Co-Delivery of Prednisolone and Curcumin in Human Serum Albumin Nanoparticles for Effective Treatment of Rheumatoid Arthritis

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Background: Prednisolone (PD) is extremely effective for treating rheumatoid arthritis (RA). However, it distributes nonspecifically throughout the body and its use is associated with serious side effects, which prompted us to compound it into a phytomedicine for greater efficacy and safety.

Methods: We combined PD with curcumin (CU), an effective monomer from traditional Chinese medicine, and human serum albumin (HSA) in a nanoparticulate system (N-PD/CU) to compensate for the poor bioavailability of PD and CU. N-PD/CU was prepared by high-pressure homogenization, and its characteristics were evaluated in vitro. Next, we investigated its toxicity and mechanism of anti-inflammatory to macrophages. Finally, its pharmacokinetics, biodistribution and therapeutic efficacy were assessed in rats with adjuvant-induced arthritis (AIA).

Results: N-PD/CU showed a narrow size distribution around 150.4 ± 2.4 nm, a polydispersity index of 0.22 ± 0.02 and drug loading efficiency (DLE) of 88.75 ± 1.82% for PD and 85.79 ± 1.43% for CU. N-PD/CU showed sustained release of both drugs in vitro. N-PD/CU had no toxicity to macrophages in vitro on concentrations between 0.1 and 1.2 μmol/mL. In activated macrophages, N-PD decreased levels of pro-inflammatory cytokines, while N-CU increased levels of anti-inflammatory IL-10, and N-PD/CU exhibited best therapeutic effect in vitro, suggesting co-delivery of PD and CU may synergistically control the course of RA. In AIA rats, N-PD/CU accumulated in inflamed joints through the effect of extravasation through leaky vasculature and subsequent inflammatory cell-mediated sequestration (ELVIS effect) in inflammatory lesion and showed higher therapeutic efficacy than single-loaded nanoparticles, either free drug on its own, or a simple mixture of the two drugs.

Conclusion: This codeelivery system based on HSA is a promising platform for combination chemotherapy in RA.

Keywords: co-delivery, prednisolone, curcumin, human serum albumin, rheumatoid arthritis

Introduction
Rheumatoid arthritis (RA) is a chronic systemic autoimmune disease of unknown etiology. It is characterized by symmetric, invasive inflammation in multiple joints of the hands and feet, causing pain and restricting movement.1 As the disease progresses, the risk of bone damage and cartilage destruction increases, leading to substantial disability.2,3

Pro-inflammatory cytokines such as TNF-α, IL-1β and IL-6 drive progression of RA.4,5 IL-10, on the contrary, is an anti-inflammatory cytokine that can mitigate inflammation and immune responses in the disease.6 An effective therapy against
the disease is glucocorticoids (GCs) such as prednisolone (PD), which can inhibit secretion of pro-inflammatory cytokines. PD, like most GCs, shows poor bioavailability because of its nonspecific distribution throughout the body, which increases the risk of adverse effects.

This has motivated interest in using botanicals as adjunct therapy to prevent the development of arthritis and other chronic diseases. For example, the hydrophobic polyphenol curcumin (CU), from the rhizome of the herb Curcuma longa L., shows remarkable anti-inflammatory properties, and has shown efficacy against various diseases, including inflammatory disorders. CU can attenuate RA by promoting the production of IL-10, which itself further reduces production of pro-inflammatory cytokines. However, like PD, CU is hydrophobic, which limits their clinical application and their inability to target inflammatory lesion, cause adverse effects after administration. It may be that combining PD and CU can improve overall efficacy against RA, but this likely requires co-delivering them using a targeted system that can ensure that they act specifically in inflamed tissues, thereby reducing risk of adverse effects.

An excellent nanocarrier candidate for these drugs is human serum albumin (HSA), which is nontoxic and biocompatible, and has high loading capacity for various drugs, allowing delivery of higher doses while keeping adverse events tolerable. In the present study, we examined whether we could deliver PD and CU effectively using HSA nanoparticles to inflamed joints in rats with adjuvant-induced arthritis (AIA). Our goal was to target the drugs to inflamed joints via the recently described phenomenon of extravasation through leaky vasculature and subsequent inflammatory cell-mediated sequestration (ELVIS effect). The ELVIS effect is, analogous to the classical enhanced permeability and retention (EPR) effect in solid tumors.

We show here that HSA-based nanoparticles loaded with both PD and CU can be prepared by high-pressure homogenization and that in AIA rats, they accumulate specifically in inflamed joints and show greater anti-inflammatory efficacy than the free drugs together or on their own.

Materials and Methods

Materials

Prednisolone (purity ≥ 99%) was supplied by Chengdu Aike Pharmaceutical (Chengdu, China); curcumin (purity ≥ 99%), oleic acid (OA) and purified yolk lecithin (lipoid E80), Lipoid Co., Ltd (Ludwigshafen, Germany); HSA and soybean oil, Beijing Biotopped Science & Technology (Beijing, China);Incomplete Freund’s adjuvant (10mg/mL), Chondrex (Washington DC, USA). 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) and Dioctadecyl tetramethyl Indodicarbocyanine (DID) were supplied by Solarbio Science &Technology Co, Ltd (Beijing, China). Enzyme-linked immunosorbent assay (ELISA) kits to assay levels of TNF-α, IL-1β, IL-6 and IL-10 in cell supernatant were purchased from Thermo Fisher (Vienna, Austria). ELISA kits to assay levels of TNF-α, IL-1β, IL-6 and IL-10 in serum of rats were obtained from Bioscience (San Diego, CA, USA). All other chemical reagents and solvents were of analytic grade or above.

Cells and Animals

RAW264.7 cells (Chinese Academy of Sciences, Shanghai, China) were cultured in RPMI 1640 medium (Hyclone Laboratories, Omaha, NE, USA) and supplemented with 10% (v/v) fetal bovine serum (Hyclone) and 1% (v/v) penicillin/streptomycin (Solarbio Science & Technology, Beijing, China). Cells were incubated in a 5% CO₂ environment at 37°C.

Male Sprague-Dawley rats (200 ± 20 g) were supplied by the Laboratory Animal Center of Southwest Medical University (Luzhou, China). All experimental rats were housed in common animal rooms and fed with standard feed, with indoor temperature 25 ± 2°C and relative wet degree 50 ± 10%. Animal experiments were approved by the Animal Ethics Committee of Southwest Medical University (Permit No. 20160126). During the whole animal experiments, laboratory animal guideline for ethical review of animal welfare for the animal was followed by according to National Standard of China.

Preparation of Drug-Loaded Nanoparticles

The lipophilicity of PD and CU was enhanced by forming PD-OA and CU-OA as described. Briefly, PD (12 mg) and CU (6 mg) were dissolved in acetone (6 mL), to which was added 75 μL OA (10%, w/v). The system was mixed well and stirred up for one hour at room temperature, then acetone was removed by vacuum rotary evaporation. Prepared PD-OA and CU-OA, E80 (90 mg), and cholesterol (90 mg) were dissolved in chloroform and mixed together, then HSA
solution at a concentration of 1.3% (w/v) was added. Then the mixture was subjected to high-speed shearing for 1 min (Bio-Homogenizer, Biospec Products, USA), followed by homogenization into a emulsion at 18,000 psi for 10 cycles (AH 100D; ATS Engineering, Canada). Finally, the organic solvent was removed by vacuum rotary evaporation. The resulting nanoparticles, loaded with both drugs, were referred to as N-PD/CU. The same method was used to prepare blank nanoparticles without either drug, as well as nanoparticles with each drug on its own (N-PD, N-CU).

Characteristics of Drug-Loaded Nanoparticles

The size and zeta potential of drug-loaded HSA nanoparticles were measured at room temperature using a Malvern Zetasizer Nano ZS90 (Malvern Instruments, Malvern, UK), using equal volumes of nanoparticles suspended in ultrapure water. Samples of nanoparticles were also analyzed by transmission electron microscopy (H-600, Hitachi, Japan). Samples were diluted in ultrapure water and stained with phosphotungstic acid for approximately 30 s. Encapsulation efficiency (EE) and drug loading efficiency (DLE) were measured by HPLC.

Drug Release from HSA Nanoparticles in vitro

Dialysis was used to analyze the release of PD and CU from N-PD/CU. Dialysis bags (molecular weight cut-off, 10 kDa) were filled with 1.5 mL of N-PD/CU or the equivalent amounts of either free drug, and the bags were immersed in 10 mL of phosphate-buffered saline (PBS) containing 0.2% (w/v) Tween 80 with stirring at 100 rpm at 37°C. At fixed times, a sample of release medium (300 µL) was extracted and the same volume of fresh medium was added back to keep the final volume constant. Concentrations of each drug in the release medium were determined by HPLC (Agilent 1260, USA).

Cytotoxicity of Nanoparticles Toward RAW264.7 Cells in vitro

RAW264.7 cells were seeded into 96-well plates at a density of 1×10^4 per well and incubated in 5% CO₂ at 37°C. After 24 h, the original medium was replaced with 200 µL of solutions containing various concentrations of blank nanoparticles, N-PD/CU, single-loaded nanoparticles, each free drug on its own, or a simple mixture of both free drugs. Cells were incubated for another 48 h, then the numbers of living cells were determined using the MTT assay. Relative cell viability was calculated based on the absorption of treated wells relative to that of control wells incubated with drug-free medium.

Pharmacodynamics in vitro

RAW264.7 cells in logarithmic growth phase were digested into a single-cell suspension, inoculated into 6-well plates (5×10^4 cells per well), then cultured for 24 h. Cells were incubated in a 5% CO₂ atmosphere at 37°C. Then LPS was added to a final concentration of 100 ng/mL and cultured for 24 h to activate the cells. The medium was replaced with 2 mL of fresh medium containing free PD, free CU, N-PD, N-CU, N-PD/CU, or the simple combination of PD and CU (total drug concentration in all cases was 200 µg/mL). After incubation for 48 h, the medium was collected, centrifuged at 2000 rpm for 4 min, and the supernatant was assayed for the inflammatory cytokines TNF-α, IL-6, IL-1β and IL-10, using ELISA kits according to the manufacturer’s protocols.

Pharmacokinetics and Biodistribution of Nanoparticles

Fifty male Sprague-Dawley rats (200 ± 20 g) were used to establish a rat model of AIA. Heat-killed mycobacteria were suspended in incomplete Freund’s adjuvant (10 mg/mL), then each rat was injected subcutaneously with 100 µL at the base of the tail. Every day, we evaluated the articular index (AI) score for the paws of each rat as described. After 14 days, we chose the AIA rats with an AI score >4 points to study the in vivo pharmacokinetics and distribution. The selected rats were fasted and randomly assigned to 10 groups (n=5). Five groups received a single injection of N-PD/CU containing 5.32 mg/kg PD and 2.55 mg/kg CU, while the other five groups received a single injection of a simple mixture of PD and CU at the same concentrations as in the nanoparticles. Blood samples were collected at 0.25, 0.5, 1, 2 and 4 h, and concentrations of each drug in serum were determined by HPLC. Meanwhile, various tissues were collected (heart, liver, spleen, lung, kidney and joints), homogenized in a volume of 0.9% NaCl equal to twice the tissue weight, and assayed for each drug by HPLC.

Distribution of N-PD/CU in Arthritic Rats

Nine AIA rats were depilated over the entire body using cream, and then given a tail vein injection of saline, free DID or DID/N-PD/CU (5 µg DID per rat). At 4, 12, and 24
h, animals were anesthetized with 10% chloral hydrate and imaged using a real-time fluorescence imaging system (Fx Pro/FX, Bruker, USA).

**Therapeutic Efficacy of Nanoparticles in Arthritic Rats**

On days 14, 16, 18, 20 and 22 after arthritis induction, rats in each group (n=3) were injected intravenously with N-PD/CU, N-PD, N-CU, either free drug on its own, a simple mixture of the two free drugs, or saline. The PD dose was 5.32 mg/kg, and the CU dose was 2.55 mg/kg. Three healthy rats not receiving any treatment served as a naive control.

On every other day starting on day 14 after arthritis induction, swelling was assessed in each limb of all animals as described and we summed all scores for each animal to obtain an overall score.

On day 22 after arthritis induction, right ankle joints were dissected from rats in each of the groups. Hind paws were fixed in 4% paraformaldehyde for 72 h, decalcified with 15% neutral EDTA solution for 60 days at room temperature, then embedded in paraffin. Thin sections were cut and stained with hematoxylin-eosin.

**Pro-Inflammatory Cytokine and Anti-Inflammatory Expression in Serum of Arthritic Rats After Nanoparticles Treatment**

On day 22 after arthritis induction, blood was collected from arthritic rats in each of the groups. Serum levels of pro-inflammatory cytokines (TNF-α, IL-1β and IL-10) and the anti-inflammatory cytokine IL-10 were measured using ELISA kits according to the manufacturer’s instructions.

**Statistical Analysis**

Statistical analysis was performed using SPSS 19.0 (IBM, Chicago, IL, USA). Differences between two groups were assessed for significance using Student’s t test, and differences among more than two groups were assessed using ANOVA. Differences associated with p<0.05 were considered significant. All results were expressed as mean ± SD.

**Results**

**Characterization of Drug-Loaded Nanoparticles**

Dynamic light scattering showed the average hydrodynamic diameter of N-PD/CU to be 150.4 ± 2.4 nm; N-PD, 136.90 ± 5.0 nm; and N-CU, 148.0 ± 3.1 nm (Figure 1A–C). Blank nanoparticles were smallest (Figure 1D). The polydispersity index (PDI) was less than 0.3 for all nanoparticles (Figure 1A), indicating uniform size. Average zeta potential was −21.6 ± 1.9 mV for N-PD/CU, −18.6 ± 1.9 mV for N-PD, −25.2 ± 2.6 mV for N-CU, and −26.5 ± 2.2 mV for blank nanoparticles (Figure 2A). Transmission electron microscopy showed that all types of nanoparticles were monodisperse and uniformly spherical (Figure 1E), with sizes of 100–200 nm. Drug encapsulation efficiency of N-PD/CU was 88.75 ± 1.82% (w/w) for PD and 85.79 ± 1.43% (w/w) for CU, for a total drug loading efficiency of 2.36 ± 0.04% (w/w) (Figure 2A). Encapsulation and drug loading efficiencies of N-PD were, respectively, 91.61 ± 1.82% and 2.30 ± 0.03%, N-CU showed slightly lower encapsulation and drug loading efficiencies.

Average zeta potential was −21.6 ± 1.9 mV for N-PD/CU, −18.6 ± 1.9 mV for nanoparticles containing only PD, −25.2 ± 2.6 mV for nanoparticles containing only CU, and −26.5 ± 2.2 mV for blank nanoparticles (Figure 2A). Transmission electron microscopy showed that all types of nanoparticles were monodisperse and uniformly spherical (Figure 1E), with sizes of 100–200 nm. Drug encapsulation efficiency of N-PD/CU was 88.75 ± 1.82% (w/w) for PD and 85.79 ± 1.43% (w/w) for CU, for a total drug loading efficiency of 2.36 ± 0.04% (w/w) (Figure 2A). Encapsulation and drug loading efficiency of N-PD were respectively 91.61 ± 1.82% and 2.30 ± 0.03%, which were both a bit higher than N-CU.

**Drug Release from Nanoparticles in vitro**

Compared to the free drug solution, PD and CU were released more slowly from N-PD/CU. By around 10 h, nearly 100% of free CU was released from the corresponding free drug solution, compared to 24 h for PD. In contrast, by 72 h, only 49.22 ± 1.95% of PD had been released from N-PD/CU (Figure 2B), similar to 48.72 ± 1.24% of CU (Figure 2C). These results indicate that N-PD/CU allows controlled drug release.

**Nanoparticles Cytotoxicity in vitro**

Cell viability remained about 94% after exposure to HSA concentrations between 0.1 and 1.2 μmol/mL (Figure 3). Cells exposed to N-PD/CU showed similar viability as those exposed to blank nanoparticles, suggesting that the HSA-based drug carrier is nontoxic. In fact, cells exposed to N-PD/CU showed higher viability than cells exposed to a simple mixture of both free drugs.
Pharmacodynamics in vitro

To observe the anti-inflammatory effects of N-PD/CU on activated macrophages, levels of inflammatory cytokines were measured after various treatments. The N-PD/CU group showed significantly lower levels of TNF-α and IL-6 in the supernatant after spinning down cultured cells than other groups (Figure 4A and C, *p<0.05, **p<0.001 or ***p<0.0001), and it showed significantly lower levels of IL-1β than the N-CU group (Figure 4B, *p<0.05). Conversely, the N-PD/CU group showed significantly higher levels of IL-10 than other groups except N-CU (Figure 4D, *p<0.05 or **p<0.001). These results suggest that N-PD/CU can inhibit the production of pro-inflammatory cytokines and promote the release of anti-inflammatory IL-10 to a greater extent than single-loaded nanoparticles, either free drug on its own, or a simple mixture of the two drugs.

To explore in greater detail how the drug cargoes, PD and CU, work against RA in activated macrophages, we analyzed the effects of N-PD and N-CU on inflammatory cytokine levels. N-PD led to significantly lower levels of TNF-α and IL-1β than N-CU compared to PBS (Figure 4A and B, *p<0.05, **p<0.001, ***p<0.0001), while N-CU led to significantly higher levels of IL-10 than N-PD compared to PBS (Figure 4D, *p<0.05). These results suggest that PD primarily inhibits the production of pro-inflammatory cytokines TNF-α and IL-1β, while CU effectively promotes the production of anti-inflammatory cytokine IL-10.

Biodistribution and Pharmacokinetics of Nanoparticles

Encapsulating PD and CU in nanoparticles significantly increased their concentrations in liver, spleen, and joints at 15 min, 1 h, 2 h and 4 h after injection into AIA rats.
In fact, the nanoparticles increased drug concentration in joints relative to the simple drug mixture by 2.1-fold at 15 min (*p < 0.05), 3.5-fold at 1 h (**p < 0.001), 6.6-fold at 2 h (***p < 0.0001), and 8.4-fold at 4 h (**p < 0.001). AUC$_{0-4}$h of PD in joints was significantly higher with N-PD/CU (6.61 ± 0.57 μg/g·h) than with the simple mixture of both drugs (1.92 ± 0.10 μg/g·h, *p < 0.05).

Furthermore, there was no significant difference in drug levels between the two treatment groups in heart or lung at any of the time points (Figure 5A, p > 0.05). Levels of the drugs were similar between the two treatment groups at 0.25, 1 and 2 h (Figure 5A, p > 0.05), but the level of N-PD/CU was higher than the level of the simple mixture at 4 h.

Biodistribution of N-PD/CU was significantly higher than N-CU in heart, liver, spleen, and joints at 15 min, 1 h, 2 h and 4 h (Figure 5B). AUC$_{0-4}$h was significantly higher for N-PD/CU (2.20 ± 0.25 μg/g·h) than for the drug mixture (0.71 ± 0.04 μg/g·h, *p < 0.05).

By analyzing drug concentration-time profiles, we found that encapsulating PD into N-PD/CU led to much higher PD concentrations in the plasma at early time points (Figure 6A). The area under the curve (AUC) and C$_{max}$ were, respectively, 2.62- and 2.31-fold higher for N-PD/CU than for the simple mixture of both free drugs. Similar results were obtained for CU (Figure 6B). Furthermore, t$_{1/2}$ for N-PD/CU was 2.36 times higher than for N-PD and 1.25 times higher than for N-CU (Figure 6C).

Figure 7 compares the pharmacokinetics of N-PD/CU with the pharmacokinetics of the simple mixture of two drugs in the plasma of AIA rats. In the case of both drugs, AUC and C$_{max}$ were significantly higher for N-PD/CU than for the simple mixture. In animals treated with N-PD/CU, Re$_{joint}$ and Ce$_{joint}$ were 3.44 and 2.13 for PD and 3.10 and 2.82 for CU. These results indicate that HSA-based nanoparticles lead to greater joint accumulation of both drugs.

Distribution of N-PD/CU in Arthritic Rats
Real-time fluorescence imaging showed that DID/N-PD/CU accumulated in joints more than free DID (Figure 8). At 4 h and 12 h, DID/N-PD/CU showed detectable fluorescence in
the hind limbs of AIA rats, whereas free DID did not. Even at 24 h, most free DID fluorescence was in internal organs. These results suggest that N-PD/CU can target inflammatory lesions, confirm with the results of pharmacokinetics that HSA-based nanoparticles lead to greater joint accumulation.

**Therapeutic Efficacy of Nanoparticles in vivo**

Arthritic rats were treated by tail vein injection with saline, free PD, free CU, N-PD/CU, N-PD or N-CU. Each group was monitored for changes in body weight, swelling.

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**Figure 4** Expression of pro-inflammatory cytokines and anti-inflammatory cytokine in cell-free supernatant of activated macrophage treated with different preparations. Levels of (A) TNF-α, (B) IL-1β, (C) IL-6 and (D) IL-10 were assayed. Data shown are mean ± SD (n=3). *p<0.05, **p<0.001, ***p<0.0001 vs cells treated with N-PD/CU, †p<0.05, ††p<0.001, †††p<0.0001 vs cells treated with PBS.

**Figure 5** Biodistribution profiles of N-PD/CU and PD/CU. (A) The concentration of PD in different plasma, organs and joints at different time points in AIA rats treated by N-PD/CU and PD/CU. (B) The distribution of CU in different plasma, organs and joints at different time points in AIA rats treated with N-PD/CU and PD/CU. Data represent the mean ± SD (n=5). *p<0.05, **p<0.001, ***p<0.0001 vs animals treated with PD/CU.
scores and histopathology. Histological sections of joints in the right hind limb were analyzed under the microscope, using healthy rats as a control group.

Body weight in rats injected with N-PD/CU increased continuously from day 14 (booster injection) to day 22; their body weight was similar to that of healthy, untreated rats (Figure 9A). Mean arthritis scores were lowest in animals treated with N-PD/CU (Figure 9B), which showed barely noticeable swelling of joints. In the joints of rats treated with saline, histological examination of right hind limbs revealed massive synovial cell lining, extensive pannus formation and severe bone destruction (Figure 9C); the histopathology score was highest (>7.0) in this group. Animals treated with either free drug showed much thicker synovial cell lining and greater bone destruction than animals treated with the corresponding single-loaded nanoparticles. Animals treated with a simple mixture of the two drugs showed less synovial cell lining and less severe bone destruction, with a histopathology score of 4.0. Rats treated with N-PD/CU showed minor pannus formation and obvious bone protection, with a histopathology score of 2.0.

Effects of Nanoparticles on Pro-Inflammatory and Anti-Inflammatory Cytokine Production in vivo

The cytokines TNF-α, IL-1β, and IL-6 drive progression of rheumatoid arthritis, while anti-inflammatory cytokines slow progression. Levels of all three pro-inflammatory cytokines in serum were significantly lower in animals treated with N-PD/CU than in animals in other groups (*p<0.05, **p<0.001 or ***p<0.0001, Figure 10A–C). Conversely, IL-10 levels in serum were significantly higher in animals treated with N-PD/CU than in animals in other groups (**p<0.001 or ***p<0.0001, Figure 10D).

| parameters       | N-PD/CU       | PD     | CU     | PD     | CU     |
|------------------|---------------|--------|--------|--------|--------|
| $AUC_{0-\infty}$ ($\mu$g/g.h) | 63.29 ± 5.13$^a$ | 30.28 ± 1.14$^b$ | 24.17 ± 3.13 | 14.12 ± 1.24 |
| $C_{\text{max}}$ (μmol/g) | 49.62 ± 6.60$^a$ | 26.62 ± 0.69$^b$ | 21.51 ± 3.58 | 14.01 ± 1.34 |
| $T_{1/2}$ (h)        | 1.20 ± 0.27$^a$ | 0.67 ± 0.04$^b$ | 0.73 ± 0.18 | 0.56 ± 0.14 |

Abbreviations: $AUC_{0-\infty}$: area under drug concentration-time curve, $C_{\text{max}}$: maximum drug concentration.

$^a$ $p<0.05$ versus PD of PD/CU group.

$^b$ $p<0.05$ versus CU of PD/CU group.

Figure 6: Plasma concentration-time curves of N-PD/CU and PD/CU. (A) Concentration-time curves of PD. (B) Concentration-time curves of CU. Data represent the mean ± SD (n=5). (C) Pharmacokinetic parameters of N-PD/CU and PD/CU (n=5).
These results suggest greater therapeutic efficacy with N-PD/CU than with single-loaded nanoparticles, either free drug on its own, or a simple mixture of the two drugs.

**Discussion**

HSA is an attractive drug carrier, offering high loading capacity for a wide variety of compounds, especially water-insoluble drugs. We found that the encapsulation and drug loading efficiency of HSA-based nanoparticles were similar regardless of whether they carried PD and CU separately or together. Both drugs were released slowly and sustainably from N-PD/CU, which aids in maintaining therapeutic levels of the drugs in circulation.

Liposomes are perhaps the most extensively studied carrier for targeted delivery of drugs to inflamed joints in...
RA treatment. However, hydrophobic drugs can be encapsulated only into the bilayer of liposomes rather than into their hydrophilic cores. This translates to rapid release in vivo because the lipid bilayer is destabilized.\textsuperscript{11,31} In our study, we prepared HSA-based nanoparticles by high-pressure homogenization, which utilizes high-shear cavitation to form a new disulfide bond in the free sulfhydryl group of albumin, which allows cross-linking into nanoparticles.\textsuperscript{32} In this approach, the drugs formed a tight complex with HSA, with the drugs mainly in the core, E80 and cholesterol helped disperse and stabilize the drug-albumin complexes. These HSA-based nanoparticles avoided the rapid drug release of typical liposomes. In other approaches,\textsuperscript{33,34} researchers have used hyaluronic acid to load hydrophobic drugs such as CU, and phospholipids was used to enhance the stability of the drug-loading system too, obtaining higher drug loading and better stability in vivo than liposomes.

In RA, the formation of synovial hyperplasia vasospasm is associated with early cartilage and bone erosion, and the interface between this vasospasm and cartilage contains a large number of activated macrophages, also known as M1 macrophages. Such macrophages drive the progression of RA by continuously secreting a variety of pro-inflammatory cytokines such as TNF-$\alpha$, IL-1$\beta$, and IL-6. This makes macrophages a potential therapeutic target in inflammatory diseases.\textsuperscript{35,36} In our work, we used LPS-activated macrophages as an inflammatory model to explore the pharmacodynamic efficacy of N-PD/CU and anti-inflammatory mechanism of PD and CU separately. N-PD/CU inhibited pro-inflammatory cytokines and promoted production of anti-inflammatory cytokine IL-10 more effectively than other preparations. PD inhibited the production of pro-inflammatory cytokines TNF-$\alpha$ and IL-1$\beta$ to a greater extent than CU, while CU effectively promoted the production of anti-inflammatory cytokine IL-10. These results suggest that CU can synergize with PD for combination therapy of RA by simultaneously inhibiting the level of pro-inflammatory cytokines and increasing the level of anti-inflammatory cytokines.

PD and CU reduced the level of IL-6 to similar extents, which is consistent with some reports that CU can strongly

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\caption{Therapeutic efficacy of nanoparticles in vivo. (A) Body weight variation after given different treatments. (B) Mean arthritis scores were calculated for the joints in the right hind limb following different treatments. Data shown are mean ± SD (n=3). (C) Photomicrographs of histological sections of ankle joints from animals after different treatments. Arrows indicate finger-like pannus formation; asterisks, bone destruction. Bar, 100 nm.}
\end{figure}
reduce IL-6 levels in inflammation. Consistent with anti-inflammatory effects, CU reduced oxidative damage in the synovial fluid and improved the production of IL-10, it can also down-regulate pro-inflammatory cytokines such as IL-1, IL-8, as well as monocyte chemotactic protein 1–1 (MCP-1), inducible nitric oxide synthase and nitric oxide. CU can also modify the activity of T and B cells, macrophages, neutrophils, dendritic cells and natural killer cells. It inhibits the proliferation of lymphocytes and their secretion of IL-4, IL-5, and granulocyte-macrophage colony-stimulating factor.

N-PD/CU accumulated in joints to a much greater extent than free PD/CU, suggesting that encapsulating PD and CU with HSA can improve their bioavailability. The joint-targeting ability of N-PD/CU is due to the ELVIS effect. Therefore, the therapeutic efficacy of PD will be increased, and the side effects caused by systemic distribution will be reduced.

Our results provide further evidence that encapsulating GCs into biomimetic carriers can improve their therapeutic efficacy in AIA rats. These positive results are also due to the stability, high accumulation and sustained drug release of N-PD/CU. Co-delivering GCs with CU reduces the dose of GCs needed, using non-toxic and biocompatible HSA as a nanocarrier reduces the toxicity of drugs to cells in vitro compare to free drugs. Therefore the side effects of GCs reduce, which may improve the clinical application of GCs. Future studies should verify the reduction in side effects in vivo as well as the optimal ratio of PD and CU for combination therapy.

Conclusion
We successfully prepared HSA-based nanoparticles for codeelivery of PD and CU by high-pressure homogenization. The N-PD/CU exhibited synergistic anti-inflammation effect against macrophages and accumulated to a greater extent at inflamed joint sites than single-loaded nanoparticles or a simple mixture of the drugs. This resulted in superior therapeutic effect in rats with adjuvant-induced arthritis. HSA-based nanoparticles show promise for combination chemotherapy.

Abbreviations
OA, oleic acid; lipoid E80, purified yolk lecithin; RA, rheumatoid arthritis; PD, prednisolone; CU, curcumin; HSA, human serum albumin; N-PD/CU, Co-delivery of prednisolone and curcumin in human serum albumin nanoparticles; N-PD, prednisolone in HSA-based nanoparticles.

Figure 10 Production of pro-inflammatory cytokines and anti-inflammatory cytokines in arthritic rats treated with different preparations. Healthy, untreated animals served as a control (Normal). Serum levels of (A) TNF-α, (B) IL-1β, (C) IL-6 and (D) IL-10 were assayed. Data shown are mean ± SD (n=3). *p<0.05, **p<0.001, ***p<0.0001 vs animals treated with N-PD/CU.
nanoparticles; N-CU, curcumin in HAS-based nanoparticles; EE, encapsulation efficiency; DLE, drug loading efficiency; blank, blank HSA-based nanoparticles; MTT, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide; DID, Diocadecyl tetramethyl Indodicarbocyanine; ELISA, Enzyme-linked immunosorbent assay; ELVIS, extravasation through leaky vasculature and subsequent inflammatory cell-mediated sequestration; EPR, enhanced permeability and retention; PBS, phosphate-buffered saline.

Acknowledgements
This work was supported by Sichuan Science and Technology Program (2017RZ0048) and the National Natural Science Foundation of China (Grant No. 81803478).

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

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The authors report no conflicts of interest in this work.

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