Development and characterization of a novel nanoemulsion drug-delivery system for potential application in oral delivery of protein drugs

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Background: The stability of protein drugs remains one of the key hurdles to their success in the market. The aim of the present study was to design a novel nanoemulsion drug-delivery system (NEDDS) that would encapsulate a standard-model protein drug – bovine serum albumin (BSA) – to improve drug stability.

Methods: The BSA NEDDS was prepared using a phase-inversion method and pseudoternary phase diagrams. The following characteristics were studied: morphology, size, zeta potential, drug loading, and encapsulation efficiency. We also investigated the stability of the BSA NEDDS, bioactivity of BSA encapsulated within the NEDDS, the integrity of the primary, secondary, and tertiary structures, and specificity.

Results: The BSA NEDDS consisted of Cremophor EL-35, propylene glycol, isopropyl myristate, and normal saline. The average particle diameter of the BSA NEDDS was about 21.8 nm, and the system showed a high encapsulation efficiency (>90%) and an adequate drug-loading capacity (45 mg/mL). The thermodynamic stability of the system was investigated at different temperatures and pH levels and in room-temperature conditions for 180 days. BSA NEDDS showed good structural integrity and specificity for the primary, secondary, and tertiary structures, and good bioactivity of the loaded BSA.

Conclusions: BSA NEDDS showed the properties of a good nanoemulsion-delivery system. NEDDS can greatly enhance the stability of the protein drug BSA while maintaining high levels of drug bioactivity, good specificity, and integrity of the primary, secondary, and tertiary protein structures. These findings indicate that the nanoemulsion is a potential formulation for oral administration of protein drugs.

Keywords: nanoemulsion, drug-delivery system, protein drug, oral administration, stability

Introduction

Many protein targets and protein drugs have recently been discovered, with approximately a dozen therapeutic protein drugs receiving regulatory approval each year around the world.1,2 Over 125 protein drugs are already in the US market, and the value of advanced drug-delivery systems in this market had reached $121 billion by 2010.3 Although many varieties of protein drugs have been designed for therapeutic purposes, several challenges remain, including enzymatic susceptibility, stability during storage period, and efficacy after administration.4 Thus, protein stability is one of the major hurdles for a drug’s successful entry into the market.5 At present, many of the marketed protein drugs are mainly designed for injective delivery. Because of the thermal instability of protein drugs, they need to be stored and transported at cold temperatures in order to ensure long shelf life.5,6
To overcome the crucial problem of protein instability, stabilizers such as nonionic surfactants, cyclodextrin derivatives, poloxamers, glycerol, ethylene glycol, glucose, sucrose, and trehalose have been used. However, the effects of these stabilizers vary among different proteins. With a view to enhancing the stability of therapeutic peptides and proteins, many studies have focused on the use of such drug-delivery systems as microspheres, nanoparticles, liposomal systems, and traditional emulsions. However, these drug-delivery systems have several drawbacks, such as low encapsulation efficiency and drug loading, use of inorganic solvents such as methylene chloride and acetone (which induce toxicity), low yield and high production costs. A nanoemulsion-drug delivery is a novel, thermodynamically stable delivery system that can enhance the stability of a protein drug by protecting it from oxidative and enzymatic degradation. Bovine serum albumin (BSA) has often been used as a standard and model protein drug in drug development and delivery, using carriers such as microparticles, nanospheres, nanoparticles, and gels.

The nanoemulsion-drug delivery system (NEDDS) is an isotropic system in which two immiscible liquids are combined to form a single phase by application of appropriate surfactants or by mixing to form a droplet with a diameter range of approximately 0.1–100 nm. Such systems can also be formed by heterogeneous dispersion of oil droplets in an aqueous medium stabilized using surfactant molecules. In recent years, the focus on NEDDS has increased because of its distinctive structure and properties, including a small droplet size, possibility of fabrication with biocompatible materials, increased dissolution rate and solubility, improved diffusion across an unstirred aqueous layer, and enhanced mucosal permeability. NEDDS possesses high kinetic and long-term thermodynamic stability and transparency. Therefore, it has been widely used to improve the stability of drugs such as ramipril, celecoxib, hyaluronic acid, β-cypermethrin, β-carotene, β-cyclodextrin and Bacillus anthracis protective antigen.

NEDDS possesses a good storage capacity and thermodynamic stability for drug delivery. In this study, we chose BSA as our model drug. And also, we prepared and optimized BSA NEDDS by using a phase-inversion method and pseudoternary phase diagrams. The delivery system was evaluated using transmission electron microscopy, atomic force microscopy, Nano ZS90 (Malvern Instruments, Malvern, UK). We also tested the following characteristics of the system stored at room temperature for 180 days: thermodynamic stability, specificity, bioactivity and stability of this system and model protein drug, BSA, including integrity of the primary, secondary, and tertiary structures.

Materials and methods

Materials

BSA (MW 66 kDa; fat, purity 98.3%; lot 071K7602) was purchased from Sigma-Aldrich Chemicals (St Louis, MO) and used without further purification. Isopropyl myristate (IPM) was purchased from Croda (Goole, UK). Cremophor EL 35 (polyethoxylated castor oil) was obtained from BASF (Mumbai, India). Propylene glycol and Tween 20 were obtained from Shanghai Chemical Reagent Corporation (Shanghai, China). Low molecular protein marker (14.4–97.4 kDa) and prelyte protein ladder marker (17–170 kDa) were purchased from Fermentas International (Burlington, Canada). A micro-BCA protein assay kit was purchased from Wuhan Boster Bioengineering, Wuhan, China. Rabbit anti-mouse BSA monoclonal antibody was purchased from Beijing Saichi Shengwu Keji (Beijing, China). Horseradish peroxidase (HRP) goat anti-mouse immunoglobulin G (IgG) was purchased from Beijing Zhongshan Golden Bridge Biotechnology (Beijing, China). High-purity Milli-Q water (Millipore, Billerica, MA) was used for all solutions, with a resistance value of >18.2 MV/cm. All other chemicals used in the study were of analytical reagent grade.

Pseudoternary phase diagram construction

On the basis of the results of preliminary experiments, we chose IPM as the oil phase, Cremophor EL-35 as the surfactant, propylene glycol as the cosurfactant, and normal saline as the aqueous phase. The surfactant and the cosurfactant were mixed (Smix) in different mass ratios (1:1, 2:1, and 3:1). The pseudoternary phase diagrams for these mixtures were created using titration at room temperature to ascertain the ideal concentration ranges of the components. Nine different combinations of oil and Smix (1:9, 2:8, 3:7, 4:6, 5:5, 6:4, 7:3, 8:2, and 9:1) were added drop-wise to the aqueous phase by using gentle agitation. Transparent and easily flowing nanoemulsion was observed. Using Origin 7.0 software (OriginLab, Northampton, MA), we generated the physical state of the nanoemulsion by creating a pseudo-three-component phase diagram, with the first axis representing the aqueous phase, the second axis representing the oil, and the third axis representing the mixture of the surfactant and cosurfactant at a fixed mass ratio.
Formulation of the BSA NEDDS

From the pseudoternary diagram and visual observations, a precisely defined composition was selected for the nanoemulsion-drug delivery system. Briefly, Smix was prepared by mixing Cremophor EL 35, propylene glycol in the ratio 2:1 (w/w) and the mixture was stirred at 600 rpm at 25°C. Next, IPM was added to Smix to obtain the desired ratio of 9:1 (Smix–IPM) and stirred for 5 minutes, after which the mixture was cooled to room temperature. This was followed by the addition of aqueous phase (normal saline) to the Smix–IPM mixture while stirring, such that the final concentration of aqueous phase in the mixture was 20% w/w. The contents were stirred for 15 minutes to ensure nanoemulsion formation, which was monitored by checking the optical clarity and physical properties of the system. To prepare the novel BSA NEDDS, a blank formula and different concentrations of BSA power and aqueous solution, being 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, and 60 mg/mL, were used following the same protocol and method to the pseudoternary phase diagram construction, with the exception of water being replaced by BSA as the aqueous phase.

Characterization of the BSA NEDDS

Sample solutions were imaged using transmission electron microscopy (Tecnai-10 TEM; Hillsboro, OR). A drop of the particle suspension was placed onto a carbon-coated copper grid obtained from Ted Pella (Redding, CA) to yield a thin liquid film. The films were negatively stained with 1% phosphotungstic acid solution (w/w; pH 7.4) for 1 minute. A high-resolution atomic-force microscope (IPC-208B; Chongqing University, Chongqing, China) was used to characterize the molecular morphology of the BSA NEDDS. Approximately 0.05 mL of the sample solution was dropped onto microscopic glass slides with gold coating to fracture the film, which was then dried and tested. The measurement was performed under the following conditions: tungsten probes (force constant, 0.06 N m); scan range, 10.5 nm × 10.5 nm; imaging mode, tapping mode; and scanning method, point by point scanning at room temperature. The G3DR software (Changqing, China) was used to process the data. The excess phosphotungstic acid solution was removed using filter paper, and stained samples were characterized at 80 kV. The average size and zeta potential of the BSA NEDDS were tested at 20°C by using a dynamic light-scattering instrument (Zetasizer) equipped with an argon ion laser.

Drug-loading and encapsulation efficiency

For sample preparation, triplicate aliquots of the blank and novel BSA NEDDS (1 g, approximately, 1 mL) were placed in sterile 2-mL eppendorf centrifuge tubes and diluted to a protein concentration of 250–500 mg/mL with normal saline. Two volumes of absolute ethanol were added to approximately 0.3 mL of blank or sample, and the mixture was shaken to ensure complete emulsification, then centrifuged at 13,000 × g for 30 minutes. The supernatant was removed, and the precipitate was mixed with 0.3 mL of the above solutions. The amount of BSA incorporated in the nanoemulsion was determined by centrifuging the BSA-loaded nanoemulsion and separating the supernatant. The precipitate was discarded, and the BSA content of the solution was then analyzed by using the commercially available bicinchoninic acid kit. The following equations were used in calculating for the percentage of drug loading and encapsulation efficiency:

\[
\text{Drug loading (\%)} = \left( \frac{\text{Total weight of protein drug}}{\text{Total weight of sample}} \right) \times 100%
\]

\[
\text{Encapsulation efficiency (\%)} = \left( \frac{\text{Actual/Theoretical protein drug loading}}{100} \right)
\]

Stability study of the BSA NEDDS

Thermodynamic stability

Thermodynamic stability was tested and revised according to the methods described by Shafiq et al.26 Briefly, selected formulations were centrifuged at 13,000 × g for 30 minutes at 4°C and 25°C, maintain for 48 hours. Six cycles of centrifugation were performed, and the appearance of these formulations was investigated by using indexes including turbidity, phase separation, precipitation, drug separation, demulsification, and creaming.

Effects of different temperatures and pH values

Effects of the temperatures and pH on particle size and zeta potential of the BSA NEDDS were measured directly. Effects of the increasing temperature in the range 20°C–80°C on particle size were determined by Nano ZS90 at 20°C. Effects of increasing pH in the range 1–14 on particle size and zeta potential were measured by Nano ZS90 at 20°C.

Short- and long-term stability

To examine the physical stability of this system, short- and long-term stability studies were carried out on the selected formulations. Samples were divided into three vials after production and stored at room temperature for 180 days. On days 0, 30, 60, 90, and 180, all samples were examined and centrifuged for 30 minutes at 13,000 × g. Turbidty, phase separation, precipitation, drug separation, breaking, and creaming were evaluated, and also the primary structural...
integrity was again assessed by sodium dodecyl sulfate polyacrylamide gel electrophoresis. Furthermore, the main physical characteristics, including particle size, polydispersity index, zeta potential, electrophoresis mobility, and conductivity were determined using photon correlation spectroscopy on days 0, 30, 60, 90, and 180 at 20°C.

**Stability study of the BSA enclosed in the BSA NEDDS**

**Relative content and bioactivity assay**

The relative content and bioactivity of four samples stored at room temperature for 0, 30, 90, and 180 days were determined. BSA was extracted from the nanoemulsion by using the drug-loading and encapsulation efficiency protocol described previously. The BSA content of the BSA NEDDS, which were stored at room temperature for 0, 30, 90, and 180 days, were detected using microbicinchoninic acid methods. Relative bioactivity was determined by enzyme-linked immunosorbent assay. The methods were performed as follows: 96-well flat-bottom plates (Costar, WA) were coated overnight at 4°C with 50 µg/mL of theoretical protein drug above four samples, blocked with a blocking buffer (10 mM PBS, pH 7.4, with 0.05% Tween 20), and kept at overnight at 4°C. Goat anti-mouse BSA monoclonal antibody (1:300) was used as the primary antibody. And HRP-conjugated goat anti-mouse IgG (1:10,000) was used as the secondary antibody. O-phenylenediamine was used as a substrate. The absorbance of the solutions was then determined using Bio-Rad model 680 microplate reader (Bio-Rad, Hercules, CA) set at a wavelength of 490 nm.

**Evaluation of primary structural integrity and specificity**

To determine the primary structural integrity and specificity of the BSA NEDDS, sodium dodecyl sulfate polyacrylamide gel electrophoresis and Western blotting were performed. The supernatant and precipitate of a blank nanoemulsion, the supernatant and precipitate of the BSA nanoemulsion, the supernatant and precipitate of a BSA emulsion, and native BSA aqueous solution were separated using 10% sodium dodecyl sulfate polyacrylamide gel electrophoresis. The gels were electroblotted onto polyvinylidene fluoride filter membranes at 25 mV for 15 minutes by using a TransBlot semidyry blotter (Bio-Multiphor II electrophoresis unit; GE Healthcare, Amersham, UK), after which they were blocked using a buffer (10 mM PBS, pH 7.4 with 0.05% Tween 20) and kept overnight at 4°C. Subsequently, the membranes were hybridized with goat anti-mouse BSA monoclonal antibody (1:300) for 1 hour at 37°C. HRP-conjugated goat anti-mouse IgG was used as the secondary antibody (1:10,000). Binding reaction was detected using diaminobenzidine color-development reagent (0.05% diaminobenzidine and 0.005% hydrogen peroxide in distilled water).

**Evaluation of the secondary structure**

To verify the integrity of the secondary structure of the BSA in this NEDDS, circular dichroic spectra of the BSA aqueous solution and three batches of the BSA NEDDS placed at room temperature for 30, 90, and 180 days were acquired at room temperature by using a Chirascan spectropolarimeter (Jasco 810, Easton, MD) equipped with ultraviolet light (190–250 nm) with a 0.01° path length and a step size of 1 nm. The lamp housing was purged with nitrogen, and an average of three scans was obtained. BSA was extracted from the NEDDS by using the previously mentioned protocol.

**Evaluation of the tertiary structure**

To verify the integrity of the tertiary structure of BSA in this NEDDS, we used Fourier transform infrared (FTIR) spectroscopy. FTIR spectra of the BSA aqueous solution and three batches of the BSA NEDDS placed at room temperature for 30, 90, and 180 days were obtained using a Spectrum One spectrophotometer (Lambda 950; PerkinElmer, Waltham, MA) separately 500, 550, 600, and 200 times diluting with water. Four samples were prepared using the potassium bromide disk method and examined using the transmission mode. A spectral range of 400–4000 cm⁻¹ with a resolution of 2 cm⁻¹ (accumulation of 64 scans) was used in order to obtain good-quality spectra. By monitoring the peak frequencies and locations of certain bands, all possible chemical changes and interactions were determined.

**Statistical analysis**

Values are expressed as means ± standard deviation. Statistical significance was determined using one-way analysis of variance by the least-significant-difference test. *P* < 0.05 was considered significantly different.

**Results**

**Construction of pseudoternary phase diagrams**

Pseudoternary phase diagrams were separately constructed for each Smix ratio (Figure 1). A Smix with a ratio (Km) of 2:1 in a mixture of Cremophor EL-35 (surfactant), propylene glycol (cosurfactant), isopropyl myristate (oil), and normal saline (aqueous medium) showed a large nanoemulsion area.
Figure 1 Pseudoternary phase diagrams of the BSA NEDDS, composed of the following constituents: oil, isopropyl myristate; surfactant, Cremophor EL-35; and cosurfactant, propylene glycol; drug-loading value was 45 mg/mL. (A) Surfactant and cosurfactant mixture ratio (Smix), 1:1; (B) Smix, 2:1; (C) Smix, 3:1. 

Note: The red area represents the nanoemulsion delivery system region.

Abbreviations: BSA, bovine serum albumin; NEDDS, nanoemulsion drug-delivery system; IPM, isopropyl myristate.

(Figure 1B). Figure 1C shows that the 3:1 Smix formula generated the smallest nanoemulsion area. The area of nanoemulsion increased with Smix proportions of 1:1 and 2:1 (Figure 1A and B). On the basis of these pseudoternary phase diagrams, we chose the Smix of Cremophor EL-35/propylene glycol at a ratio of 2:1, which showed the largest nanoemulsion area, as the main formulation for the BSA NEDDS.

Preparation of the BSA NEDDS formulation

It is important to determine the surfactant concentration accurately and to use the optimum concentration in the formulation. According to the results of the different concentrations and dosages of BSA, the BSA aqueous solution was added to the mixture of Cremophor EL-35, propylene glycol, and isopropyl myristate. BSA (45 mg/mL) was dissolved in the optimized mixture of 18% Cremophor EL-35, 9% propylene glycol, 3% isopropyl myristate (all w/w) and slowly added to normal saline under gentle stress to produce a transparent nanoemulsion-delivery system using the phase-inversion method.

Characterization of the BSA NEDDS

The droplets in the novel BSA NEDDS appeared dark and the surroundings were bright, such that a “positive” image was observed using transmission electron microscopy (Figure 2A). We randomly selected 500 droplets on the electron micrograph, and calculated the average particle size according to the scale and droplet-diameter measurements and statistical data. The results showed that the droplet particle sizes were mostly within the range of 1–100 nm, with an average size of approximately 21.0 nm. Examination of the atomic-force microscopy images revealed that most of the droplets exhibited spherical morphology, with a mean diameter of approximately 20 nm (Figure 2B). The droplets were solid, with surfaces mostly rugged with sags and crests. No droplets aggregated, especially in the BSA stabilized with the nanoemulsion, suggesting its stability, although proteins in the neighboring droplets might have been in contact with each
other. The differences observed among these morphological evaluations were essentially due to the fact in the Nano ZS90, the hydrodynamic radius of the particle was measured. On the contrary, transmission electron microscopy and atomic-force microscopy were performed on nanoemulsions of surface ions and solvent and nonsolvent molecules. Consequently, the actual mean diameter (obtained by transmission electron microscopy) was generally far smaller than the hydrodynamic diameter. Figure 2C shows that the average particle size in the novel BSA NEDDS was 21.8 nm, 75% of the particles were ≤50 nm and 90% <60 nm in size, and the range of distribution was very narrow. We determined the polydispersity index (PDI) value, which measures the spread of the particle-size distribution, in which a small value indicates a narrow particle-size range.22 In general, the novel BSA NEDDS exhibited a relatively limited size distribution, with a PDI value of 0.170, indicating a close size distribution and good stability of the nanoemulsion due to the reduced Ostwald ripening. Figure 2D shows that the average particle size in the novel BSA NEDDS was –24.21 mV and remained in the −40-mV to 40-mV range. The viscosity, refractive index, and dispersant refractive index were 0.8872 cP, 1.59 \( n_D \), and 1.330, respectively.

**Drug-loading and encapsulation efficiency**

The bicinchoninic acid method is a basic, classical technique used to detect BSA content. The standard curve equation is \( y = 0.0008x + 0.1636 \), \( r^2 = 0.9948 \), and the linear range is 25–1000 µg/mL. The drug-loading content and encapsulation efficiency of the BSA NEDDS can be measured in comparison to a standard curve. In this study, we obtained high encapsulation efficiency (>90%) and enough drug loading (45 mg/mL).

**Stability study of the BSA NEDDS**

**Thermodynamic stability**

After six cycles, the BSA NEDDS did not show any phase separation, turbidity, precipitation, drug separation, demulsification, or creaming. These observations suggest that this system has good thermodynamic stability at temperatures...
between 4°C and 25°C. Thermostability also differentiates nanoemulsion from emulsions, which are kinetically stable and eventually show phase separation. Therefore, the formulations obtained in our study were tested for thermodynamic stability by using centrifugation and heating–cooling and freeze–thaw cycles. Thermodynamic stability analysis showed no turbidity, phase separation, precipitation, drug separation, demulsification, or creaming before or after centrifugation (13,000 × g, 30 minutes) within the temperature range of 4°C–25°C.

**Effects of different temperatures and pH values**

Effects of the increasing temperatures or pH values on the particle size and zeta potential were determined using the Nano ZS90, as shown in Figure 3. Figure 3A shows the effect of increasing temperature on particle size of the BSA NEDDS. The particle size did not change significantly (it remained in the 20–30-nm range) at temperatures up to 80°C and increased rapidly at higher temperatures (>80°C), but remained under 100 nm. Figure 3B shows that no obvious change in zeta potential was observed within the pH range of 1–13, and the value remained between −40 mV and 20 mV. The average particle size of the novel BSA NEDDS remained at 20–30 nm at different pH values (Figure 3B). These results showed that the average particle size did not obviously change on using high temperature up to 80°C (Figure 3A) and that the zeta potential remained in the −40-mV to 40-mV range at different pH values (Figure 3B). The average particle size of the novel BSA NEDDS remained within the range of 20–30 nm at different pH values (Figure 3B).

**Short- and long-term stability**

The stability test showed that the novel BSA NEDDS did not change in visible appearance (ie, flocculation, stratification, precipitation, or creaming and emulsification) either

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**Figure 3 (A and B)** Effect of the different conditions on the BSA NEDDS. (A) Effect of particle size on the increasing temperature of the BSA NEDDS; (B) Effect particle size and zeta potential on the increasing pH values of the BSA NEDDS.

**Abbreviations:** BSA, bovine serum albumin; NEDDS, nanoemulsion drug-delivery system.
in the natural state or after high-speed centrifugation (13,000 × g, 30 minutes). The main physical characteristics results – particle size, polydispersity index, condition, zeta potential, electrophoresis mobility and melting point – measured using the Nano ZS90 at 20°C are shown in Figure 4A–F. The BSA NEDDS, which was stored at room temperature for 0, 30, 60, 90, and 180 days, was very stable, and the particle size of this system was within the range of 20–30 nm (Figure 4A). Fluctuations in particle size were very small, and there was no change observed even after storage at room temperature for 180 days. The zeta potential remained between −20 mV and −30 mV (Figure 4B). Melting point and mobility and condition (Figure 4C–E) results showed no obvious changes either. There were also no visible changes in the main physical characteristics of the BSA NEDDS.

Figure 5A shows that the sodium dodecyl sulfate polyacrylamide gel electrophoresis gels of the BSA aqueous solution and this system stored at room temperature for 30, 60, 90, and 180 days generated a clear band in lanes 1–5. At the same time points, BSA aqueous solution showed no obvious bands in lanes 6–9. These results imply that no changes occurred in the BSA NEDDS when it was stored at room temperature for 180 days.

Stability study of the BSA in the BSA NEDDS

Relative content and bioactivity analysis

Figure 5B showed no differences in the relative content among the fresh sample and the samples stored at room temperature for 30, 60, 90, and 180 days. The relative content and bioactivity of these samples did not show any decrease after storage at room temperature for 180 days. The results showed that BSA in the BSA NEDDS has very high stability. BSA retained high bioactivity when the BSA NEDDS was stored at room temperature for 180 days.

Primary structural integrity and structural specificity

The results of the evaluation of structural integrity are shown in Figure 6A, which shows that the aqueous BSA solution (lane 10), the supernatant nanoemulsion (lane 6), and the BSA emulsion (lane 8) each had one bright, clear band, each with a molecular weight of 66.4 kDa. The supernatant and precipitate of the blank nanoemulsion or emulsion

![Graphs](https://example.com/graph.png)

**Figure 4 (A–F)** Long-term stability results for the BSA NEDDS. (A) Particle size; (B) polydispersity index; (C) conductivity; (D) zeta potential; (E) electrophoresis mobility; (F) melting point.

**Abbreviations:** BSA, bovine serum albumin; NEDDS, nanoemulsion drug-delivery system.
had no obvious bands (lanes 1–4). The precipitate of this nanoemulsion-drug delivery system (lane 7) and the emulsion (lane 9) of BSA had one very narrow and dark band at 66.4 kDa. The results for lanes 7 and 9 show that the surfactant and cosurfactant partly precipitate the protein, but the supernatants of the nanoemulsion and emulsion gave more intense bands, so most of BSA in the nanoemulsion and emulsion was present in the supernatant. Meanwhile, the results showing that the supernatant and precipitate of the blank nanoemulsion or emulsion had no band showed that sodium dodecyl sulfate polyacrylamide gel electrophoresis can be adapted to the evaluation of structural integrity.

Thus, the structural integrity of the novel BSA NEDDS was well maintained and showed no obvious degradation. In the structural specificity evaluation by Western blotting (Figure 6B), the protein lanes are similar to those observed in Figure 6A, and these bands are close to the 66-kDa markers. Once again, one clear band was evident in the BSA aqueous solution (lane 10), the nanoemulsion supernatant (lane 6), and the BSA emulsion (lane 8), while the supernatant and precipitate from the blank nanoemulsion and emulsion had no obvious bands (lanes 1–4). The precipitates of the novel BSA NEDDS (lane 7) and emulsion (lane 9) lacked clear bands. Meanwhile, the results showing that the supernatant...
and precipitate of the blank nanoemulsion or emulsion (lanes 1–4) had no bands suggested that Western blotting methods can also be adapted to the evaluation of structural specificity. Overall, good structural integrity and specificity of the BSA NEDDS are demonstrated in Figure 6A and B. For example, no obvious degradation was detected in the BSA NEDDS, but the supernatant and precipitate of the blank nanoemulsion and emulsion lacked obvious bands (Figure 6A and B). The supernatant of the nanoemulsion and emulsion showed obvious, intact bands (Figure 6A and B). These results showed good immunogenicity of the novel BSA NEDDS without obvious degradation of BSA.

Integrity of the secondary structure

To correlate the small changes in the aromatic environment observed with a possible change in the secondary structure of BSA, we assessed circular dichroic spectra in the ultraviolet region, which corresponds to the protein backbone. Figure 7A–D shows the circular dichroic spectrum of the BSA aqueous solution in comparison with the circular dichroic spectrum of the BSA extracted from nanoemulsion stored at room temperature for 30, 90, and 180 days. Both spectra showed clear minima of a similar magnitude at 209 nm and 222 nm, which are indicative of the known α-helical conformation of BSA. The reduced maximum at 210 nm indicated a relative loss of the helical structure of BSA following encapsulation and extraction from this NEDDS. In addition, the ratio of α-helix, β-sheets, turns and random is not changed compared with that of the native aqueous BSA solution in Figure 8, in the secondary structure. These results suggest that BSA underwent adsorption at the interface within the emulsion system, which is commonly associated with some degree of unfolding. Importantly, the similarity characterize peak was about 210 nm and also its main secondary structure ratio of α-helix and β-sheets did not change between the native aqueous BSA solution and its NEDDS when stored at room temperature for 180 days in the circular spectra. Turn and random of BSA showed part change compared the BSA NEDDS with native aqueous BSA solution.

Integrity of the tertiary structure

The BSA stability of the tertiary structure was measured by FTIR spectroscopy. In Figure 9 and Table 1, FTIR spectra of BSA aqueous solutions showed peaks at 3445.54 cm$^{-1}$, 2073.25 cm$^{-1}$, 1651.09 cm$^{-1}$, 1543.11 cm$^{-1}$, 1455.79 cm$^{-1}$, 1403.38 cm$^{-1}$, 1308.93 cm$^{-1}$, 1245.23 cm$^{-1}$,
Figure 7 (A–D) Secondary structural spectra of the BSA in the NEDDS. (A) BSA aqueous solution; (B) BSA NEDDS stored at room temperature for 30 days; (C) BSA NEDDS stored at room temperature for 90 days; (D) BSA NEDDS stored at room temperature for 180 days.

Note: Arrows represent BSA absorption peaks.

Abbreviations: BSA, bovine serum albumin; NEDDS, nanoemulsion drug-delivery system.

1162.07 cm\(^{-1}\), 1114.44 cm\(^{-1}\), 1019.18 cm\(^{-1}\), 409.45 cm\(^{-1}\), and 401.15 cm\(^{-1}\). The BSA NEDDS stored at room temperature for 30 days showed peaks at 3445.98 cm\(^{-1}\), 1923.02 cm\(^{-1}\), 1274.41 cm\(^{-1}\), 1455.12 cm\(^{-1}\), 1519.29 cm\(^{-1}\), 1332.75 cm\(^{-1}\), 1269.24 cm\(^{-1}\), 1185.89 cm\(^{-1}\), 949.74 cm\(^{-1}\), and 880.00 cm\(^{-1}\). The characteristic band of \(
\text{O-COO (asymmetric)}\), \(\text{O-COO (symmetric)}\), and \(\text{C-O-C}\) respectively. The characteristic band of the BSA proteins at 1650 cm\(^{-1}\) and 1530 cm\(^{-1}\) are due to the \(\text{C=O stretching band of the carboxyl functional group on the}\)
tryptophan moiety as well as the carboxylate group. The FTIR spectroscopy results showed that the tertiary structure of BSA did not change in comparison with that of the BSA in the aqueous solution and the three samples stored at room temperature for 30, 90, and 180 days.

**Discussion**

In this study, we successfully developed the BSA NEDDS with a high encapsulation efficiency (>90%) and drug-loading (45 mg/mL) using the phase-inversion method and the pseudoternary phase diagram. The entrapment efficiency and drug loading of this delivery system was measured by a microbicinchoninic acid assay, which is a traditional method for measuring the protein content of a drug.27 The BSA loaded into the nanoemulsion was separated by centrifugation, and the supernatant was analyzed by a microbicinchoninic acid assay. Centrifugation is normally used to determine the stability of a delivery system after emulsion/nanoemulsion formation. Importantly, it is a standard for distinguishing between nanoemulsion and traditional emulsions. It is also the easiest and most widely used and effective method for separating molecules. We converted the nanoemulsion to an emulsion with absolute ethanol and separated BSA by high-speed centrifugation (13,000 × g, 30 minutes). For comparison, soy protein isolate emulsions and gels were centrifuged in a microcentrifuge tube at 10,000 × g for 20 minutes at room temperature.28 These results also confirmed that this delivery system is thermodynamically stable in these particular concentrations of oil, surfactant, and water, with no phase separation, creaming, or cracking.29 These results show that the stability of a protein drug can be improved greatly in our research, which we attribute to the following characteristics of this system. First, we used a novel nanoemulsion-drug delivery system, which is one of the most promising techniques for the delivery of drugs. Its advantages include higher storage stability and thermodynamic stability.30–32 We chose the nanoemulsion-drug delivery system formula to apply to a protein drug because...
it has a gel phase that forms through the preparation process. Second, we obtained the largest area of this nanoemulsion-drug delivery system by using pseudoternary diagrams to determine the best formula. Phase diagrams that provided information about the boundaries between different phases as a function of composition and other variables, such as temperature, were constructed. From this information, structural organization can be inferred. Therefore, the choice of oil and surfactant and the mixing ratio of oil to surfactant/cosurfactant are key issues in producing the nanoemulsion. The use of the phase-inversion method, a spontaneous low-emulsification-energy method also called the titration method, to prepare the BSA NEDDS is also important. Such systems can be prepared simply by mixing oil, water, surfactant, and cosurfactant in the appropriate proportions, followed by mild agitation. The order of component mixing is generally not critical, because the nanoemulsion-drug delivery system is formed spontaneously. Nevertheless, the driving forces are small, and it can take a long time for the system to reach equilibrium. Aqueous titration or spontaneous emulsification has been successfully investigated for preparing nanoemulsions of many lipophilic drugs in previous articles. Some studies on the nanoemulsion-delivery system have frequently used high-energy methods, especially homogenization. Homogenization pressure significantly influences the properties of emulsions because the shear forces and turbulence produced during homogenization, both of which are pressure-dependent, affect the particle size and size distribution. It has also been shown that disruption of any of these interactions will shift this delicate balance and destabilize the protein. The chemical and physical stability of proteins can also be compromised by environmental factors, such as pH, ionic strength, temperature, high pressure, nonaqueous solvents, metal ions, detergents, adsorption, and agitation and shearing. We chose the phase-inversion method, which is a low-energy emulsification method, to avoid destabilizing the protein drug. Finally, the BSA NEDDS formula does not include any physical reagents such as inorganic solvents (such as methylene chloride or acetone) that might destroy the protein drug, in contrast to other delivery systems. We used simpler preparation techniques compared to those used to generate microparticles, nanospheres, and nanoparticles, which may also contribute to the enhanced stability of the protein drug.

It is known that the stability of protein drugs remains one of the key hurdles to their successful entry into the market. Therefore, many factors must be considered for these drugs: structural integrity and specificity of the primary, secondary, and tertiary protein structures, relative content, and biological activity. Many traditional methods are very sensitive and useful for evaluating the structural integrity and specificity of proteins, including centrifugation, micro-BCA method, sodium dodecyl sulfate polyacrylamide gel electrophoresis, Western blotting, enzyme-linked immunosorbent assay, circular dichroic spectrometry, and FTIR spectrometry. In this study, the blank nanoemulsion or emulsion did not interfere with the identification of BSA in the nanoemulsion. The essential characteristics and thermodynamic stability of the novel BSA NEDDS were also evaluated. BSA has been widely used with many drug-delivery systems such as nanoparticles, microparticles, emulsions, gels, and liposomes because it is a common and model protein. In this study, a novel BSA NEDDS was designed and successfully prepared by the phase-inversion method and using pseudoternary phase diagrams. The results of stability tests, including assessments of particle size and zeta potential under different temperatures or pH values and the integrity of the primary, secondary, and tertiary structures of BSA, have shown that the BSA NEDDS is very stable and greatly enhances the stability of the protein drug BSA. These data will enhance the reliability of the nanoemulsion-drug delivery system for use as an oral protein drug-delivery system and change the route of drug delivery, ie, injection, and the requirement of freezing drugs. In future, we intend to study some more advanced

<table>
<thead>
<tr>
<th>Sample</th>
<th>BSA aqueous solutions stored for 30 days</th>
<th>BSA NEDDS stored for 30 days</th>
<th>BSA NEDDS stored for 90 days</th>
<th>BSA NEDDS stored for 180 days</th>
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<tbody>
<tr>
<td>Peak 1</td>
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<td>Peak 2</td>
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<td>Peak 3</td>
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<td>2929.87</td>
<td>2924.37</td>
<td>2539.36</td>
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<tr>
<td>Peak 4</td>
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<td>1643.72</td>
<td>2888.64</td>
<td>2412.35</td>
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<tr>
<td>Peak 5</td>
<td>1455.79</td>
<td>1547.08</td>
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</tr>
<tr>
<td>Peak 6</td>
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<tr>
<td>Peak 7</td>
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<td>Peak 8</td>
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<tr>
<td>Peak 9</td>
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<td>1351.77</td>
<td>1420.07</td>
<td>1455.12</td>
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<tr>
<td>Peak 10</td>
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<tr>
<td>Peak 11</td>
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<td>1269.24</td>
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<td>1103.03</td>
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<td>Peak 17</td>
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Notes: Values are expressed as wavenumbers (cm⁻¹); peak values (bold, underscore) represent BSA absorption peaks.
methods involving high-performance liquid chromatography, capillary electrophoresis, and protein drugs such as human serum albumin. Also, the stability of the BSA nanoemulsion-delivery system stored at different temperatures of 4°C, 25°C, 40°C, and 45°C and 65°C should be researched further.

Conclusion
In the present study, a NEDDS for oral administration of the standard-model protein drug BSA was successfully developed. The optimum nanoemulsion formulation was a mixture of Cremophor EL-35, propylene glycol, and isopropyl myristate at a ratio of 18:9:3 (w/w/w). The BSA NEDDS showed a narrow particle-size distribution, with a PDI value of 0.170, a spherical average particle size of 21.8 nm, and zeta potential of −24.8 mV. This nanoemulsion-drug delivery system is very stable with high encapsulation efficiency and 45 mg/mL drug loading. It also ensured good integrity of the primary, secondary, and tertiary structures; the system showed good specificity and bioactivity when stored at room temperature for 180 days. These results show that the NEDDS can greatly enhance the stability of a protein drug while retaining good integrity, specificity, and bioactivity. Thus, this delivery system can greatly enhance stability and be employed for oral administration of protein drugs.

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

References


