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Chain length effect on drug delivery of chrysin modified mPEG-PCL micelles

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Four chrysin modified mPEG-PCL block copolymers with different chain length of mPEG and PCL blocks were synthesized and self-assembled into micelles to load anticancer drug doxorubicin (DOX). The effect of block chain length on drug delivery was investigated. The four block copolymers were characterized by 1H NMR, GPC and DSC. The drug loading contents of all the micelles were higher than 20%, the mPEG2k-PCL5k-CHS micelles showed the highest drug loading content and encapsulation efficiency of 26.8% and 93%, respectively. The micelles were spherical with size increasing after drug encapsulation, the mean size of drug loaded micelles was around 100 nanometers. The π-π stacking interaction between micelles and DOX was evoked. mPEG2k-PCL5k-CHS micelles exhibited the best profile in sustained-release. The cellular uptake and IC50 revealed that the DOX loaded mPEG2k-PCL5k-CHS micelles showed the best anticancer activity in vitro.

Introduction

Polymeric micelles have been attracted great interest to biomaterials scientists and pharmacists as nanocarriers for anticancer drug delivery due to their advantages in long circulation and easy functionalization.1 The unique core-shell architecture of polymeric micelles trapped lipophilic anticancer drugs in the hydrophobic cores to enhance the solubility of anticancer drugs and protect drugs to avoid degradation via enzymes in the delivery process in vivo.2 The clinical trials3 of polymeric micelles based nanomedicine encouraged more and more researchers and pharmaceutical companies to focus on the development of polymeric micelle formulations.

Amphiphilic poly(ethylene glycol)-poly(ε-caprolactone) (PEG-PCL) diblock copolymers could self-assembly into polymeric micelles in aqueous solution. As biodegradable and biocompatible polymers, PEG-PCL were extensively investigated for anticancer drug delivery.4 Exciting progresses were received in PEG-PCL polymeric micelle drug delivery systems for the specificity of hydrophobic PCL segments as cores, which were favorable for drug loading as well as release diffusion due to the coexistence of flexible amorphous and rigid crystal domains with low glass transition temperature (about -60°C) and crystal melting temperature (about 60°C).3 All kinds of anticancer drugs including doxorubicin, paclitaxel, camptothecin and 5-fluorouracil have been reported to load in PEG-PCL micelles for the treatment of different types of cancers on animal models.

To polymeric micelles, the balance between hydrophilic and hydrophobic segments decided the self-assembly behaviors and properties of micelles.5 With the adjustment of compositions and hydrophilic/hydrophobic balance of amphiphiles, the morphology of polymeric micelles could be controlled in spherical,10 rod,11 flow-like12 and worm13. The balance between hydrophilic and hydrophobic segments also affected drug loading, release profile and anticancer activity.14 The chain length of PEG in PEG-PCL micelles was reported to act an important role in the fabrication of drug delivery systems.15

Amphiphilic copolymers aggregated together via self-assembly to form polymeric micelles,16 the weak physical interactions within polymeric micelles were considered not strong enough to maintain the aggregation, the dissociation was possibly occurred during the delivery in blood stream, thus, crosslink of shells or cores was carried out to stabilize the polymeric micelles,5 however, the solidified micelles were partially non-degradable and the hydrodynamic characteristics of polymeric micelles were seriously affected.17,18 In our previous work, we developed a new strategy to stabilize polymeric micelles with introducing π-π stacking interaction between hydrophobic moieties and anticancer drugs. A series of polymeric micelles were fabricated and the promising stabilization was exhibited.14,19-24 Recently, small molecules with different π-conjugated moieties of cinnamic acid, coumarin derivative and chrysin were immobilized on the terminal groups of PCL segments in PEG-PCL micelles to evoke π-π stacking interaction with anticancer drug doxorubicin, significant drug loading properties and anticancer activities both in vitro and in vivo were exhibited in chrysin modified PEG-PCL micelles.

In this paper, the chain length effects of chrysin modified PEG-PCL micelles on the drug delivery of doxorubicin were investigated in details. Four PEG-PCL diblocks copolymers with the block molecular weights of 2000 and 5000 were synthesized. The influence of both hydrophilic and hydrophobic chain length
on the movement of copolymer chains, self-assembly, drug release profiles and in vitro anticancer activity of polymeric micelles were investigated.

**Materials and method**

### Materials

α-methoxy-poly(ethylene glycol)s (mPEG, Mw =2000 and 5000 g/mol), stannous octoate (Sn(Oct)), ε-caprolactone (CL) and Hoochst were purchased from Sigma-Aldrich Co. Chrysin (CHS), methylbenzenesulfonfyl (TsCl) and triethylamine (TEA) were purchased from Asta Tech Pharmaceutical (Chengdu, China) and used as received. Doxorubicin hydrochloride (DOX·HCl, Zhejiang Hisun Pharmaceutical China) was deproteinated according to the method previously reported. 

### Characterizations

The 1H NMR spectra were performed on Bruker Avance II NMR spectrometer at 600 MHz using CDCl₃ as solvents with 0.5% tetramethylsilane as the internal standard. Gel permeation chromatography (GPC) was employed to characterize the molecular weight and the molecular weight distribution. GPC analysis was performed on a SDV Linear M5µ8Χ300 mm column (Polymer Standard Service, Mainz, Germany) with chloroform as the eluent (1 mL/min) and polystyrene standards for column calibration. The eluent was analyzed with a Waters HPLC system equipped with a model 1515 pump, a 717 plus autosampler, and a 2414 refractive index (RI) detector. The thermal properties of block copolymers were determined by differential scanning calorimetry (DSC, Q2000 TA Instruments), about 5 mg samples in hermetically closed aluminum pans were subjected to a heat1cool1heat program from 180 to +120 °C at a heating rate of 10 °C/min and a cooling rate of 5 °C/min. The X-ray diffraction was recorded on X’Pert ProMPD. Dynamic light scattering (DLS) measurements were carried out using a dynamic light scattering spectrometer (Malvern ZetasizerNano ZS) to determine the size and size distribution of micelles. Transmission electron microscopy (TEM, FEI Tecnai G2 F20 S-TWIN) was used to observe the morphology of micelles. The TEM samples were prepared by dropping the fresh solution onto Quantifoil holey carbon foil (Micro Tools GmbH, Germany) and dried overnight at room temperature. The π-π interactions between DOX and micelles were determined by UV-vis absorption (Spectord 200 PLUS) and Fluorescence spectra (HITACHI F-700).

### Synthesis of mPEG-PCL diblock copolymers

mPEG-PCL diblock copolymers were synthesized by ring-opening polymerization of ε-CL with mPEG homopolymer as macroinitiator and Sn(Oct)₂ as the catalyst. Prescribed amount of mPEG, ε-CL and Sn(Oct)₂ (ca. 0.1% of ε-CL in molar amount) were mixed in a round-bottom flask connected with a vacuum joint. The mixture was degassed in vacuum. The flask was sealed off and placed in an oil bath and polymerized at 130°C for 48 h.

The product was cooled at room temperature, dissolved in chloroform and purified by precipitating into large amount of cold diethyl ether. The precipitated product was vacuum-dried at 40 °C.

### Synthesis of mPEG-PCL-tosyl and mPEG-PCL-CHS

mPEG-PCL and tosyl chloride were dissolved in CH₂Cl₂ in an ice bath. TEA was added dropwise in the solution. The solution was stirred at the room temperature for 24 h and washed with 0.5 M HCl and saturated brine for three times. The organic phase was dried with anhydrous MgSO₄ overnight. The filtrate was concentrated and precipitated in cold anhydrous diethyl ether. The precipitate (mPEG-PCL-tosyl) was vacuum-dried at 40 °C.

Chrysin and K₂CO₃ were dissolved in DMF under nitrogen atmosphere. The mPEG-PCL-tosyl dissolved in DMF was added into the mixture solution with stirring. The reaction was stirred for 12 h at room temperature. The solution was filtrated and the filtrate was concentrated and precipitated in large amount of ethyl ether. The precipitate was dialyzed against deionized water using a 2000 MW cutoff tubing (MWCO 2000, Spectra/Por, USA). The mPEG-PCL-CHS was received after freeze-drying (yield =90%).

### Preparation of drug loaded micelles

The mPEG-PCL-CHS amphiphile (10 mg) and DOX (4.3 mg) were dissolved in 1 mL of DMSO with ultrasound for 0.5 h. The solution was dropped into 10 mL deionized water under strongly stirring overnight. The mixture was dialyzed against deionized water at 4 °C for 12 h in a dialysis membrane tube (Spectra/Por MWCO=5000). The solution was removed from the dialysis tube with centrifugation and lyophilized. The content of DOX was determined by fluorescence measurement (excitation at 485 nm) in DMSO using calibration curve obtained from DOX/DMSO solutions with different DOX concentrations. The whole procedure was processed in dark. The drug loading content (DLC) and encapsulation efficiency (EE) were calculated from the following formulae:

\[
DLC(\%) = \frac{[\text{weight of drug in micelle}]}{\text{weight of drug in feeding}} \times 100% \\
EE(\%) = \frac{[\text{weight of drug in micelle}]}{\text{weight of drug in feeding}} \times 100%
\]

### Release profile

DOX loaded micelles were dispersed in PBS (1 mL, ionic strength = 0.01 M, pH = 7.4). The release experiments were employed under sink conditions in order to ensure the good solubility of DOX. The mixture was put in dialysis membrane tubings (Spectra/Por MWCO = 1000). The tubings were immersed in vials containing 25 mL of phosphate buffered saline (PBS) solution and put in a shaking bed at 37 °C. 1 mL of PBS solution was taken out and the same volume PBS was added to the vials at prescribed time intervals. The released DOX was determined by a fluorescence detector with excitation wavelength at 485 nm and emission wavelength at 550 nm. 

### Cytotoxicity test

The cytotoxicity of blank micelles was tested by CCK8 assay. The cytotoxicity of blank micelles was tested by CCK8 assay. The cytotoxicity of blank micelles was tested by CCK8 assay.
against NIH 3T3 fibroblasts and HepG2 cells. The NIH 3T3 fibroblasts and HepG2 cells were separately inoculated into 96-well plates with \(5 \times 10^5\) cells per well in 100 \(\mu\)L of medium. After 24 h incubation, the medium was removed and replaced with 100 \(\mu\)L of medium containing different concentrations of blank micelles. The micelles were incubated with cells for 48 h. The medium was removed and the wells were washed with PBS (pH=7.4). 10 \(\mu\)L of 5 mg/mL CCK8 solution in PBS (pH=7.4) was added to each well. After incubated for 4 h, the medium containing unreacted CCK8 was removed carefully and the absorbance was measured.

**In vitro anticancer activity study**

HepG2 cells were separately inoculated into 96-well plates with \(5 \times 10^5\) cells per well in 100 \(\mu\)L of medium for 24 h. DOX·HCl and DOX loaded micelles solutions in DMEM were added to the plates and incubated for 48 h. The cell viability was measured by CCK8 assay.

**Cellular uptake**

The cellular uptake and distribution of drug loaded micelles in cells were detected using both flow cytometry and confocal laser scanning microscopy (CLSM). For CLSM studies, HepG2 cells were seeded on 35 mm diameter glass dishes at a cell density of \(1 \times 10^5\) cells per well and incubated for 24 h. HepG2 cells were incubated with DOX·HCl and DOX loaded micelles for 1 and 3 h (DOX concentration = 10 \(\mu\)g/mL), the cell nucleus were dyed by Hoechst. The cells were washed by PBS and taken photos under CLSM (Leica TC P SP5). DOX was excited at 480 nm with the emission at 590 nm. For flow cytometry studies, HepG2 cells were seeded on 6-well plates at a density of \(2 \times 10^5\) cells per well and incubated for 24 h. HepG2 cells were incubated with DOX·HCl and DOX loaded micelles for 1 and 3 h (DOX concentration = 10 \(\mu\)g/mL). The cells were digested with trypsinization and rinsed with PBS. The cells were resuspended in PBS after centrifugation (1000 rpm/min, 5 min) and measured for the fluorescence intensity (excitation: 480 nm; emission: 590 nm) on a BD FACS Calibur flow cytometer (Beckton Dickinson).

**Results and discussion**

We found that the introduction of \(\pi\)-conjugated moieties in polymeric micelles was favorable for the improvement of stability and drug loading content, and the chrysin modified mPEG-PCL micelles exhibited the optimal properties in anticancer drug delivery. Herein, PCL and mPEG fragments with various molecular weights were used to adjust the properties of the diblock copolymers with the immobilization of chrysin on the terminal group. Four diblock copolymers were fabricated micelles to deliver anticancer drug. The molecular weights of 2000 and 5000 to both mPEG fragments and PCL blocks were used to investigate the chain length effect on drug delivery.

The \(^1\)H NMR spectra of the four chrysin modified diblock copolymers were shown in Figure 1. The assignments of the protons from δ = 6.5 to 8 ppm were attributed to CHS (7, 8, 9, 10, 11, 12). The signals of 1 and 2 at the chemical shifts δ = 3.35 to 4.5 ppm were attributed to the protons in mPEG segments. The proton signal in PCL blocks was 3, 4, 5 and 6.

The molecular of the PCL block was calculated by comparing the integrals of characteristic peaks of the PCL block at δ= 2.25 ppm (triplet, -C (-O)-CH=) and mPEG block at δ = 3.35 ppm (singlet, -OCH3).

The molecular weights of the four diblock copolymers were also tested by GPC, only one peak was observed in all the GPC spectra (Figure 2), no unreacted mPEG was observed in the spectra. The calculated molecular weights of both \(^1\)H NMR and GPC results were summarized in Table 1. The Mns of mPEG-PCL-CHS copolymers calculated from \(^1\)H NMR spectra were comparable to the designed molecular weights. As the molecular weight tested by GPC was the relative molecular weight to the polystyrene standard samples, the Mns tested by GPC were deviated away from the theoretical values. It was interesting that the copolymers with mPEG5k blocks showed very narrow polydispersities. The PDIs of mPEG5k-PCL2k-CHS and mPEG5k-PCL5k-CHS copolymers were 1.06 and 1.05, respectively.
Table 1. The properties of mPEG-PCL-CHS micelles.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Molecular weight</th>
<th>Micelle diameter (nm)</th>
<th>DLC (%)</th>
<th>EE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mn&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Mn&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Mw/Mn</td>
<td>Blank</td>
</tr>
<tr>
<td>mPEG2k-PCL2k-CHS</td>
<td>4800</td>
<td>6700</td>
<td>1.14</td>
<td>15.5 ±1.1</td>
</tr>
<tr>
<td>mPEG2k-PCL5k-CHS</td>
<td>7500</td>
<td>12300</td>
<td>1.38</td>
<td>25.3 ± 2.8</td>
</tr>
<tr>
<td>mPEG5k-PCL2k-CHS</td>
<td>7100</td>
<td>10900</td>
<td>1.06</td>
<td>30.3 ± 2.0</td>
</tr>
<tr>
<td>mPEG5k-PCL5k-CHS</td>
<td>10200</td>
<td>14700</td>
<td>1.05</td>
<td>39.4 ± 1.6</td>
</tr>
</tbody>
</table>

<sup>a</sup> Calculated from <sup>1</sup>H NMR spectra.
<sup>b</sup> Tested by GPC.

The crystallization of the four diblock copolymers was investigated by DSC and XRD (Figure 3). Figure 3A presented clear melting endothermal peaks in the thermograms of mPEG-PCL-CHS block copolymers with different chain length. The melting temperatures (T<sub>ms</sub>) of mPEG2k-PCL2k-CHS, mPEG2k-PCL5k-CHS, mPEG5k-PCL2k-CHS and mPEG5k-PCL5k-CHS copolymers were 48.4, 43.0 (51.8), 54.7 and 55.5 °C, and the ΔHs were 106.0, 69.0, 114.5 and 105.8 J/g. The T<sub>ms</sub> of mPEG and PCL blocks were nearly the same around 50 °C, it was hard to observe the respective melting endothermal peaks of mPEG and PCL except mPEG2k-PCL5k-CHS. In the thermogram of mPEG2k-PCL5k-CHS, two melting points of 43.0 and 51.8 °C appeared, which were attributed to the blocks of mPEG2k and PCL5k, respectively. As we knew that the T<sub>m</sub> of semi-crystalline polymer commonly increased with increasing molecular weight. The discrepancy of T<sub>m</sub> between mPEG2k and PCL5k was large enough to be observed in the thermogram of mPEG2k-PCL5k-CHS. The crystals of mPEG and PCL blocks were mixed together in mPEG-PCL-CHS and they could not be separated in DSC results. XRD was an effective tool to discover the crystals of mPEG and PCL blocks. In the XRD spectra of the four copolymers (Figure 3B), the characteristic crystal peaks of PCL blocks at 2θ = 21.5° and 23.8° were obvious, mPEG showed two strong peaks at 2θ = 19.1 and 23.4°. The peaks at 23.4 and 23.8 degree were overlapped to exhibit a strong peak. The crystallinities calculated from XRD spectra were 72.3%, 60.3%, 76.0% and 70.3% for mPEG2k-PCL2k-CHS, mPEG2k-PCL5k-CHS, mPEG5k-PCL2k-CHS and mPEG5k-PCL5k-CHS copolymers, which was consist with the ΔHs in DSC results. The size and morphology of mPEG-PCL-CHS micelles were tested by DLS and TEM (Figure 4). The polymeric micelles were monodisperse in DLS result. TEM image of micelles showed the micelles were in spherical shape. The sizes of the four mPEG-PCL-CHS micelles tested by DLS were summarized in Table 1. The mean sizes of mPEG2k-PCL2k-CHS, mPEG2k-PCL5k-CHS, mPEG5k-PCL2k-CHS and mPEG5k-PCL5k-CHS micelles were 15.3, 25.3, 30.3 and 39.4 nanometers. Anticancer drug doxorubicin was encapsulated in the mPEG-PCL-CHS micelles, the mean sizes of the four drug loaded micelles were 56.6, 106.5, 60.4 and 61.3 nanometers, which were enlarged after drug encapsulation. Drug loading content and encapsulation efficiency were also measured, the DLCs of all the four micelles were higher than 20%. All the encapsulation efficiencies were higher than 85%. Within all the four species, mPEG2k-PCL5k-CHS micelle exhibited the highest DLC and EE, it possessed the largest mean size probably due to the highest DLC within all the drug loaded micelles. The mean diameters of both blank and drug loaded micelles in TEM images were calculated, the results were consistent with those of DLS results.

Figure 3. DSC (A) and XRD (B) spectra of mPEG-PCL-CHS block copolymers.
The interaction between DOX and mPEG-PCL-CHS micelles was investigated. The π-π stacking interaction within the drug loaded micelles was tested by fluorescence spectra and UV-vis absorption (Figure 5). All the four DOX/mPEG2k-PCL2k-CHS, DOX/mPEG2k-PCL5k-CHS, DOX/mPEG5k-PCL2k-CHS and DOX/mPEG5k-PCL5k-CHS micelles exhibited remarkable decrease in the fluorescence intensity of emission band with the same exciting wavelength of 485 nm and DOX concentration of 50 µg/mL, respectively. The significant quenching was the evidence of π-stacked DOX, which indicated that the DOX was packed into the micelles. The DOX/mPEG2k-PCL5k-CHS exhibited the greatest extent quenching and the minimum quenching appeared in DOX/mPEG5k-PCL2k-CHS. The more quenching extent in fluorescence revealed the stronger π-π stacking interaction within DOX loaded micelles. Red shift was observed in DOX loaded micelles (Figure 5b). The red shift of DOX was generally due to π-π stacking or ground-state electron donor-acceptor interaction between DOX and carriers. The drug loaded micelles with longer PCL chain length showed wider red shift.

mPEG-PCL micelles were extensively reported as carriers to deliver anticancer drugs, they were non-toxic to cells. In order to investigate the cytotoxicity of chrysin modified mPEG-PCL micelles, NIH/3T3 fibroblasts and HepG2 liver cancer cells were incubated with mPEG-PCL-CHS micelles with different concentrations. The cell viabilities of the two cell lines were presented in Figure 6, the cell viabilities of mPEG-PCL-CHS micelles to both NIH/3T3 and HepG2 cells were higher than 90% even the concentration of micelles was as high as 500 µg/mL, which was much higher than the concentration applied in vitro and in vivo. The results revealed that the four mPEG-PCL-CHS micelles were non-toxic to cells.

The in vitro release profiles of DOX loaded micelles was carried out in physiological condition (pH=7.4) at 37 °C. The burst release was found in all the four samples in the first 10 hours, however, the extent of burst release was different. The release rate decreased with the sequence of mPEG5k-PCL2k-CHS, mPEG5k-PCL5k-CHS, mPEG2k-PCL2k-CHS and mPEG2k-PCL5k-CHS. DOX/mPEGk-PCL5k-CHS exhibited the best profile in sustained release (Figure 7A). The drug release mechanism in these polymeric micelles was diffusion control. The drug diffused from the hydrophobic cores of micelles to the medium. The burst release was attributed to the drug absorbed in the hydrophilic PEG layer, which diffused easier and faster to the medium. Interestingly, the release rate sequence of drug loaded micelles was consistent with the sequence of fluorescence quenching in Figure 5A, it supported the conclusion that more extent quenching implied stronger π-π stacking interaction between DOX and micelles, which prevented the release of DOX from micelles and resulted in low drug release rate. The release profiles indicated that the sustained release of DOX was achieved...
and the drug release rate could be controlled by modulating the chain length of mPEG and PCL blocks.\textsuperscript{15}

The DOX loaded micelles were incubated with HepG2 cells to evaluate the in vitro anticancer activity (Figure 7B). The in vitro anticancer efficiency of DOX loaded micelles was dose-dependent. The IC\textsubscript{50}s (the concentration of anti-drug that killed 50% of cells) of free DOX·HCl, DOX/mPEG2k–PCL5k–CHS, DOX/mPEG2k–PCL5k–CHS, DOX/mPEG5k–PCL2k–CHS and DOX/mPEG5k–PCL2k–CHS micelles were 0.1, 0.25, 0.42, 0.65 and 2.04 µg/mL, respectively. As water-soluble anticancer drug, DOX·HCl diffused into cells and chelated into the DNA backbone to destroy DNA replication,\textsuperscript{25} the diffusion of DOX·HCl was faster in cellular internalization to the endocytosis of drug loaded micelles. The IC\textsubscript{50}s of the four drug loaded micelles were coincident with the release profile. Lower drug release rate was corresponded to lower IC\textsubscript{50}s, DOX/mPEG2k–PCL5k–CHS micelles exhibited the best anticancer efficiency.

![Figure 6. Cytotoxicity of mPEG1PCL1CHS micelles incubated with 3T3 fibroblasts (A) and HepG2 cancer cells (B).](image)

HepG2 cells without any treatment and with the treatment of blank micelles were used as the negative controls in flow cytometry test (Figure 8, F and G), there was only autofluorescence of the cells, which was not changed with time extension. The fluorescence intensity of DOX·HCl was the highest, and the following sequence was DOX/mPEG2k–PCL5k–CHS, DOX/mPEG2k–PCL2k–CHS, DOX/mPEG5k–PCL5k–CHS and DOX/mPEG5k–PCL2k–CHS. With longer incubation time (3 h), HepG2 cells treated with DOX/mPEG2k–PCL5k–CHS still showed the strongest fluorescence within all the four drug loaded micelles. The flow cytometry results were in agreement with the results of confocal fluorescence microscopy study.

![Figure 7. The release profiles (A) and inhibition effect of DOX loaded mPEG-PCL-CHS micelles to HepG2 cells.](image)

Figure 7. The release profiles (A) and inhibition effect of DOX loaded mPEG-PCL-CHS micelles to HepG2 cells.
length were synthesized via ring-opening polymerization of ε-CL with mPEG as macroinitiator, the terminal group of PCL segment was modified with chrysin to introduce π-π stacking interaction with anticancer drug doxorubicin. The chain length effects of chrysin modified mPEG-PCL diblock copolymers on the crystallization behavior of copolymers, self-assembly of micelles, drug loading, release profiles, cellular uptake and in vitro anticancer activity of drug loaded micelles were investigated. All the four micelles exhibited promising drug loading content higher than 20%, the mPEG2k-PCL5k-CHS micelles showed the highest drug loading content and encapsulation efficiency of 26.8% and 93%, respectively. The micelles were spherical with the size around 100 nanometers. The π-π stacking interaction between micelles and doxorubicin was evoked after drug encapsulation. The release rate decreased in the sequence of drug loaded mPEG5k-PCL2k-CHS, mPEG5k-PCL5k-CHS, mPEG2k-PCL2k-CHS and mPEG2k-PCL5k-CHS micelles. DOX/mPEGk-PCL5k-CHS exhibited the best profile in sustained release. The results of cellular uptake and IC_{50} revealed that the mPEG2k-PCL5k-CHS micelles exhibited the best in vitro anticancer activity.

Conclusions

Four amphiphilic mPEG-PCL copolymers with different chain lengths were synthesized via ring-opening polymerization of ε-CL with mPEG as macroinitiator, the terminal group of PCL segment was modified with chrysin to introduce π-π stacking interaction with anticancer drug doxorubicin. The chain length effects of chrysin modified mPEG-PCL diblock copolymers on the crystallization behavior of copolymers, self-assembly of micelles, drug loading, release profiles, cellular uptake and in vitro anticancer activity of drug loaded micelles were investigated. All the four micelles exhibited promising drug loading content higher than 20%, the mPEG2k-PCL5k-CHS micelles showed the highest drug loading content and encapsulation efficiency of 26.8% and 93%, respectively. The micelles were spherical with the size around 100 nanometers. The π-π stacking interaction between micelles and doxorubicin was evoked after drug encapsulation. The release rate decreased in the sequence of drug loaded mPEG5k-PCL2k-CHS, mPEG5k-PCL5k-CHS, mPEG2k-PCL2k-CHS and mPEG2k-PCL5k-CHS micelles. DOX/mPEGk-PCL5k-CHS exhibited the best profile in sustained release. The results of cellular uptake and IC_{50} revealed that the mPEG2k-PCL5k-CHS micelles exhibited the best in vitro anticancer activity.

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Notes and references


