Research Article

Polymer-Stabilized Micelles Reduce the Drug Rapid Clearance In Vivo

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Micelles are self-assembled nanoscaled aggregates from amphiphilic unimers and can be used to encapsulate hydrophobic drugs. However, the dynamic exchanging of unimers between micelles and bulk solution often leads to micelle destabilization and subsequent leaking of the encapsulated substances. Thus, we incorporated a hydrophobic polymer into the micellar core for interlacing the unimers and stabilizing the micelle structure. The polymer-stabilized and non-polymer-stabilized micelles have similar physicochemical properties including small sizes (~35 nm), negative surface charges (~−35 mV), and high drug contents (~15%). Drugs encapsulated in polymer-stabilized micelles are released in a slower rate than are non-polymer-stabilized micelles. From in vivo pharmacokinetic studies, drugs loaded in polymer-stabilized micelles have lower clearance and higher plasma concentration and lower volume distribution than non-polymer-stabilized micelles have. In conclusion, polymer-stabilized micelles can reduce rapid drug clearance via strengthening of the micellar structure and increase in the available drug amount in plasma, thus broadening pharmaceutical applications of micelles.

1. Introduction

Nanotechnology is widely employed in drug delivery systems (DDSs) [1, 2]. In order to improve drug availability and to overcome the barriers of the biological system, the DDSs are generally designed to have sizes falling within the range of 1–100 nm [3]. In addition, other preferred properties of the DDSs would include biodegradability, biocompatibility, high loading capacity, good stability, high content of the drug in a final preparation, prolonged circulation, and ability to accumulate in required areas [4–6]. Common types of DDS include liposomes [7, 8], micelles [9–11], and polymeric nanoparticles [12–14].

PEG-PE micelles prepared from conjugates of polyethylene glycol (PEG) and phosphatidylethanolamine (PE) have a core-shell structure which can encapsulate hydrophobic drugs such as paclitaxel, parthenolide, tamoxifen, camptothecin, and budesonide [11]. Steric PEG block chains on the surface of micelles can decrease the opsonin protein binding in vivo and therefore block the capture by the reticuloendothelial system (RES) efficiently [15]. The low critical micellization concentration (CMC) value of PEG-PE compounds (in a 10−5 to 10−6 M range) indicates that PEG-PE micelles may maintain their integrity even upon strong dilution (e.g., by the large blood volume upon intravenous administration) [9]. PEG-PE micelle sizes can be adjusted from 7 nm to 40 nm by altering PEG with different lengths [16]. These aforementioned small size and PEGylated surface make PEG-PE micelles able to accumulate in subcutaneous tumour [17] or the infract zone [18]. The PEG-PE micelles can be further modified with a functional group or antibody moiety on the surface to enhance their targeting efficiency [19]. PEG-PE micelles possess advantages such as small size, high solubility for hydrophobic drugs, low CMC, sustained release of drugs, targeting delivery, and potential for sterilization by filtration [20] for being a promising drug carrier [11].

However, this type of micelles is dynamic and in equilibrium with individual unimers that constantly undergo
exchange between the bulk and the micelles [21, 22]. The
entrapped drugs could leak out while micelles repeatedly
disintegrate and reassemble [23]. Such drug leaking issue
will limit its pharmaceutical applications. Therefore, we
designed a polymer-stabilized PEG-PE micelle which can
maintain the micelle integrity and reduce the drug leakage.
PLGA (poly(lactic-co-glycolic acid)), a hydrophobic biode-
gradable polymer, was selected to be incorporated into the
inner core of the micelles to interlace the carbon chains of
PEG-PE unimers and was presumed to decrease unimer
disassembly which may stabilize the PEG-PE micelles. A
schematic diagram is illustrated in Figure 1. The polymer-
stabilized PEG-PE micelles should have better stability than
non-polymer-stabilized PEG-PE micelles, and they can hold
entrapped drug molecules inside for longer periods. Since
usually hydrophobic drug release from micelles will be
cleared rapidly, such design can provide more available and
accessible drugs in vivo.

Clarithromycin (CL) is a clinically used macrolide anti-
biotic and is selected as a model drug to be loaded into
the PLGA-incorporated PEG-PE micelles (referred to as
P-MCL) and the non-PLGA-incorporated PEG-PE micelles
(referred to as MCL). All the physicochemical characteriza-
tion, release profile in vitro, and pharmacokinetics in vivo
of these two types of micelles were performed and ana-
ysed to examine whether polymer-incorporated micelles
could act as a better drug carrier than non-polymer-
incorporated micelles.

2. Materials and Methods

2.1. Materials. Clarithromycin (CL), roxithromycin (ROX),
poly(D,L-lactic-co-glycolicacid) (PLGA, lactide:glycolide = 50:50, MW 40 kD–75 kD), ammonium acetate, potas-
sium dihydrogen phosphate, phosphoric acid, phosphate-
buffered saline (PBS), xylazine, and formic acid were pur-
chased from Sigma-Aldrich (Saint Louis, MO, USA) and
used as supplied. 1,2-Distearoyl-sn-glycero-3-phosphoetha-
nolamine-N-[methoxy(polyethylene glycol)-2000] (PEG-
PE) was purchased from Avanti Polar Lipids (Alabaster,
AL, USA). Fetal bovine serum (FBS) was purchased from
Thermo Fisher Scientific (Waltham, MA, USA). Zoletil was
purchased from Virbac (Carros, France). Oasis WCX μElu-
tion SPE plates were products of Waters (Milford, MA,
USA). Acetonitrile, chloroform, and methanol were HPLC-
grade products of Honeywell International (Morristown,
NJ, USA).

2.2. Preparation of P-MCL and MCL. P-MCL was prepared
by an oil-in-water emulsion solvent evaporation method
similar to that reported previously [24]. Briefly, 7.5 mg of
CL, 10 mg of PLGA, and 20 mg of PEG-PE were dissolved
in 1 mL of chloroform and was added into 3 mL of H2O with
moderate stirring. The mixed solution was emulsi-
fied by son-
cication over an ice bath using a probe sonicator (XL-2000,
Misonix, NY, USA) at 8 W output for 60 seconds. Chloro-
form was then evaporated from the colloidal suspension by
a rotary evaporator. The suspension was centrifuged at
14000×g for 30 minutes to discard the undesired big aggre-
gates. The supernatant was collected and washed three times
using an Amicon Ultra-4 centrifugal filter (30 kD MWCO,
Merck Millipore, Billerica, MA, USA). For the preparation
of MCL, the procedures were the same as in P-MCL, except
that no PLGA was mixed in the first step.

2.3. Optimization of PLGA Incorporation. To determine the
optimal PLGA incorporation amount for stabilizing micelles,
P-MCLs with di
fferent feeding ratios of PLGA/PEG-PE
(0.25, 0.5, and 1) were prepared. These P-MCLs were then
incubated in 10% FBS with mild shaking at 37°C for the
stability test. At 0, 17, 24, 41, and 65 hours, the size of each
P-MCL was measured by DLS (dynamic light scattering)
analysis using 90Plus Particle Size Analyzer (Brookhaven
Instruments Corp., Holtsville, NY, USA).

2.4. Transmission Electron Microscopy (TEM) Imaging. TEM
was performed on a Japan Hitachi H-7650 instrument with
an acceleration voltage of 80–120 kV. TEM samples were
prepared by administering the micelle suspension onto a
copper grid followed by negative staining for 10 seconds at
room temperature with freshly prepared and sterile-filtered
2% (w/v) uranyl acetate aqueous solution.
2.5. Zeta Potential Measurements. Zeta potential was performed by a ZetaPALS analyser (Brookhaven Instruments Corp., Holtsville, NY, USA) with a 35 mW diode laser (660 nm in wavelength) and a detector with a scattering angle of 90 degrees. The measured results were analysed by a PALS zeta potential analyser (Brookhaven Instruments Corp., Holtsville, NY, USA).

2.6. Determination of Encapsulation Efficiency and Loading Content of CL. CL encapsulation efficiency (EE) was determined by applying (1), where \( M_{\text{encapsulated}} \) is the mass of CL entrapped within the micelles and \( M_{\text{add}} \) is the mass of the CL added in the preparation of micelles.

\[
\text{EE (\%)} = \frac{M_{\text{encapsulated}}}{M_{\text{add}}} \times 100\% .
\]

The CL loading content is defined as the mass fraction of CL in the micelles. The lyophilized micelles were weighted and then dissolved in 50% acetonitrile solution, and CL was quantified by an Agilent (Santa Clara, CA, USA) high-performance liquid chromatograph (HPLC) equipped with a C_{18} column (250 mm × 4.6 mm id, 5 \( \mu \)m) at an absorption wavelength of 205 nm. The mobile phase was composed of 50 mM potassium dihydrogen phosphate (pH 7.0): acetonitrile \( = 60 : 40 \) (v/v) with a flow rate of 1.0 mL/min.

2.7. CL In Vitro Release Profiles. 1 mg of micelles suspended in 1 mL of phosphate-buffered saline (PBS) or 10% fetal bovine serum (FBS) solution was placed into a dialysis tube (20 kD MWCO, Spectrum Laboratories, Rancho Dominguez, CA, USA). The dialysis tube was placed in 6 mL of PBS or 10% FBS and gently stirred with a magnetic stirrer in an incubator at 37°C. 0.5 mL of aliquots was taken at fixed time intervals from the medium outside the dialysis tube, and the medium was immediately replenished with the same amount of PBS or 10% FBS. Quantitation of CL released in PBS was measured by HPLC described above, while CL released in 10% FBS was quantified by UPLC/MS/MS analysis. The release ratio (\% released) was calculated by (2), where \( M_{\text{released}} \) is the mass of the released CL and \( M_{\text{encapsulated}} \) is the mass of CL encapsulated within the micelles.

\[
\%\text{released} = \frac{M_{\text{released}}}{M_{\text{encapsulated}}} \times 100\% .
\]

2.8. Pharmacokinetics Study. Male Sprague-Dawley rats (250–300 g) were anesthetized with zoletil (25 mg/kg, im) and xylazine (10 mg/kg, im), and body temperature was maintained at 37°C with a heating pad. Polyethylene catheters were inserted into the jugular vein for blood sampling. Rats were injected intravenously with an equivalent CL dose (22 mg/kg) of P-MCL, MCL, and free CL (the CL powder was dissolved in 0.9% NaCl-injectable solution by adding a few drops of 85% phosphoric acid). 500 \( \mu \)L of blood samples was collected into K\(_2\)EDTA tubes (BD, Franklin Lakes, NJ, USA) via the jugular vein at postinjection time points (0.05, 0.25, 0.5, 1, 2, 4, 8, and 24 hours). Plasma samples were obtained after centrifugation at 4000 rpm for 15 minutes and were stored at −80°C until analysis.

2.9. Plasma Sample Extraction. Plasma sample extraction was performed as previously reported [25]. Briefly, 10 \( \mu \)L of ROX solution (6 \( \mu \)g/mL in 50% acetonitrile) as the internal standard (IS), 60 \( \mu \)L of acetonitrile, and 50 \( \mu \)L of plasma sample were mixed. After centrifugation at 3500 rpm for 10 min, 15 \( \mu \)L of supernatant was transferred to a 96-well SPE plate conditioned with methanol and water. The extracted samples were eluted with 150 \( \mu \)L of acetonitrile and evaporated using a SpeedVac concentrator (Refrigerated Vapor Trap RVT4104, Thermo Fisher Scientific, Waltham, MA, USA). The residue was reconstituted in 150 \( \mu \)L of 50% acetonitrile and 5 \( \mu \)L of aliquot was injected into the UPLC/MS/MS system. The peak area ratio of CL to IS was calculated to determine the CL concentration.

2.10. UPLC/MS/MS Condition. UPLC/MS/MS condition was followed as previously reported [25]. Briefly, chromatographic separation was performed on a Waters ACQUITY UPLC system (Milford, MA, USA) with a Waters BEH C\(_{18}\) column (50 mm × 2.1 mm, 1.7 \( \mu \)m id, Milford, MA, USA) maintained at 40°C. The mobile phase consisted of solvent A (acetonitrile) and solvent B (50 mM ammonium acetate) with a flow rate of 0.3 mL/min. The initial mobile phase composition was 50% solvent A, changed linearly to 10% solvent A (0–1.5 min) and held for 0.2 min (1.5–1.7 min), followed by a return to the initial condition within 0.1 min (1.7–1.8 min), and kept for 0.2 min (1.8–2.0 min) for the column equilibrium. The UPLC system was connected to an ACQUITY TQD tandem mass spectrometer (Waters, Manchester, UK) as an electrospray ionization interface monitoring in positive ionization mode. Quantitation was performed using multiple reaction monitoring to study precursor → product ion transitions of the...
analyte and internal standard of m/z 748.77 → 83.04 and 837.84 → 116.05 for CL and ROX, respectively.

2.11. Pharmacokinetics Analysis. The plasma concentration-versus-time data of CL from the mean of three rats was analysed to determine the pharmacokinetic parameters using a noncompartmental model in WinNonlin Professional version 5.2 (Pharsight, Mountain View, CA, USA). The area under the curve (AUC), half-life (t_{1/2}), clearance, mean residence time (MRT), and volume distribution (V_d) were evaluated. The management of animal experiment and animal care and use of NHRI has been accredited by the AAALAC International.

3. Results and Discussion

3.1. Results

3.1.1. Optimization of PLGA Incorporation. To understand how much PLGA should be incorporated to stabilize micelle structure, P-MCLs with different PLGA/PEG-PE feeding ratios were prepared and incubated in 10% FBS. Their size changes were monitored by DLS (Figure 2) to examine the stability of P-MCLs over time. When the PLGA/PEG-PE feeding ratio was 0.5, the size of P-MCL remained almost the same around 75 nm from 0 to 65 hours at 37°C. In contrast, with the 0.25 feeding ratio, the size of P-MCL increased dramatically from 80 to 275 nm after 17 hours and exhibited an obvious cloudy appearance. The similar size increment (size change from 80 to 380 nm after 41 hours) and cloudy appearance were observed while the feeding ratio was 1. Therefore, the optimal PLGA/PEG-PE feeding ratios was determined as 0.5.

3.1.2. Characterization of P-MCL and MCL. P-MCL and MCL were prepared by a single emulsification-solvent evaporation method and were observed under TEM with negative stain. P-MCL has a spherical shape (Figure 3(a)), and MCL has a fluffy irregular shape (Figure 3(b)). Table 1 summarizes the general physicochemical properties of P-MCL and MCL. They have similar average diameters (below 40 nm), high CL loading contents (about 15%), and negative surface charges (around −35 mV).

3.1.3. In Vitro Release Profiles of P-MCL and MCL. The in vitro CL release profiles from P-MCL and MCL in different release media are given in Figure 4. In PBS (Figure 4(a)), during the initial 6 hours, 30% and 50% of the loaded CL were released from P-MCL and MCL, respectively. After 6 hours, the CL was released in a much slower manner from both P-MCL and MCL. After incubation for 48 hours, the CL released from P-MCL and MCL amounted to 51.3% and 81.1%, respectively. When the media was switched to 10% FBS (Figure 4(b)), there was no obvious burst release of CL from P-MCL and MCL. At 6 hours, only 7% of the loaded CL was released from P-MCL, while CL released from MCL was about 12%. The CL was gradually released from both P-MCL and MCL during the observation period for 72 hours. Overall, P-MCL had a slower CL release rate than MCL in both PBS and 10% FBS solution.

3.1.4. Pharmacokinetics. Pharmacokinetic profiles of three types of CL administration (P-MCL, MCL, and free CL) were obtained in male Sprague-Dawley rats. Figure 5 shows the CL concentrations in plasma. CL concentrations from all the three administrations decreased rapidly within the first hour and then decreased in a one-exponential-decay

![Figure 3: TEM images of (a) P-MCL and (b) MCL negatively stained by uranyl acetate (scale bars: 100 nm).](image)

**Table 1: Physicochemical properties of P-MCL and MCL.**

| Sample | Mean diameter (nm) | Zeta potential (mV) | Loading content (%) | Entrapment efficiency (%) |
|--------|--------------------|---------------------|---------------------|--------------------------|
| P-MCL  | 35.1 ± 14.3        | −34.2 ± 0.8         | 15.7 ± 0.9          | 45.7 ± 5.2               |
| MCL    | 32.0 ± 9.1         | −37.2 ± 1.3         | 15.1 ± 0.7          | 57.9 ± 5.5               |
P-MCL had the highest CL concentration in plasma at every time point (0.05, 0.25, 0.5, 1, 2, 4, 8, and 24 hours). The pharmacokinetic parameters were analysed and are presented in Table 2. The AUC value was in the order of P-MCL (24.41 ± 1.51 h·μg/mL) > MCL (13.92 ± 2.09 h·μg/mL) > free CL (9.66 ± 3.21 h·μg/mL), showing that the blood CL concentrations of P-MCL and MCL were 2.5-fold and 1.4-fold higher, respectively, than that of free CL. The clearance and $V_d$ were in the order of free CL > MCL > P-MCL. The clearance rates of P-MCL and MCL were 2.7-fold and 1.5-fold lower, respectively, than that of free CL. The AUC and clearance of P-MCL were 1.8-fold higher and 1.8-fold slower, respectively, than those of MCL.

3.2. Discussion. The stability issue of premature drug release from micelles resulting from the dynamic exchange of unimers or the adsorption of serum proteins is one of the obstacles for developing an effective micellar nanocarrier in pharmaceutical applications [26, 27]. The premature drug leakage could decrease drug therapeutic efficiency and lead to undesirable toxicity to other tissues. Therefore, it is important to enhance the stability of micelles for preventing premature drug release in the physiological environment.

In the present study, we incorporated hydrophobic PLGA into PEG-PE micelles to enhance the micelle stability. While the PLGA/PEG-PE feeding ratio is 0.5, P-MCL shows the best stability property in 10% FBS with no size increment over time, suggesting that no aggregation occurred (Figure 2). With this optimal amount of PLGA incorporation, P-MCL has a slower CL release rate than MCL has in both PBS and 10% FBS solutions (Figure 3), indicating its better micelle stability. In the pharmacokinetic study, P-MCL showed a higher AUC and a lower clearance than did MCL (Table 2), which means that P-MCL was cleared more slowly and supplied more available CL in vivo. Since free CL was cleared very fast, the lower clearance of P-MCL may be contributed by the less drug leakage due to the better protection from the higher micelle stability. The hydrophobic PLGA copolymer here plays a crucial role for increasing the hydrophobic interaction to the PE carbon chains of the PEG-PE unimers.
Despite of the decrease in the critical micellar concentration by adjusting the chemical composition of unimers [28, 29], micelle stability can be generally enhanced by chemical modification or by physical interaction approaches [30]. Among chemical modification approaches, the hydrophilic or hydrophobic segment of micelle unimers can be cross-linked to improve micelle integrity [23]. The cross-linked micelles had higher AUC than non-cross-linked micelles had in murine blood [31, 32]. Using the cross-linking strategy may reduce degradation or collapse of micelles, but whether the cross-linked products are still biocompatible and biodegradable is still a concern [33]. Meanwhile, drugs may not be releasable when they are entrapped in a nondegradable vehicle (only the released drugs are essential for the therapeutic efficacy) [27].

Instead of chemical modification, physical mixing of hydrophobic materials within the micelles is a convenient approach to enhance micelle stability. For example, a hydrophobic vegetable oil was introduced into pluronic micelles which exhibited less degradation than did original pluronic micelles upon dilution [34]. Physical mixing methods provide a simple way to improve micelle stability via increased noncovalent forces like hydrophobic interaction, aromatic π-π interaction, hydrogen bonding, and ionic interaction between unimers, drugs, and mixing materials [30]. Moreover, these kinds of noncovalent interactions may provide enough force to stabilize micelles, yet drugs can be released controllably from moderate entrapment.

Here we selected the physical mixing strategy to improve micelle stability by PLGA incorporation. Because of the convenience of physical mixing, we can further change the hydrophobic blending materials like different molecule weights or different L/G ratios of the PLGA or can adjust PEG lengths by switching PEG-lipid materials for optimized drug encapsulation or micelle stability.

4. Conclusions

Micelles composed of PEG lipids have emerged as promising drug delivery vehicles for hydrophobic drugs to enhance the drug availability or delivery efficiency. However, PEG-lipid micelles have a stability issue, in that unimers are in a dynamic equilibrium between micelles and bulk solution, resulting in an uncontrollable drug release or leakage. When a hydrophobic polymer such as PLGA was incorporated into the micellar core, the carbon chains of PEG-PE unimers could be interlaced by PLGA which reduce the dynamic exchange of PEG-PE unimers and thereby reduce the burst release and release rate of entrapped drugs. Moreover, micelles with improved stability by PLGA interlacing exhibited lower clearance, higher area under plasma concentration curve, and lower volume distribution than did nonstabilized micelles in a murine pharmacokinetic study, providing more advantages in controllable micellar delivery systems.

Conflicts of Interest

The authors report no declarations of interest.

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