Preparation and study of the physicochemical properties of psoralen-doxorubicin nanostructured lipid carriers

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The purpose of the present study was to prepare psoralen-doxorubicin nanostructured lipid carriers (PSO-DOX-NLC) with better physicochemical properties for further study on its multidrug resistance reversal activity on leukemia cells. PSO-DOX-NLC was synthesized by emulsion evaporation and solidification at a low temperature. PSO-DOX-NLC morphology and particle size were examined using transmission electron microscopy, and Zeta potential was determined using photon correlation spectroscopy; encapsulation efficiency was also investigated. The results showed that PSO-DOX-NLC possessed a rounded structure, with a mean (± SD) particle size of 128.7±1.8 nm, a polydispersity index of 0.22±0.01, a Zeta potential of 20.17±0.31 mV, a psoralen encapsulation rate of 76.56% and a DOX encapsulation efficiency of 89.48%. Thus, PSO-DOX-NLC prepared by emulsion evaporation and solidification at a low temperature had good physicochemical properties.

Key Words: Doxorubicin; Nanostructured lipid carriers; Physicochemical property; Psoralen

METHODS

Preparation of PSO-DOX-NLC

PSO-DOX-NLC was prepared by emulsion evaporation and solidification at a low temperature. Taking solid lipid GMS: ATO5 (1:1) in the formula dosage, 20% liquid lipid of 1349, lecithin and PSO, dissolving in 20 mL absolute ethyl alcohol with 78°C water bath, and then obtained the organic phase. While taking a proper amount of DOX and F68 to dissolve in 1 mL pure water and heating to 78°C, the inner water phase was separated, and then taking Tween80 in formula dosage to dissolve in 20 mL pure water, the external water phase was obtained. While stirring at 1000 rpm, 1 mL internal phase was injected into the stirring organic phase and was stirred for 10 min to form colostrum. The colostrum was injected into the stirring external water phase with same temperature, maintaining stirring and emulsifying to form multiple emulsion. The multiple emulsion was poured into 20 mL 4°C ice water rapidly, when it volatilized to 5 mL, then solidified in low temperature for 1 h and filtered through a 0.45 μm filtermembrane. Finally the PSO-DOX-NLC was obtained.

Chromatographic conditions

C18 as chromatographic column (250 mm × 4.6 mm, 5μm), methanol-sodium acetate buffer (pH was adjusted to 3.6 with glacial acetic acid) (65:35) as the mobile phase, 1.0 mL/min as the flow rate, 254 nm as the ultraviolet detection wavelength, 25°C as the column temperature.

Specific inspection study

Adding the proper amount of blank NLC and PSO-DOX-NLC to the mobile phase for ultrasonic demulsification and filtering with 0.45 μm filtermembrane, then taking PSO solution, DOX solution and the mixed reference solution of PSO-DOX, all in a specific concentration, to sample respectively according to 2:2:1 chromatographic condition.
Linear relationship study

Precisely adding an appropriate amount of reference solution to 10 mL volumetric flasks, diluting with methanol to obtain concentrations of 1.0, 2.0, 4.0, 8.0, 16.0 and 32.0 μg/mL mixed reference solution of PSO-DOX. With peak area (Y) as the ordinate, mass concentration (X, μg/mL) as the abscissa, HPLC was used to investigate the peak area and linear regression was made.

Precision test

Taking the reference solution of 4μg/ml, 16μg/ml and 32μg/ml which are in the low, medium and high concentration respectively, investigating their peak area 5 times a day by HPLC in parallel and continuing for 5 days, then calculated the precision RSD.

Recovery test

Taking 1 mL blank SLN, adding 0.8 mL, 3.2 mL and 6.4 mL reference solution into 10 mL volumetric flasks respectively, diluting with mobile phase to 10 mL, and obtained the low, medium and high concentration samples. Then breaking emulsion via ultrasound, filtering with 0.45 μm membrane filtration, investigate their peak area for the average recovery using high-performance liquid chromatography (HPLC), and calculated the ultrafiltration recovery rate.

Stability test

A reference solution of PSO-DOX (8 μg/mL), was sampled at 0 h, 2 h, 4 h, 6 h, 12 h and 24 h, respectively, for calculation of peak area.

Determination method of encapsulation efficiency

The encapsulation efficiency was determined by ultrafiltration in the experiment. Adding an appropriate amount of PSO-DOX-NLC solution into an ultrafiltration centrifuge tube, then investigated the content of free drug (W_f) by HPLC after centrifuging in 12,000 rpm for 20 min. Adding another 1.0 mL PSO-DOX-NLC solution into the 10 mL volumetric flask, diluting with the mobile phase to 10 mL, filtering with 0.45 μm membrane filter, and investigated the content of total drug (W_t) using HPLC. The encapsulation efficiency was then calculated using the following formula:

$$EE\% = \frac{W_{total} - W_{unencapsulated}}{W_{total}} \times 100\%$$

Investigation of morphology

Putting a proper amount of PSO-DOX-NLC on a copper net with carbon film, and dyeing with 2% phosphotungstic acid, then observing under transmission electron microscopy after natural drying.

Determination of particle size and electric potential

The PSO-DOX-SLN solution was diluted and its mean particle size and electric potential were determined using a Zeta size nanolaser particle size analyzer (NanoZS90, Malvern Instruments, United Kingdom).
Figure 2) Transmission electron micrograph of psoralen-doxorubicin nanolipid carriers

RESULTS

Specific inspection study

Under the chromatographic conditions described above, all samples were effectively separated, indicating that the method had high specificity, which is shown in Figure 1.

Linear relationship study

The standard curve equation of PSO was \( Y=1.1\times10^{5}X+14742 \) (\( r=0.9998 \)) and that of DOX was \( Y=54195X+1877.9 \) (\( r=0.9999 \)), which suggested that the standard curves for PSO and DOX were in good linear in the concentration ranging from 1 μg/mL to 32 μg/mL.

Precision, recovery and stability test

The results showed that all the RSD of the precision test were under 3%, the recovery of the three concentrations of PSO and DOX were between approximately 95% and 105% and RSD<3%, The RSD of PSO and DOX were 0.79% and 1.55%, respectively (n=6), suggesting that the precision, recovery and stability of the method were good.

Morphology of PSO-DOX-NLC

The morphology of the PSO-DOX-NLC is shown in Figure 2. The particles had a spherical shape, were well dispersed and basically had no adhesion.

The determination of particle size and Zeta potential

The mean (± SD) particle size of the nanoparticles was 128.7±1.8 nm, PDI was 0.22 0.01 (n=3), and the potential was (20.17±0.31) mV, as shown in Figure 3.

Discussion

NLC were prepared by the addition of liquid lipid on the basis of solid lipid nanoparticles. The effect of efflux drug of the solid lipid in the perfect lattice was decreased, and the loading rate and encapsulation efficiency were improved (5,6). In the present experiment, the composite nano structure with high encapsulation efficiency and good physicochemical properties was prepared by emulsion evaporation and solidification at a low temperature, and the method of simultaneous detection of PSO and DOX encapsulation efficiency was established.

Existing research (7) shows that poloxamer with space steric effect, can prevent nanoparticle aggregation. Therefore, in the present experiment, an appropriate amount of F68 was added to the inner water phase and improved the stability of collostrum, while Tween80 was added to the external water phase to prepare nanoparticles with good appearance and properties. Lecithin, not only a lipid material, but also a type of emulsifier, can improve the stability of nanoparticles. GMS, commonly used as the lipid material, make the nanoparticles have uniform particle size and good dispersion degree and, at the same time, it is a W/O emulsion. Besides, nanoparticles prepared with ATO5 also show good stability and high entrapment efficiency. All experiments simultaneously adopted two types of the solid lipid mentioned above.

Figure 3) Particle distribution and Zeta-potential of PSO-DOX-NLC

Adriamycin, with a certain water solubility, was dealt with anionic polymer materials (8) and dehydrochlorination in the preliminary experiment. However, the stability of the adriamycin nanoparticles prepared this way was inferior to that prepared by multiple emulsion method. Therefore, the multiple emulsion method was chosen.

According to the principle of similar compatibility, fat soluble drugs can more easily be prepared into nanoparticles (9). However, the experimental results showed that encapsulation efficiency of adriamycin was inferior to that of PSO. The reason may be that adriamycin, a type of cationic drug, tends to combine with lipid materials and adhere to its surface, which has an impact on the release of the drug. The specific mechanism needs further research.

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