value of MLP was 30 times higher than that of CPT-11. These results indicate that the MLP exhibited an increase in net drug exposure and systemic circulation time in the blood.

Very recently, we have identified a novel NIR dye, iDSPE, which are able to insert into the lipid bilayers of liposomes to evaluate the blood retention and distribution of liposomes conveniently.34 To evaluate the real-time biodistribution and tumor targeting efficiency of liposomes in HT-29 tumor-bearing mice, various formulations were injected into tumor-bearing mice via tail intravenous injection. The results for real-time imaging and biodistribution of ir623 and the iDSPE-embeded MLP (iMLP) were observed in HT-29 tumor-bearing mice by ex/in vivo imaging system (Figure 6). Obviously, after intravenous injection of free ir623, nonspecific distribution of fluorescence all over the body was observed within 2 hrs, and with rapidly decreased fluorescence signals within 48 hrs injection. While iMLP group exhibited detectable NIR fluorescence signals at 24 hrs postinjection, which were mainly concentrated at the tumor sites and retained long term in the tumor (Figure 6A). After living imaged, the mice were sacrificed and tumor tissue as well as other main organs were harvested for the ex vivo fluorescence imaging, also revealing the obvious tumor accumulation of the iMLP (Figure 6B). These results indicated that the MLP showed an excellent tumor-targeting capability and a potential to help extend drug retention in blood correlate well with the pharmacokinetic studies. Therefore, nanoliposomes may be an ideal delivery platform for the passive accumulation of drugs in tumor lesions.

![Figure 5](image) In vivo plasma concentration-time profiles of SN38 after intravenous injection of CPT-11, moeixitecan (M), and moeixitecan-loaded liposomal nanoparticles (MLP) in SD rats (n=3).

In vivo antitumor activity in human colon xenograft tumors

The effectiveness of MLP in suppressing of tumor growth was evaluated in a HT-29 colorectal xenograft model. The group with saline solution and that received CPT-11 served as the control. As shown in Figure 7A, the saline group showed a rapid increase in tumor growth in HT-29 tumor-bearing mice over the whole period of the experiment with tumor volume reaching approximately 1025±139 mm³ at day 15. Meanwhile, the tumor growth was remarkably inhibited after treated with CPT-11, moeixitecan, and MLP as compared to saline solution. When the treatment was finished, the mice were sacrificed and the tumors were

![Table 3](image) The main pharmacokinetic parameters in SD rats after intravenous injection of CPT-11, moeixitecan (M), and moeixitecan-loaded liposomal nanoparticles (MLP) (n=3)

| Parameters                  | CPT-11        | M            | MLP           |
|-----------------------------|---------------|--------------|---------------|
| Dose (mg/kg)                | 5             | 5            | 5             |
| AUC_{0-∞}(h×ng/mL)          | 118±8         | 774±129      | 3454±352      |
| C_max (ng/mL)               | 56.1±10.7     | 137±8.0      | 616±43.4      |
| t_max                        | 0.083±0.000   | 0.229±0.197  | 0.265±0.190   |
| T_{1/2}(h)                  | 1.730±0.106   | 2.420±0.195  | 4.641±0.103   |
| MRT (h)                     | 2.443±0.089   | 4.641±0.103  | 6.700±1.470   |

Note: Data are presented as mean±standard deviation.
Figure 6 Biodistribution of ir623 and iMLP in HT-29 tumor-bearing mice. (A) Representative images of in vivo whole-body imaging of mice at 0.5, 4, 8, 24, and 48 hrs postintravenous injection of ir623 and iMLP. (B) The ex vivo optical images of the tumors and other major organs from the sacrificed mice (n=3).
Abbreviation: iMLP, iDSPE-embedded moeixitecan-loaded liposomal nanoparticles.

Figure 7 Anti-tumor effect on HT-29 tumor-bearing mice treated with moeixitecan-loaded liposomal nanoparticles (MLP). (A) Tumor volume changes as a function of time after intravenous injection of CPT-11, moeixitecan (M), and MLP. (B) Body weight profile of tumor-bearing mice after treatment. (C) Typical excised tumors at the time of sacrifice. (D) Statistical results for tumor weight.
weighed. The tumor weights of mice were consistent with the tumor volume results: the overall inhibition rates of CPT-11, moeixitecan, and MLP were 46.06%±6.30%, 54.41%±7.30%, and 66.86%±6.54%, respectively (p<0.01, Figure 7C and D). Treated with liposomally formulated moeixitecan with a low dose of SN38 (2.5 mg/kg) had an enhanced effect, resulting in 66.86% tumor inhibition compared with CPT-11 treatment (46.06%, p<0.05). Moreover, MLP were even more effective than our previously developed prodrug moeixitecan that only led to 54.41% in the inhibition rates (p<0.05). The increased in vivo antitumor activity of MLP is likely due to the sustained drug release kinetics, improved systemic circulation, and preferential tumor accumulation via the enhanced permeability and retention (EPR) effect. The results are consistent with in vitro cytotoxicity indicating that MLP exerted excellent therapeutic activity better than CPT-11 on the xenograft tumors.

Figure 8 Safety evaluation of moeixitecan-loaded liposomal nanoparticles (MLP).
Notes: (A) H&E stained images of organ sections separated from mice. Magnification, ×200 (B) blood parameters panel data from mice injected i.v. with saline (control, CT), CPT-11, moeixitecan (M), and moeixitecan-loaded liposomal nanoparticles (MLP). Mean parameters value±SD, n=5. RBC count (a), WBC count (b), neutrophil count (c), lymphocyte count (d), monocyte count (e), and HCT value (f).
Abbreviations: WBC, white blood cell; RBC, red blood cell; HCT, hematocrit.
Safety evaluation
The variations in the body weights of mice in all the treatment group were plotted against time as shown in Figure 7B. Obviously, the body weights of mice did not decrease significantly compared with control group. Afterward, we evaluated the toxicity of different formulations by histological studies on excised tissue sections with H&E staining. As shown in Figure 8A, we did not observe any appreciable abnormality in major organs. Finally, we further evaluated hematological affects by routine peripheral blood examination. The blood cell counts were assessed to investigate the hematological toxicity against MLP. There was no significant variation in the counts of RBC, hemoglobin, or in any other routine peripheral blood tests of CPT-11 treatment groups and MLP treatment group when compared with the control group (Figure 8B). However, the blood cells counts were significantly increased in the moeixitecan treatment groups compared to the control group and the MLP treatment group. The reductive toxicity for MLP might be attributed to liposomal incorporation can encapsulate drug for providing more selective delivery or targeting to the tumor tissue by reduced accumulation in the normal tissues. Thus, these results indicated the MLP, at the test dose for in vivo administration, was relatively safe. Although animal experiments with higher doses of MLP were not explored, raised doses should be expected to improve antitumor activity.

Conclusion
In summary, we successfully designed stable liposomal formulations containing a lipophilic prodrug of SN38, moeixitecan. The MLP not only overcome various drawbacks of CPT-11 and SN38, but also resolve various limitations of the reported SN38-incorporating drug delivery systems, especially the premature drug release. The moeixitecan was able to incorporate into liposomes with high and stable drug loading, significantly improved solubility, stability, and sustained drug release kinetics. Furthermore, the moeixitecan-loaded liposomes exhibited significant cytotoxicity with selective promotion of cell apoptosis against HT-29 cell lines compared with CPT-11, and showed more therapeutic effect in a colorectal xenograft model than the clinically approved CPT-11 and moeixitecan. Simultaneously, the in vivo pharmacokinetics of the MLP were improved relative to moeixitecan and the toxicities were lower to normal tissues in tumorbearing mice. Particularly, the results of MD simulations indicated that folding moeixitecan molecules were preferably accumulated in the hydrophobic acyl chain region of the lipid bilayers, which caused packing and ordering of lipids increasing the stability of lipid bilayers and protect stabilization of the lactone form from hydrolysis. Both experimental are in good agreement with the theoretical data. The MD simulations provide an insight into the nature of moeixitecan-lipid membrane interactions at the molecular level and could be used for optimizing the liposomal formulations. Therefore, our results suggest that the moeixitecan-loaded liposomes could be a promising chemotherapeutic drug better than CPT-11 for clinical applications.

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Disclosure
The authors report no conflicts of interest in this work.

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