Research Article

Enhancement of Oral Bioavailability of Puerarin by Polybutylcyanoacrylate Nanoparticles

Lixia Zhao,1,2 Anchang Liu,1,2 Min Sun,1,3 Jinsong Gu,4 Haigang Wang,2 Shuang Wang,1 Jing Zhang,1 Chenyu Guo,1 Rui Duan,1 and Guangxi Zhai1

1 School of Pharmaceutical Sciences, Shandong University, Jinan 250012, China
2 Department of Pharmacy, Qilu Hospital of Shandong University, Jinan 250012, China
3 Department of Pharmacy, Central Hospital of Zibo, Zibo 255036, China
4 Department of Biotechnology, College of Life Science and Technology, University of Jinan, Jinan 250022, China

Correspondence should be addressed to Guangxi Zhai, professorgxzhai@yahoo.cn

Received 14 June 2011; Accepted 6 July 2011

Abstract

The interest using novel drug delivery systems to improve oral bioavailability of drug with poor solubility is increasing. In this study, a new oral delivery system, polybutylcyanoacrylate nanoparticles (PBCNs), was introduced to improve the oral bioavailability of puerarin (PUE). PUE-loaded PBCN was successfully prepared by anionic polymerization method. Characterization of PUE-loaded PBCN was evaluated with morphology, size, zeta potential, and in vitro release study. The PBCN loading PUE exhibited a spherical shape under transmission electron microscopy with an average size of 159.4 nm, and the zeta potential was $-15.0$ mV. The in vitro release of PUE-loaded PBCN showed an initial burst release followed by a sustained release. Physicochemical state of PUE in PBCN was investigated by differential scanning colorimetry, X-ray diffraction, and Fourier transform infrared spectroscopy. The results indicated that PUE in PBCN was in a noncrystalline state. The oral pharmacokinetic study in rats showed that the relative bioavailability of PUE-encapsulated PBCN to the crude PUE was more than 550%. It can be concluded that PBCN as an oral drug carrier can significantly improve the oral bioavailability of PUE.

1. Introduction

The bioavailability is an important parameter showing the degree and rate of drug molecules entering blood circulation, indicating the effectiveness and safety of an extravascular administration formulation. It can be influenced by drug formulations, food, and physiological factors. However, up to 50% of orally administered drugs present formulation problems related to poor solubility (0.011 M) [1]. In recent years, many specific pharmaceutical approaches such as micro- and nanotechnology have been developed to improve the bioavailability [2]. As one promising delivery system with improved bioavailability, polybutylcyanoacrylate nanoparticles (PBCNs) have attracted considerable attention [3]. It is a type of solid colloidal particles ranging in size from 10 nm to 1000 nm made of biodegradable polymers [4, 5] which could significantly enhance the oral absorption of some drugs such as thymopentin [6]. The PBCNs that served as the oral carriers can prevent the destruction of drugs (e.g., peptide drugs) by the acid and enzymes in gastrointestinal tract [7, 8], improve the drug absorption through Peyer’s patches (PP), the immunization-related tissue in small intestine of human and animals which accounts for about 25% of the intestinal mucosa, and other intestinal lymphoid tissue into the blood circulation [9], and prolong the residence time in vivo because of the small particles and bioadhesive property [6].

Puerarin (PUE, Figure 1) is the naturally occurring isoflavone C-glycoside extracted from the roots of *Pueraria lobata* (Willd.), *Ohwi*, and *P. thomsonii* Benth, which is widely used in China for the treatment of cerebrovascular and cardiovascular disease. Although PUE has definite therapeutic effectiveness, its clinical application is limited by its poor solubility, short elimination half-life, and poor oral bioavailability. The commercial available preparation for PUE is its injection which has poor patient compliance...
because of the high frequency of injecting administration [10]. The oral preparation containing *Pueraria lobata* extraction is Yufengningxin tablet which needs to be orally given five tablets each time and three times each day. Therefore, the development of a suitable oral preparation is a research focus of PUE. In the past few years, several pharmaceutical techniques were adopted to improve the oral absorption of puerarin, such as microemulsion [10–12] and solid lipid nanoparticles [13]. However, up to date no study on PBCN for enhancing the oral bioavailability of PUE has been reported.

In this study, PUE-loaded PBCN (PUE-PBCN) was prepared by anionic polymerization method in order to improve the oral bioavailability of PUE and was characterized with morphology, size, zeta potential, and in vitro release study. Physicochemical state of PUE in PBCN was investigated by differential scanning colorimetry (DSC), X-ray diffraction (XRD), and Fourier transform infrared (FTIR) spectroscopy. Additionally, the pharmacokinetics of PUE-loaded PBCN was studied in rats.

2. Materials and Methods

2.1. Materials. PUE was purchased from Nanjing Zelang Medical Technology Co., Ltd. (Nanjing, China). Dextran 70 (D-(+) glucose) and poloxamer 188 (F68) were provided by the Shanghai Treechem Biotech Co., Ltd. (Shanghai, China). N-Butylcyanoacrylate (BCA) was a gift sample from Beijing Suncon Medical Adhesive Co., Ltd. (Beijing, China). All other reagents used in the study were of analytical regent grade.

2.2. Preparation of PBCN. Based on the previous reports with some modifications [14–16], PUE-PBCN was prepared using open anionic polymerization of n-butyl-2-cyanoacrylate (BCA) monomer in acidic medium (0.01 M HCl), containing 0.5% (w/v) of dextran 70 and 0.5% (w/v) of poloxamer 188 as stabilizer and emulsifier. The monomer BCA was injected drop by drop into the stirred medium including PUE (the weight ratio of drug to polymer was 1 : 2). Polymerization was conducted for 4 hours under magnetic agitation at 400 rpm and room temperature. Subsequently, sodium hydroxide solution (1 M) was used to adjust the pH of resulting suspension to 6.8 ± 0.1. The above mixture was stirred for an additional 1 h at the same conditions, and the reaction was gradually completed. The PUE-PBCN suspension was obtained after the above system was filtered through 0.8 μm filter membrane.

2.3. Characterization of PBCN

2.3.1. Morphology. PUE-loaded PBCN was observed under a transmission electron microscope (TEM, JEM-1200EX, JEOL, Tokyo, Japan) using the negative-staining method. One drop of diluted PUE-PBCN was added to a copper grid to form a thin liquid film, and then the film was negatively stained by adding 2% (w/v) phosphotungstic acid (PTA, pH 7.0). Extra droplet was instantly removed with filter paper, and then the grid was dried at room temperature as a TEM sample. The film was examined under TEM and photographed.
2.3.2. Particle Size and Zeta Potential. The particle size of PUE-PBCN was analyzed using a particle sizer (Zetasizer 3000 HAS, Malvern Instruments Ltd., Malvern, Worcestershire, UK) with photon correlation spectroscopy (PCS) at a fixed angle of 90° at a temperature of 25°C. It was conducted with He-Ne laser of 3 mW at a wavelength of 633 nm, and the particle size analysis data were evaluated using the volume distribution.

Zeta potential of PUE-PBCN was determined using TV microscopic electrophoresis system (DXD-II, Optics Co., Ltd., Jiangsu, China) at room temperature.

2.3.3. Encapsulation Efficiency (EE) and Drug Loading (DL). The measurement for encapsulation efficiency and drug loading of PUE-PBCN was carried out with centrifugation ultrafiltration method according to the previous reports [17, 18]. The free PUE was separated from PUE-PBCN by centrifugal filter tubes (Amicon Ultra-4, Millipore, Ireland) with a molecular cut off of 10 kDa. Briefly, 2 mL of PUE-loaded nanoparticle suspension was added into centrifugal filter tube and centrifuged at 4000 rpm for 20 min at room temperature using a centrifuge (Biofuge primo R, Heraeus, Hanau, Germany). The initial total amount of PUE in the suspension system and the amount of free drug in the filtrate were measured using UV at the wavelength of 251 nm. The encapsulation efficiency and drug loading in the PBCN were calculated as, respectively follows:

\[
EE\% = \frac{W_{\text{total drug}} - W_{\text{free drug}}}{W_{\text{total drug}}} \times 100\%, \quad (1)
\]

\[
DL\% = \frac{W_{\text{total drug}} - W_{\text{free drug}}}{W_{\text{polymer}}} \times 100\%, \quad (2)
\]

where “W_{free drug}” is the amount of PUE unloaded in PBCA nanoparticles, “W_{total drug}” is the initial total amount of PUE in the suspension system, and “W_{polymer}” is the weight of butyl-cyanoacrylate monomer.

2.3.4. DSC and XRD Analysis. The physical state of the drug entrapped in the PBCN was characterized using a differential scanning calorimeter (CDR-4P, Shanghai Tianping Instrument Ltd., Shanghai, China) and an X-ray diffractometer (D/max r-B, Rigaku Co., Tokyo, Japan). Prior to analysis, PUE-PBCN and blank PBCN used for DSC and XRD were obtained by freeze-drying without any freeze-dried protectants.

For DSC measurement, about 10 mg of samples (PUE, PUE-PBCN, or blank PBCN) were sealed in the aluminum pan and heated at a scanning rate of 10°C/min from 30 to 400°C under dry nitrogen atmosphere at a flow rate of 0.2 mL/min.

X-ray diffraction patterns were determined for PUE-PBCN, blank PBCN, and pure PUE with a Cu line as the source of radiation. A radiation at 40 kV voltage and 40 mA current was used, and diffractograms were performed with a scanning rate of 2°/min over a 2θ range of from 6° to 40°.

2.3.5. Fourier Transform Infrared (FTIR) Spectroscopy. FTIR analysis was performed to provide further information on the drug-polymer relationship using an FTIR spectrometer (Thermo Electron Scientific Instruments Corp.). FTIR spectra of PUE or BCA monomer or PUE-PBCN (with or without drug) were recorded in KBr pellets or KBr cell on an FTIR spectrometer with resolution of 2 cm⁻¹. A total of 64 scans were used and data were recorded over the range 4000–400 cm⁻¹.

2.3.6. In Vitro Release Study. The in vitro release of PUE from the nanoparticles was studied by dialysis against phosphate buffer solutions (PBS) with 12–14 kDa molecular cutoff bag pH 7.4 [19]. PUE-PBCN suspension or PUE propylene glycol solution (containing 3 mg/mL of PUE) was placed into dialysis bags, respectively. Then the bags were suspended in flasks containing 150 mL of PBS as dissolution medium at 37°C in shaking water bath at 100 rpm. 1 mL of dissolution medium was withdrawn at regular time intervals, and the same volume was added with fresh release medium. The concentrations of PUE in dissolution medium were measured using UV spectrophotometer at 251 nm. All experiments were performed in triplicates.

2.4. Pharmacokinetics Study. The study on the pharmacokinetics of PUE-PBCN was performed in male Wistar rats (200 ± 20 g) supplied by the Medical Animal Test Center of Shandong University [20]. All animal experiments complied with the requirements of the National Act on the use of experimental animals (China). The animals were divided randomly into 2 groups (n = 5), housed in an environmentally controlled breeding room (temperature 25 ± 2°C, humidity 60 ± 5%, 12 h dark/light cycle) for 7 days and fasted overnight before experiment with free access to water. Group 1 was orally administrated PUE suspension (30 mg/kg, PUE dispersed in 0.5% sodium carboxymethylcel lulose (CMC-Na) solution), and Group 2 was given PUE-PBCN at the same dose of 30 mg/kg through the same route. 0.3 mL of blood was withdrawn from the subclavian vein at 0 min, 5 min, 10 min, 20 min, 40 min, 1 h, 2 h, 4 h, 6 h, 8 h, 12 h, and 24 h. The samples were placed into heparinized tubes and isolated immediately by centrifugation at 4000 rpm for 10 min. The plasma obtained was stored at −20°C before analysis.

2.5. Liquid Chromatography Tandem Mass Spectrometry (LC-MS) Analysis. Extraction of PUE from plasma was conducted as follows: 25 μL of internal standard solution (genistein, 1 μg/mL) and 75 μL of methanol were added to 25 μL of plasma and the resulting mixture was vortexed for 1 min and then centrifuged at 11,000 rpm for 5 min to obtain a clean supernatant. An aliquot (20 μL) of supernatant was injected into the LC-MS for analysis [21].

PUE in plasma was determined by LC-MS analysis using an Agilent 1200 system equipped with an autosampler,
a vacuum degasser unit, and a binary pump. The HPLC system was coupled to an Agilent G6410B triple-quadrupole mass spectrometer (Agilent Technologies, USA), equipped with electrospray ionization (ESI) run by Agilent Mass Hunter Workstation B.01.03.

Mass spectrometric analysis was performed in multiple reaction monitoring (MRM) mode by monitoring ion transitions at m/z 415.1 → 295.1 for PUE and 269.1 → 133.0 for genistein (IS), with spray gas pressure of 350 Pa, protective air of nitrogen gas at a flow rate of 10 L/min, dwell time of 200 ms, capillary voltage of 4000 V, fragment electric voltage of 150 V for PUE and 135 V for genistein, and collision energy of 20 eV for PUE and 25 eV for genistein. Chromatographic peaks of analyte and IS were identified on the basis of retention time and MRM response.

Chromatographic separation was performed using a ProntoSIL C18 column (150 mm × 3 mm, 3 μm, Bischoff, Germany). The mobile phase consisted of a mixture of 10 mM ammonium acetate in water containing 0.1% formic acid and methanol (20:80, v/v). The separation was performed under isocratic conditions with a constant flow rate of 0.6 mL/min. The analytical method was validated according to “Guidance for Industry-Bioanalytical Method Validation” [22]. Pharmacokinetic parameters were evaluated using Pharmacokinetic program DAS 2.0 (supplied by Chinese Pharmacological Society).

2.6. Statistical Analysis. The statistical analysis of the data was carried out using Student’s t-test with P < 0.05 as the minimal level of significance.

3. Results and Discussion

3.1. Fabrication of PUE-PBCN. PUE-PBCN was prepared by anionic polymerization based on the reported method [3]. Based on the reported mechanism of polymerization [3, 23], the PUE-PBCN may fabricate as follows (Figure 2). Firstly, the surfactant such as F68 dissolved in acidic media above the critical micelle concentration (CMC) and the monomer-swollen micelles were formed as micelles took up BCA monomer. Secondly, the anionic radicals present in the media entered the swollen structures to initiate the polymerization, resulting in the formation of the primary particles. The newly formed polymer particles continued to grow in size by absorbing monomer molecules until the monomer completely disappeared. These primary particles were stabilized by the surfactant (F68) and stabilizer (Dextran 70), which prevented the aggregation of formed particles by steric repulsion. Concurrently, PUE, a high hydrophobic drug, might be gradually dispersed in the hydrophobic part of the swollen micelles and further incorporated in polymer nanoparticles by emulsion during the polymerization. The mean encapsulation efficiency and drug loading of PUE-PBCN suspension were 48.75% and 15.02%, respectively.

3.2. Characterization of PUE-PBCN. As shown in Figure 3, PUE-PBCN was spherical in shape with a smooth surface under TEM, which was in consistent with the reported result of PBCN loading curcuminoids [3].

It can be seen from Figure 4 that more than 85% of PUE-PBCN was between 100 nm to 300 nm, indicating that the size range of PBCN was narrow. The mean diameter was 159.4 nm with polydispersity index (PDI) of 0.139.

In the present study, the mean zeta potential of PUE-PBCN was −15.00 mV, which was similar to the report of Rivastigmine-loaded PBCN [5].

3.3. DSC and XRD Analysis. DSC, characterizing the thermal behavior of polymer and drug correlated to their melting and recrystallization, was used to investigate the physicochemical
state of PUE in the formulation [24]. The DSC thermograms of pure PUE, blank PBCN, and PUE-PBCN were shown in Figure 5. The wide endothermic peak at 112.2°C present in the thermogram of the crude PUE was not visible in that of PUE-PBCN, implying that PUE in PBCN was not in crystalline state but in an amorphous form. Blank PBCN showed an endothermic peak about 55°C, while the peak disappeared in the thermogram of PUE-PBCN, which might be due to drug interfering in the heat flow [25].

XRD analysis was employed to study the potential changes of the crystalline state of PUE in PBCN. The XRD patterns for PUE-PBCN, blank PBCN, and pure PUE were shown in Figure 6. Compared with the diffractograms for blank PBCN and pure PUE, almost all diffraction peaks disappeared in that of PUE-PBCN, indicating that PUE was in an amorphous form.

3.4. FTIR Analysis. FTIR analysis was proposed to evaluate possible interactions between drug and polymer carrier. Figure 7 showed the infrared spectra of BCA monomer (a), blank PBCN (b), PUE-PBCN (c), and pure PUE (d). The peak at about 2200 cm$^{-1}$ in spectra of BCA is characteristic of C≡N (stretching mode of the polymer). The C≡N peak was also present in the spectra of blank PBCN and PUE-PBCN, indicating that C≡N did not participate in the polymerization. Broader bands at about 3400 cm$^{-1}$ were observed in the spectra of blank PBCN, PUE-PBCN, and pure PUE, which were corresponding to hydroxyl radicals. Compared to the spectrum of BCA monomer, these bands in spectra of PBCN may be a result of polymerization, which can be inferred from the chemical structures of BCA and PBCA (Figures 1(b) and 1(c)) [25]. Moreover, the fingerprint region at 1600 cm$^{-1}$ and 650 cm$^{-1}$ assigned to the absorption peak of benzene did not change or shift and only got weak in the spectrum of PUE-PBCN, which could be a strong evidence showing no new chemical bond produced. Similar results were obtained in the previous report [26].

3.5. In Vitro Release Study. The in vitro release of PUE from PBCN was conformed in phosphate buffer at pH 7.4 with dynamic dialysis method. The results of drug release from PUE solution or PBCN suspension at pH 7.4 were shown in Figure 8. The PUE release from solution was found to
be much faster, nearly 100% in 5 h. In contrast, the release profile of PUE from PBCN demonstrated two phases: an initial burst release followed by a sustained release, only about 75% over the period of 24 h. This could be attributed to the fact that PUE loaded on or near the surface of PBCN was released first, and subsequently the drug embedded in the nanoparticles was released, which might release slowly by diffusion from the matrix or along with the degradation of polymer [27]. The result was in agreement with the previous reports that drug-loaded PBCN provided a controlled release pattern [28].

3.6. Pharmacokinetics Study. The mean concentrations of PUE in the plasma after oral administration of a single dose in rats were measured using a highly sensitive and specific LC/MS/MS method. The retention time of puerarin and genistein was 1.4 min and 1.8 min, respectively. The mean recovery of PUE was more than 93.0%, and the precision for interday and intraday was less than 15%. Besides, PUE in analyzed samples was stable for 24 h at room temperature, 30 days at −20°C and three freeze-thaw cycles.

The calibration curve was linear over the concentration range of 2–10000 ng/mL for PUE, and the regression equation was

\[
A = 8.01 \times 10^{-5}C + 2.35 \times 10^{-4}
\]

with the mean correlation coefficient (r) of 0.9996.

The data for pharmacokinetics in rats were analyzed with DAS 2.0, and pharmacokinetic parameters were shown in Table 1. The pharmacokinetic behaviors of PUE suspension and PUE-PBCN were described using noncompartment model. As shown in Table 1, the area under the concentration-time curve (AUC) of PUE-PBCN was 6.765 ± 2.374 mg/L·h, which was 5.56-fold greater than that of PUE suspension administration (1.216 ± 0.158 mg/L·h, \( P < 0.01 \)). The mean residence time (MRT) of PUE-PBCN (6.299 ± 0.925 h) and half-life \( T_{1/2} \) (3.937 ± 0.972) were 1.63 times those of CUR suspension (3.864 ± 0.832 h for MRT, 2.409 ± 0.791 h for \( T_{1/2} \)), while the plasma clearance (4.936 ± 1.810 L/h·kg) of CUR-PBCN was much lower relative to that of the control suspension (25.04 ± 3.489 L/h·kg). These results might be related to the small size of nanoparticles and coating of F68 on the surface to keep the nanoparticles contacting closely and long with the gastrointestinal (GI) tract [29]. The mean plasma concentration-time profiles for PUE and PUE-PBCN were presented in Figure 9. At all time points, the PUE concentrations in plasma were significantly higher for rats treated with PUE-PBCN than those treated with PUE suspension. Twenty-four hours after oral administration of PUE-PBCN, the PUE concentration in plasma was still more than 0.025 mg/L, while it was undetectable after 12 h for PUE suspension. Double peaks in the profile of PUE-PBCN may be contributed to, on the one hand, the possible enterohepatic recirculation; on the other hand, the absorption time lag after the first absorption existing due to the more time for PUE was needed to be released from PBCN and absorbed [30]. The similar phenomenon has been in agreement with the pharmacokinetic study of oral administration of PUE load SLNs [13]. These results indicated that PUE entrapped into PBCN led to the increased absorption of PUE by oral administration.
of PUE by 5.5-fold. It can be concluded that PBCN can improve the oral bioavailability. The drug-loaded PBCN was directly uptaken by the mucus and microfold cells (M cells) of the Peyer's patches and the lymphatics through the GI tract [31, 32]. In addition, PUE could be protected by incorporation into PBCN during the absorption process [33]. The high affinity of PBCN to M cells in the GI tract and increased permeability by surfactants could result in long MRT and thus leading to the enhanced bioavailability [34, 35]. The results support that PBCN is a promising delivery system for improving oral bioavailability of PUE.

**4. Conclusions**

In the present study, PUE-PBCN was successfully prepared to improve the bioavailability. The drug-loaded PBCN was sphere-like shape with a mean diameter of 159.4 nm and the zeta potential was $-15.00 \text{ mV}$. It could be concluded from DSC, FTIR, and X-ray analysis that PUE loaded in nanoparticles was in an amorphous form. The pharmacokinetic data showed that the PBCN could improve the oral bioavailability of PUE by 5.5-fold. It can be concluded that PBCN can improve the oral absorption of PUE.

**Acknowledgments**

This work was supported by the Natural Science Foundation of Shandong Province, China (Grant no. Q2007C13), the Shandong Administration of Traditional Chinese Medicine (Grant no. 2009-152), and the Science and Technology Research Project From Population and Family Planning Commission of Shandong Province, China. L. Zhao, A. Liu and M. Sun contributed equally to the work.

**References**


**Table 1: Main pharmacokinetic parameters of PUE suspension and PUE-PBCN in rats.**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Unit</th>
<th>Value (mean ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$AUC_{(0-\infty)}$</td>
<td>mg/L·h</td>
<td>1.216 ± 0.158</td>
</tr>
<tr>
<td>$MRT_{(0-\infty)}$</td>
<td>h</td>
<td>3.864 ± 0.832</td>
</tr>
<tr>
<td>$T_{1/2}$</td>
<td>h</td>
<td>2.409 ± 0.791</td>
</tr>
<tr>
<td>$V_z$</td>
<td>L/kg</td>
<td>85.41 ± 21.63</td>
</tr>
<tr>
<td>$CL$</td>
<td>L/h·kg</td>
<td>25.04 ± 3.489</td>
</tr>
<tr>
<td>$C_{max}$</td>
<td>mg/L</td>
<td>0.363 ± 0.045</td>
</tr>
</tbody>
</table>

$^{\ast\ast}$ Comparing to PUE suspension group, $P < 0.001$; $^\ast$ Comparing to PUE suspension group, $P < 0.05$. 

The relative bioavailability ($F_{rel}$) was calculated as

$$F_{rel} = \frac{AUC_{PUE-PBCN}}{AUC_{PUE}} \times 100\%.$$ (3)

The result indicated that PBCN could significantly enhance the oral bioavailability of PUE (more than 5.5-fold that of PUE suspension). The enhanced bioavailability by the PBCN formulation might be a cowork of direct uptake of nanoparticles through the GI tract, increased permeability by surfactants, and decreased degradation and clearance for PUE. The particles in nanorange could be directly uptaken by the mucus and microfold cells (M cells) of the Peyer's patches and the lymphatics through the GI tract [31, 32]. In addition, PUE could be protected by incorporation into PBCN during the absorption process [33]. The high affinity of PBCN to M cells in the GI tract and increased permeability by surfactants could result in long MRT and $T_{1/2}$, thus leading to the enhanced bioavailability [34, 35]. The results support that PBCN is a promising delivery system for improving oral bioavailability of PUE.


