Development of high drug-loading nanomicelles targeting steroids to the brain

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Abstract: The objective of this research was to develop and evaluate high drug-loading ligand-modified nanomicelles to deliver a steroidal compound to the brain. YC1 (5α-cholestan-24-methylene-3β, 5α, 6β, 19-tetraol), with poor solubility and limited access to the brain, for the first time, has been proved to be an effective neuroprotective steroid by our previous studies. Based on the principle of ‘like dissolves like’, cholesterol, which shares the same steroidal parent nucleus with YC1, was selected to react with sodium alginate, producing amphiphilic sodium alginate–cholesterol derivatives (SACDs). To increase the grafting ratio and drug loading, cholesterol was converted to cholesteryl chloroformate, for the first time, before reacting with sodium alginate. Further, lactoferrin was conjugated on SACDs to provide lactoferrin-SACDs (Lf-SACD), which was established by immune electron microscopy (IEM) and self-assembled into brain-targeting nanomicelles. These nanomicelles were negatively charged and spherical in nature, with an average size of <200 nm. The YC1 drug loading was increased due to the cholesteryl inner cores of the nanomicelles, and the higher the grafting ratio was, the lower the critical micelle concentration (CMC) value of SACD, and the higher drug loading. The in vitro drug release, studied by bulk-equilibrium dialysis in 20 mL of 6% hydroxypropyl-β-cyclodextrin solution at 37°C, indicated a prolonged release profile. The YC1 concentration in mouse brain delivered by lactoferrin-modified nanomicelles was higher than in those delivered by non-modified nanomicelles and YC1 solution. The unique brain-targeting nanomicelle system may provide a promising carrier to deliver hydrophobic drugs across the blood–brain barrier for the treatment of brain diseases.

Keywords: drug-loading, lactoferrin, nanomicelles, sodium alginate, cholesterol, brain-targeting

Introduction

YC1 (5α-cholestan-24-methylene-3β, 5α, 6β, 19-tetraol) (Figure 1) is a kind of oxysterol extracted for the first time from the soft coral Nephthea brassica in the South Sea of China.1 Our previous studies have demonstrated that YC1 plays a significant role in protecting neurons.2 However, it was found that the neuroprotection of YC1 after intravenous injection was substantially impeded by its poor solubility and nonspecific systemic distribution. Therefore, it is meaningful and necessary to develop an alternative brain-targeting delivery system of YC1 for its beneficial applications.

Recently, self-assembling micelles as promising drug delivery systems have been paid increasing attention on account of their uncomplicated preparing conditions and relatively high drug loading.3–4 According to the empirical rule of ‘like dissolves like’ (specifically, the greater the similarity between molecular structures, the higher the mutual solubility due to the stronger intermolecular attractions),5 if the inner core of nanomicelles presents...
Lactoferrin (Lf), one of the Tf family members, is a single-chain, iron-binding glycoprotein with a molecular weight (MW) of 80 kDa, which has several particular advantages as a brain-targeting ligand compared with Tf. First, the plasma concentration of Lf (238.0 ng/mL) is much lower than the saturation value of Lf receptors (LfRs) in the BBB.17,18 Second, thanks to its higher affinity with iron, Lf, by binding with iron as a complex, may cross the bovine brain capillary endothelial cells (BBCECs) more easily.19,20 Third, Lf is capable of traversing the BBCECs from the apical side to the basolateral side, suggesting that LfRs may be able to mediate more chemicals across the BBB.19 Moreover, Lf participates in the pathogenesis of brain lesions. It has been reported that the expression of LfRs increased in patients with Parkinson’s disease.21 More importantly, a marked elevation of Lf in the cerebrospinal fluid (CSF) of patients with acute stroke was observed.22 In another study, the enhanced endocytotic activity of Lf was revealed in a rat ischemia/stroke model in which transient focal ischemia was induced by using the filament occlusion of middle cerebral arteries.23 Therefore, it was concluded that such an increase in transcytosis of Lf did not involve the upregulation of the LfRs, but rather an increase of transcytosis transport.24 Accordingly, these findings suggested that Lf may be a promising brain-targeting ligand that could significantly enhance YC1, a potential therapeutic agent for central nervous system (CNS) disorders, to penetrate the BBB.

In this research, Lf-SACDs were synthesized and then self-assembling nanomicelles were prepared for loading and targeting delivery of YC1 to the brain. The hydrophilic cholesteryl inner core of the nanomicelles may increase the drug loading of YC1 owing to the structural similarity between cholesterol and YC1. Meanwhile, the Lf conjugated on the hydrophilic outer layer may enhance the LfRs-mediated endocytosis of nanomicelles, generating an elevation of YC1 cerebral distribution. The emulsification method was used to encapsulate YC1 into the nanomicelles. The particle size, zeta potential, morphology, grafting ratio, encapsulation efficiency, and the in vitro releasing profile of the YC1-loaded nanomicelles were studied. The brain-targeting activity of the YC1-loaded Lf-SACD nanomicelles was evaluated in vivo. The distribution of YC1 in major organs was characterized and the targeting parameters were calculated.

Materials and methods

Materials
Sodium alginate (Protanal LF 120M with MW of 180–250 kDa) containing 35%-45% of guluronic acid and
55%–65% of mannuronic acid was a kind gift from FMC BioPolymer (Philadelphia, PA, USA). Formamide (FA) and dimethyl formamide (DMF) were dried by molecular sieve. p-toluenesulfonic acid (pTSA) was dried at 80°C for 20 minutes in a vacuum. Cholesterol was bought from Qi Hua Armamentarium (Guangzhou, People’s Republic of China). The internal standard (ST009, 5α-androstane-3β, 5, 6β-triol), lactoferrin (Lf, from bovine), dicyclohexylcarbodiimide (DCC), N-hydroxysulfosuccinimide (NHSS), 1-ethyl-3-[3-dimethylaminopropyl] carbodiimide hydrochloride (EDC), 4-(N,N-dimethylamino)pyridine (DMAP), and 2-(4-morpholino)ethanesulfonic acid (MES) were purchased from Sigma-Aldrich (St Louis, MO, USA) and used without further purification. The anti-Lf antibody was a rabbit polyclonal antibody raised against bovine. Hydroxypropyl-β-cyclodextrin (HP-β-CD) was provided by Roquette (Lestrem, France). All the other chemicals and solvents were of chromatographic and pharmaceutical grade.

Synthesis and characterization of SACD

Cholesterol (0.66 g) and triphosgene (1.00 g) were reacted in tetrahydrofuran (THF) at 40°C for 30 minutes,25 and only one hydroxyl of cholesterol was converted into cholesteryl chloroformate. Then, according to Yang et al.,2 water-free pTSA (0.32 g) and sodium alginate (1.00 g) were dispersed in a mixture of 38 mL of FA/DMF (10:9, v/v) in a 250 mL round bottom flask, and the sodium alginate was partially hydrolyzed into alginic acid at 50°C by stirring for 30 minutes. DCC (0.40 g), DMAP (0.48 g), and the solution of chloroform containing activated cholesteryl chloroformate were then added to the suspension of sodium alginate. The reaction between alginic acid and the chloroformate of cholesterol was carried out at 25°C–30°C by stirring for 24 hours. The product was purified with 200 mL of ethanol, with stirring at 40°C, and then centrifuged. The obtained pellets were dried under vacuum for 24 hours at 40°C, dissolved in distilled water, and neutralized by adding the solution of 1.5% Na2CO3. The obtained solution was dialyzed for 3 days and further freeze-dried to collect SACD.

The 1H NMR spectra of SACD were studied by 1H nuclear magnetic resonance spectroscopy (Avance III 400 Plus; Bruker, Billerica, MA, USA); FT-IR measurement was carried out on a Fourier transform infra-red spectrometer (Tensor37, Bruker) at 0.5 cm−1 using KBr pellets.

Synthesis of Lf-SACD

Lactoferrin was conjugated to SACD through the carboxylic acid groups of alginic acid. According to Liu et al.,26 100 and 40 mM solutions of EDC and NHSS were prepared using 0.1 M MES solution, respectively. SACD was added to the 0.1 M MES (0.5 mL) with 40 μL of EDC and NHSS, and the obtained solution was incubated at 4°C for 1 hour. Then, Lf was dissolved into phosphate-buffered saline (PBS, pH 7.4) to form a 1.0 mg/mL solution, and the activated SACD (5.0 mg) solution and Lf solution (125 μL) were reacted for 24 hours at room temperature. The product was dialyzed against distilled water for 3 days using the Spectra/Pro membrane with a molecular weight cut-off of 100 kDa and the final product was obtained after lyophilization.

Characterization of Lf-SACD

Immune electron microscopy (IEM) was harnessed to confirm the successful conjugation of Lf to SACD. Briefly, Lf (0.13 mg), SACD (1.2 mg), and Lf-SACD (1.2 mg) were separated by 10% SDS-PAGE. After electrophoresis, the proteins were electroblotted to a polyvinylidene fluoride (PVDF) membrane (0.45 μm; Millipore, Billerica, MA, USA). The PVDF membrane was then blocked with 5% skimmed milk (Bio-Rad, Hercules, CA, USA) in Tris HCl-buffered saline solution (TBS, 0.1 g KCl, 4.0 g NaCl, and 1.5 g Tris, pH 7.4) for 2 hours. Lf antibodies were added and the system was incubated for 2 hours, and then reacted with the rabbit antibodies for 1 hour. The immune complexes were visualized using the electrochemiluminescence (ECL) detection reagents.

Preparation of self-aggregating nanomicelles

A microphase separation method was used to prepare nanomicelles. Briefly, 20.0 mg of SACD and 8.0 mg of YC1 were dispersed into 10 mL of tetrahydrofuran (THF), respectively, and then added to the distilled water (10 mL) drop-by-drop under stirring until THF was completely volatilized.27 Subsequently, YC1-SACD nanomicelles were formed. Lf-SACD nanomicelles were generated by a similar procedure.

Characterization of the blank and YC1-loaded nanomicelles

Micelle size and zeta potential

The average size and zeta potential of nanomicelles were determined by Zetasizer Nano ZS90 (Malvern Instruments Ltd, Malvern, UK). The concentration of the nanomicelles was about 1.0 mg/mL in distilled water.28
Analysis of morphology by transmission electron microscope
The morphology of YC1-loaded and YC-unloaded nanomicelles was observed using transmission electron microscopy (TEM, JEM2010 frozen type; JEOL, Tokyo, Japan). The sample was filtrated through a 0.45 μm Millipore membrane filter, and then 40 μL of the obtained filtrate was placed on a copper chip, which was exposed to air to dry at room temperature and then mounted on the TEM for observation.

Critical micelle concentration of SACD
The critical micelle concentration (CMC) of SACD was estimated by spectrofluorophotometer (LS55; Perkin Elmer, Waltham, MA, USA), using pyrene as a fluorescence probe. The excitation wavelength was 334 nm and the slit width was 2.5 nm. The fluorescence emission spectra were chosen between 360–500 nm with a slit width of 2.5 nm. SACD was dissolved in 0.15 mol/L of NaCl solution to form a concentration range from 9.0 to 1.0 × 10^{-3} mg/mL. Pyrene (Acros Organics, Geel, Belgium) was added to the SACD solutions to obtain the pyrene concentration of 2 μmol/L. All samples were treated by ultrasonic sound to make sure the pyrene was completely embedded into the hydrophobic microdomains. The intensity ratios of I_{373}/I_{384} were plotted as a function of logarithm concentration of the polymers. The CMC value was taken from the intersection of the tangent to the curve at the inflection with the horizontal tangent through the points at low concentrations.

Determination of grafting ratio
As described above, after 24 hours reaction of cholesterol and sodium alginate, 200 mL of ethanol was added to the solution to extract the unreacted cholesterol. After centrifugation, the volume of the supernatant was determined and diluted to an appropriate concentration to be measured by high performance liquid chromatography (HPLC, 1200; Agilent, Santa Clara, CA, USA) with an ODS column (Phenomenex, 250 × 4.6 mm, 5 μm) held in an oven at 38°C. The mobile phase consisted of methanol with a flow rate of 1.0 mL/minute. The ultraviolet detecting wavelength was 205 nm and the injection volume was 10 μL. The grafting ratio was calculated using the following formula:

\[
\text{Grafting ratio} = \frac{\text{weight of YC1 measured in total eluate}}{\text{weight of YC1 added to the reaction system}} - \frac{\text{total amount of sodium alginate}}{100}
\]

Measurement of encapsulation efficiency and drug loading
The encapsulated and unencapsulated YC1 were separated using a column of Sephadex G-50 (Amersham Biosciences, Uppsala, Sweden). Sephadex G-50 was pretreated and packed into a 5 mL syringe. Distilled water went through the syringe at a rate of 0.2 mL/minute for 1 hour to remove air bubbles. The sample was then loaded into the column, with distilled water as the mobile phase at a flow rate of 1.0 mL/minute. Elute from 0–16 minutes was collected and the concentrations were determined by liquid chromatography coupled with mass spectrometry (LC-MS, 6120; Agilent). The LC column was Agilent XDB-C18 (50 × 4.60 mm, 1.8 μm). The mobile phase, consisting of methanol/water (90/10, v/v), was delivered at a flow rate of 0.5 mL/minute. The column oven temperature was maintained at 30°C and the injection volume was 1.0 μL. The Agilent’s single quadrupole mass spectrometer (6120 series) was equipped with an atmospheric pressure chemical ionization (APCI) source. At first, the full scanning mode was used to select the characteristic ions to be monitored for YC1 and the internal standard (ST009). Next, selective ion monitoring (SIM) was used to monitor the molecular weight (MW) of 403.3 g/mol for YC1 and 273.1 g/mol for ST009. The standard curve was constructed by dissolving YC1 in methanol at the concentration range of 1.0–25 μg/mL with an r^2 value of 0.9996. Encapsulation efficiency (%) = weight of YC1 measured in total eluate/weight of total YC1 × 100; drug loading (%) = weight of YC1 measured in total eluate/weight of the blank nanomicelles.

In vitro drug release
First, the solubility of YC1 in different solvents was determined and 6% HP-β-CD solution was proved to be the most effective solvent among 30% alcohol, 40% alcohol, 50% alcohol, 1% Tween 80, 2% Tween 80, 1% Tween 80 plus 50% alcohol, 4% HP-β-CD, and 6% HP-β-CD.

Second, YC1-loaded nanomicelles (2.0 mg) purified by the Sephadex G-50 column were dissolved in 2.0 mL of distilled water. The solution was added to the dialysis membrane bag with a molecular weight cut-off of 14 kDa. The membrane bag was then immersed into 20 mL of 6% HP-β-CD solution at a shaking rate of 100 times per minute at 37°C. A volume of 500 μL release medium was taken at 0, 0.25, 0.5, 1, 2, 4, 6, 8, 10, 12, and 24 hours, respectively, and immediately replaced with the same volume of fresh release medium. The YC1 content in the release medium was determined by LC-MS as mentioned above. Each experiment was repeated in triplicate. The release rate was calculated
with the formula: $RR = (W_i/W_{\text{total}}) \times 100\%$, where RR is the drug release rate (%), $W_i$ is the measured amount of YC1 at the time point of $i$th hour in the release medium, and $W_{\text{total}}$ is the total amount of YC1 in the equal volume of nanomicelle suspensions prior to dialysis. The release characteristics were explored by the similar factor method, the zero grade model, the first grade model, the Higuchi model, and the Regter–Peppas model.

Biodistribution of YC1 in vivo

All procedures involving animals were approved by the Institutional Authority for Laboratory Animal Care and Ethics Committee. Adult Kunming mice of either sex (18–22 g, Laboratory Animal Center, Sun Yat-sen University, Guangzhou, People’s Republic of China) were injected with Lf-modified nanomicelles (LF group), SACD nanomicelles without modification (MC group), and YC1 solution (YC group) via the tail vein at a dose of 8 mg/kg, respectively. The mice were anesthetized by inhalation of diethyl ether before blood sample collection.

At 15 minutes, 30 minutes, 1 hour, 2 hours, 4 hours, and 8 hours, blood samples were collected from the ocular artery directly after eyeball removal and placed into test tubes containing sodium citrate. Plasma was immediately separated by centrifugation. After that, the mice were sacrificed and the brains, hearts, livers, lungs, and kidneys were excised. The organs were rinsed three times in physiological saline (0.9% NaCl solution) to wash away the residual blood, blotted on filter paper, weighed, and then homogenized in a 3-fold volume of saline. The separated plasma and the organ homogenates were stored at −20°C until assay.

Acetic ether (5 mL), 0.12 mg/mL ST009 (10 µL) as the internal standard solution, 20.3 mg/mL EDTA (ethylene diamine tetraacetic acid, 10 µL), and 20.9 mg/mL BHT (butylated hydroxytoluene, 2.0 µL) were added to a plasma sample (1.0 mL) and mixed by vortex for 5 minutes. The mixture was centrifuged at 4,000 rpm for 10 minutes and then the supernatant was dried by nitrogen gas flow. Subsequently, 250 µL of methanol was added to the dried sample and mixed by vortex. The solution was centrifuged at 4,000 rpm for 10 minutes and 5.0 µL of the clear supernatant was injected into the LC-MS for analysis as previously described. This procedure was repeated for processing the tissue samples.

Pharmacokinetic analysis and evaluation of targeting activity

The concentration data of YC1 in plasma and tissue samples were pooled to obtain the mean concentration of YC1. The pharmacokinetics of YC1 was described as a two-compartment open model and the pharmacokinetic parameters in the plasma and organs were obtained. The area under the concentration–time curve (AUC) was calculated using the software program 3p97 (Mathematical Pharmacology Professional Committee of China; Shanghai, People’s Republic of China). The mean residence time (MRT) was calculated by statistic moment program (DAS, Mathematical Pharmacology Professional Committee of China). To evaluate the targeting attribute of Lf-modified nanomicelles, MRT (mean residence time), overall targeting efficiency (TE), targeting index (TI), and relative overall targeting efficiency (RTE) were calculated as per the following equations (Equations 1–5) and compared with those of the MC and YC groups.\(^\text{30}\) In Equation 1, the denominator represents the total of YC1 exposure to all the tissues.

$$\text{TE} (%) = \frac{(\text{AUC}_{0\rightarrow\infty_{\text{LF}}})}{\sum_{i=1}^{\infty_{\text{group}}} (\text{AUC}_{0\rightarrow\infty_{\text{group}}})} \times 100$$  
$$\text{TI}_{\text{LF}} (%) = \frac{(\text{AUC}_{0\rightarrow\infty_{\text{MC}}})}{(\text{AUC}_{0\rightarrow\infty_{\text{MC}}})} \times 100$$  
$$\text{RTE}_{\text{LF}} (%) = \frac{(\text{TE}_{\text{MC}} / \text{TE}_{\text{MC}})}{(\text{TE}_{\text{MC}} / \text{TE}_{\text{MC}})} \times 100$$

Statistical analysis

Data were expressed as means ± SD. Statistical significance of the difference was assessed by one-way analysis of variance followed by post-hoc tests to determine the significance among groups in SPSS (v17.0; SPSS, Inc, Chicago, IL, USA). A P-value of <0.05 was considered significant.

Results and discussion

Synthesis and characterization of SACD

The synthetic route of SACD is briefly illustrated in Figure 2. In \(^1\)H NMR spectra, several proton peaks at 1.0–2.5 ppm were assigned to the cholesteryl chain of cholesterol except H3 and H6.\(^7\) Moreover, the signals between 3.6 and 4.5 ppm could be considered as the methine protons of hexuronic acid residues of the alginate main chain\(^26\) and the H3 and H6 of the cholesteryl moiety. The single peak at 4.5–4.8 ppm was attributed to the methine protons of sodium alginate. The \(^1\)H NMR spectrum of SACD (Figure 3C) displayed typical peaks of the protons on the alginate chain (Figure 3B) and the peaks (1.0–2.5 ppm) of protons on the carbons of cholesterol (Figure 3A). The difference between the spectra of SACD (Figure 3C) and the physical mixture of sodium alginate and cholesterol (Figure 3D) indicated that cholesterol was successfully introduced onto sodium alginate to produce SACD.
In the FT-IR spectrum (Figure 4), the absorption of double peaks between 1700 and 1600 cm$^{-1}$ (Figure 4C) indicated the presence of two kinds of carbonyl groups from both carboxyl groups and newly formed ester bonds. However, in the FT-IR spectrum of sodium alginate (Figure 4B), only one peak was observed between 1700 and 1600 cm$^{-1}$. This distinction suggested that esterification between sodium alginate and cholesterol occurred.

**Characterization of Lf-SACD**

The electrophoretogram of Western blot analysis was shown in Figure 5. Column A supports the position of the
markers (kDa), while column B indicates that lactoferrin had a molecular weight of 70–80 kDa. Column C proved that the controlled group, SACD, was not immunogenic, while column D showed that the SACD was modified with lactoferrin, resulting in a molecular weight of 250–330 kDa for Lf-SACD. Therefore, the IEM results confirmed that lactoferrin was successfully modified onto SACD.

**Characterization of self-assembly nanomicelles**

Particle sizes and zeta potentials of the unloaded and YC1-loaded nanomicelles were investigated, respectively. The particle sizes of more than 70% of the YC1-unloaded nanomicelles ranged from 100 to 140 nm whereas the diameters of more than 70% of YC1-loaded nanomicelles were in the 140–180 nm range. Thus, the increased size of YC1-loaded nanomicelles demonstrates that YC1 molecules were encapsulated into the nanomicelles. In addition, the zeta potential was about −100 mV for both types of nanomicelles. This could be explained by the fact that the alginate ionized a large number of anionic −COO − carrying negative charges, and thus the whole system appeared to be negative.

The TEM images (Figure 6) of the Lf-SACD self-aggregates presented some core/shell-like particles with an approximate size of 100 nm. The relatively uniform size distribution and approximately spherical and regular shape for the YC1-loaded nanomicelles confirmed that the existence of intramolecular and intermolecular hydrophobic interactions among the cholesteryl grafts made the Lf-SACD self-assemble into nanomicelles.

**CMC of SACD**

In this research, two synthesis schemes of SACD were performed, where the only difference between them was whether cholesterol was converted into cholesteryl chloroformate by triphosgene. It was revealed that the reaction between hydroxyl compounds and triphosgene would produce unstable chloroformate with high reactivity. Thus, the synthesis route in which cholesterol was pretreated by triphosgene was assumed to generate SACD with a higher grafting ratio and lower CMC.

When cholesterol was not pretreated (Figure 7A) or pretreated (Figure 7B) by triphosgene for the synthesis of SACD, the CMC values of the obtained nanomicelles were about 2.0 mg/L and 0.8 mg/L, respectively. According to Yang et al, there were several types of intra- and inter-molecular interactions controlling SACD to self-assemble, such as the hydrophobic interaction among cholesteryl grafts, the hydrophilic bonding between the backbones and water molecules as well as among sodium alginate backbones themselves. Amongst these interactions, those between cholesteryl grafts played the most important role. In other words, more cholesterol connected with sodium alginate may generate a lower CMC value for SACD. Accordingly, the nanomicelles obtained from the pretreated cholesterol possessed a lower CMC value (Figure 7B), which suggested that the SACD had a higher cholesterol grafting ratio. It is important and meaningful in practice, as a lower CMC value for nanomicelles guarantees their use in very dilute media, such as in body fluids.

**Figure 5** Electrophoretogram of Western blot analysis for (A) markers; (B) LF; (C) SACD; and (D) Lf-SACD.

*Abbreviations:* LF, lactoferrin; SACD, sodium alginate–cholesterol derivatives.

**Figure 6** Microphotograph of LF-SACD self-aggregates under transmission electron microscopy (×80,000); (A) YC1-loaded; and (B) YC1-unloaded.

*Abbreviations:* LF, lactoferrin; SACD, sodium alginate–cholesterol derivatives YC1, 5α-cholestane-24-methylene-3β, 5α, 6β, 19-tetraol.

Shelf-life stability of YC1-loaded nanomicelles from lyophilization was studied by determining both the changes of the encapsulation efficiency and monitoring the particle size changes of nanomicelles after storage at 4°C for 15 days, 30 days, and 60 days. It appears that the nanomicelles displayed considerable stability and that only negligible changes of the particle size and the encapsulation efficiency were observed.
Drug loading and the relationship between the grafting ratio and encapsulation efficiency

SACDs with distinct grafting ratios were fabricated from different ratios of cholesterol and sodium alginate. Figure 8 shows that encapsulation efficiency improves with an increase in grafting ratio, suggesting a positive association between them. It was reported that the encapsulation efficiency of nanomicelles depends on the structure of drugs and the core block of the polymers. If the hydrophobic core of the polymers is similar to the structure of the loaded drugs, then the encapsulation efficiency would increase by thousands-fold. In this study, YC1 is one kind of oxysterol with the structure of a steroid that is identical to the parent nucleus of cholesterol. Therefore, with an increase in the cholesteryl grafting ratio, the intermolecular interaction between cholesteryl and YC1 would build up and YC1 could be entrapped into the inner core more easily and with more stability. It should be emphasized that Lf-SACD nanomicelles showcased higher drug loading for hydrophobic drugs when compared with other amphiphilic polymer-based nanocarriers.

In vitro drug release

Figure 9 shows that the accumulative release rate of YC1 from the nanomicelles could reach more than 80% in the first 12 hours in the releasing medium of 6% HP-β-CD and that no dumping effect was observed in the first 5 hours. The release behavior could last for 12 hours, implying that YC1 was well encapsulated into the nanomicelles. Thus, the nanomicelles may prolong the retention time of YC1 in the brain, extending...
the potential neuroprotection time. Furthermore, different release mechanisms were proposed for YC1-loaded nanomicelles releasing in HP-β-CD. Based on the values of $r^2$ in Table 1, the release from the nanomicelles most likely fits to the Ritger–Peppas equation, which suggests that the release mechanism was due to the joint effects of drug diffusion and skeleton dissolution.

**Table 1** Models and equations of YC1 releasing from nanomicelles in 6% HP-β-CD at 37°C

<table>
<thead>
<tr>
<th>Model</th>
<th>Equation</th>
<th>$r^2$</th>
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<tbody>
<tr>
<td>Zero-order</td>
<td>$y = 3.4191x + 27.247$</td>
<td>0.7221</td>
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<tr>
<td>First-order</td>
<td>$y = -0.0897x + 4.3176$</td>
<td>0.8882</td>
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<tr>
<td>Higuchi</td>
<td>$y = 20.266x + 4.965$</td>
<td>0.9186</td>
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<td>Ritger–Peppas</td>
<td>$y = 0.5943x + 2.9473$</td>
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**Abbreviation:** YC1, 5α-cholestane-24-methylene-3β, 5α, 6β, 19-tetraol.

**Figure 10** Concentration of YC1 in plasma and tissues after iv administration of YC1-loaded nanomicelles (MC), YC1-loaded Lf-modified nanomicelles (LF), and YC1 solution (Yc) ($n = 5$).

**Abbreviations:** YC1, 5α-cholestane-24-methylene-3β, 5α, 6β, 19-tetraol; Lf, lactoferrin.
Biodistribution of drugs in mice

The distribution of YC1 in the plasma and tissues of mice after iv administration of YC1-loaded Lf-modified nanomicelles (LF), YC1-loaded nanomicelles (MC), and YC1 solution (YC) separately was investigated. The results (Figure 10) indicate that the YC1 concentration in the brain for the LF group was significantly higher than those for the other two groups; the concentration in the brain reached its maximum about 4 hours after injection. In some excretory organs such as liver and kidney, YC1 concentration of the MC and LF groups was much lower than that of the YC group. This might be ascribed to the fact that nanomicelles can attenuate the elimination and extend the residence time of YC1, and thus the drug could be targeted to a specific organ with long circulating characteristics.

Pharmacokinetic analysis and evaluation of targeting activity

Based on the pharmacokinetic analysis and evaluation of targeting activity for YC1 in mice after iv administration of YC1 solution, YC1-loaded modified or non-modified nanomicelles, clear differences between these groups were found in the YC1 concentrations in organs (Figure 10), the pharmacokinetic parameters (Tables 2 and 3), and the TE (Tables 4 and 5; Figure 11). In comparison with YC and MC groups, the brain TE of the LF group was greatly improved. For example, the brain TE was enhanced from 6.9% for the YC group to 18.7% for the LF group (Table 4), and the RTE of the LF group was about two-fold greater than that of the MC group (Table 5). Conclusively, Lf-modified nanomicelles played a rather significant role in enhancing the brain TE of YC1. The noticeable enhancement of YC1 CNS distribution through the delivery system of Lf-modified nanomicelles was mainly derived from the following reasons. A large amount of YC1 solution was distributed in the liver, lung, and kidney, and was metabolized and excreted quickly; however, the YC1-loaded nanomicelles had a significantly lower YC1 concentration in these organs, and thus YC1 was eliminated at a slower rate. This phenomenon could be attributed to the fact that nanomicelles could remarkably encapsulate the steroidal YC1 to the hydrophobic inner core owing to their structural resemblance and simultaneously endow YC1 with the properties of sustained release, targeting, and lower accessibility to metabolic enzymes. On the other hand, there were measurable differences between the MC group and LF group in brain AUC, TI, and RTE, showing that lactoferrin enhanced the cerebral targeted delivery of YC1. This proved that lactoferrin receptor-mediated endocytosis actually assisted Lf-modified YC1-containing nanomicelles crossing the BBB. In summary, these combining factors contributed to the elevation of YC1 in CNS distribution and reduction in other main tissues.

Interestingly, the YC1 content of the MC group in the lungs was much higher than that of the YC group, showing a passive

Table 3 Pharmacokinetic parameters of YC1 of YC group, MC group, and LF group in brain tissue

<table>
<thead>
<tr>
<th>Parameters</th>
<th>AUC$_{0-\infty}$ (µg h/g)</th>
<th>MRT (h)</th>
<th>t$_{1/2}$ (h)</th>
<th>VRT (h)</th>
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</thead>
<tbody>
<tr>
<td>YC group</td>
<td>2.8</td>
<td>3.9</td>
<td>2.70</td>
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</tr>
<tr>
<td>MC group</td>
<td>4.1</td>
<td>4.1</td>
<td>2.84</td>
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<td>LF group</td>
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</tbody>
</table>

Notes: LF group = YC1-loaded Lf-modified nanomicelles; MC group = YC1-loaded nanomicelles without conjugating Lf; YC group = YC1 solution.

Abbreviations: AUC, area under the curve; MRT, mean residence time; VRT, variance of mean residence time; h, hour; YC1, 5α-cholestan-24-methylene-3β, 5α, 6β, 19-tetraol; Lf, lactoferrin.

Table 4 Overall TE of LF, MC, and YC groups

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Plasma</th>
<th>Heart</th>
<th>Liver</th>
<th>Lung</th>
<th>Kidney</th>
<th>Brain</th>
</tr>
</thead>
<tbody>
<tr>
<td>YC group</td>
<td>1.2</td>
<td>8.4</td>
<td>57.2</td>
<td>9.4</td>
<td>27.1</td>
<td>6.9</td>
</tr>
<tr>
<td>MC group</td>
<td>3.4</td>
<td>11.1</td>
<td>27.3</td>
<td>21.7</td>
<td>24.0</td>
<td>12.5</td>
</tr>
<tr>
<td>LF group</td>
<td>2.1</td>
<td>8.2</td>
<td>28.1</td>
<td>13.6</td>
<td>20.5</td>
<td>18.7</td>
</tr>
</tbody>
</table>

Notes: LF group = YC1-loaded Lf-modified nanomicelles; MC group = YC1-loaded nanomicelles without conjugating Lf; YC group = YC1 solution.

Abbreviations: TE, targeting efficiency; YC1, 5α-cholestan-24-methylene-3β, 5α, 6β, 19-tetraol; Lf, lactoferrin.
targeting effect of YC1-loaded unmodified nanomicelles in the lungs (Table 4 and Figure 10). This phenomenon might be accounted for by the dual reasons that a) the lung is a highly perfused organ by circulating blood and b) the cells in the reticuloendothelial system (RES) of the lung tissue phagocy-
tosed the nanomicelles when the particle size was in the range of 30–150 nm.36 However, the LF-modified nanomicelles showed much lower YC1 concentration in the lungs due to the ability of active targeting of the preparation.

Considering the significant differences in brain AUC, TE, TI, and RTE among the three groups, it is logical to surmise that the mean residence time (MRT) and t_{1/2} of the LF group should extend far more than those of the MC and YC group. However, the retention time of YC1 in the brain for the LF group was not as long as expected. This might be ascribed mainly to two reasons: a) the blood sampling time was not long enough and there was still plenty of YC1 remaining in the brain after 8 hours (Figure 10); and b) the amount of lactoferrin successfully conjugated onto SACD was insufficient. Since only 125 µg of lactoferrin was added to 5.0 mg of SACD to react, the active brain-targeting effect was limited by the inadequate number of lactoferrin ligands on the surfaces of nanomicelles.

**Conclusion**

YC1-loaded nanomicelles, self-assembling from the amphiphilic polymer LF-SACD, have been successfully prepared for drug delivery across the BBB. Firstly, one of the most significant features of the delivery system was the hydrophobic inner core that can accommodate a relatively large amount of steroideal YC1 owning to the ‘like dissolves like’ rule. Moreover, the drug distribution in mice showed that the LF-modified nanomicelles exhibited the properties of sustained release, drug protection, and a brain-targeting effect. The presence of lactoferrin on the surface of the nanomicelles promoted the accumulation of YC1 in the brain, achieving the goal of increasing the cerebral concentration of YC1. Finally, for the first time, cholesterol was chlo-
roformylated in this research before reacting with sodium alginate, resulting in amphiphilic polymer SACD with a higher grafting ratio. This further led to increasing the drug loading, reducing the CMC value, and enhancing the stability of micelles in dilute solutions. In short, it can be concluded that LF-modified sodium alginate–cholesterol nanomicelles show a potential to be a highly efficient drug carrier of hydrophobic therapeutic agents for brain targeting.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


