Alginate nanogels-based thermosensitive hydrogel to improve antidepressant-like effects of albilflorin via intranasal delivery

Dong Xu, Tao Qiao, Yue Wang, Qiang-Song Wang and Yuan-Lu Cui

State Key Laboratory of Component-based Chinese Medicine, Research Center of Traditional Chinese Medicine, Tianjin University of Traditional Chinese Medicine, Tianjin, PR China; Tianjin Key Laboratory of Biomedical Materials, Institute of Biomedical Engineering, Chinese Academy of Medical Science & Peking Union Medical College, Tianjin, PR China

ABSTRACT
Depression is a primary public health problem. However, current antidepressants work slowly, and together with side effects. Herein, the alginate nanogels were constructed to load albilflorin (albilflorin nanogels), which further formed albilflorin nanogel loaded self-assembled thermosensitive hydrogel system (albilflorin-NGSTH) and were used to improve its antidepressant effects. The nanogel showed a nano-scaled particle size and stronger antioxidant activity. Rheological studies showed that albilflorin-NGSTH had a sol-gel transition at approximately 28 °C. Albilflorin-NGSTH quickly entered the brain by intranasal delivery, and had a continuously release for albilflorin. Preliminary results of mice behavioral despair tests found that albilflorin-NGSTH had no effects on independent exploratory behavior and anxiety of the mice, and significantly decreased immobility duration of the mice in tail suspension test (TST). Moreover, the intranasally administrated albilflorin-NGSTH at a low dose improved depressive behavior, decreased levels of proinflammatory cytokines, and repaired neuronal damage of chronic unpredictable mild stress (CUMS) rats, which indicated an excellent potential for depression therapy. The treatment of albilflorin-NGSTH on depressive disorder was achieved by regulating signal pathway related to depression. Therefore, albilflorin-NGSTH has an excellent potential for clinical application in intranasal drug delivery systems.

Introduction
Depression had impacted more than 264 million of all ages in the world. Hence, it is of great clinical value and social significance to strengthen the prevention research of depression and improve the treatment effect (WHO, 2021). Nanocarriers have attracted research attention as drug delivery system since they are an excellent carrier for delivering drug to brain, and helping to attain better therapeutic effects on the central nervous system disease (Craparo et al., 2011). Particularly, there is a good strategy that the nanocarrier targets drug to brain and to enhance the antidepressant effect of drug (Chen et al., 2018). In clinical practice, the most frequently administered antidepressants had side effects and high recurrence rates (Berton & Nestler, 2006; Voican et al., 2014), and acquired about two weeks to work (Mitchell, 2006). As a result, natural polysaccharide-based drug delivery systems provided a promising strategy for drug delivery due to their inherent large surface area, biocompatibility, and low toxicity (Zhang et al., 2019; Vanavil et al., 2020).

It was reported that intranasal delivery had a rapid effect on depressive disorder by nose-to-brain route. At present, intranasal delivery as a noninvasive method in the treatment of neuropsychiatric diseases had received increasing attention, which can deliver drugs to brain and provide a best route for drugs to the brain through the olfactory pathway (Hanson & Frey, 2008; Trotta et al., 2018; Nigam et al., 2019). In 2019, esketamine, the first antidepressant of intranasal delivery, had been approved by the Food and Drug Administration (Cristea & Naudet, 2019), which had the main benefit of the rapid onset for depression. However, the major disadvantage of intranasal delivery was short residence time and drug loss in the nasal cavity because of good fluidity of the solution. At present, some hydrogels with sustained delivery had reported for anti-cancer treatment (Chen et al., 2019; Yang et al., 2020). In particular, the application of thermosensitive and biodegradable hydrogels is a good idea that avoids initial burst release and achieve sustained release of drug (Shi et al., 2021), such as an injectable thermosensitive photothermal-network hydrogel (Liu et al., 2019) and NIR-II light-modulated thermosensitive hydrogel (Ruan et al., 2019). In addition, thermosensitive hydrogel was also used for biomimaging technique or imaging of deep tissues (Chen et al., 2020; Wu et al., 2021). Thermosensitive hydrogel based on PEO-PPO-PEO poloxamers had been reported, which showed a controlled in situ release of recombinant adeno-associated viral vectors (Madry et al., 2020). Therefore, combining with the hotspot in current study, alginate nanogels based...
thermosensitive hydrogel was designed for intranasal delivery of depressive disorder.

Albiflorin as a monoterpen glycoside was the main bioactive component of the root of *Radix Paeoniae Alba* (Ranunculaceae), which might be a traditional Chinese medicine used to treat mental illness (Wang et al., 2016). Albiflorin has a variety of pharmacological activities, which included antioxidant stress (Su et al., 2013), anti-inflammatory effects (Wang et al., 2014), and neuroprotective effects (Ho et al., 2015). Modern toxicological studies showed that albiflorin had no obvious toxicity and was a safe natural drug ingredient (Han et al., 2018). At the same time, albiflorin also produced antidepressant effects though intervention of multiple targets (Song et al., 2015). Our research results also showed that albiflorin targeted hippocampal phospholipid and tryptophan metabolism for antidepressant-like effects (Wang et al., 2021). However, albiflorin maintained a low blood–brain concentration in vivo after oral administration, which was because gut microbiota transformed albiflorin to benzoic acid (Zhao et al., 2018). Furthermore, albiflorin presented a low oral bioavailability (12.09%) (TCMSP, 2015). According to the report that depression with patients had increased levels of markers of inflammation and oxidative stress (Lindqvist et al., 2017), as a result, it was a good choice that albiflorin was used to prepare nanogel-thermosensitive in situ hydrogel system that targeted albiflorin to brain and further increased its antidepressant effects.

In this study, we prepared alginate nanogels based thermosensitive hydrogel, which was used as an intranasal drug delivery system. To this aim, the alginate nanogels was constructed to load albiflorin (albiflorin nanogels) via the reverse microemulsion method. Subsequently, the albiflorin nanogels were loaded into thermosensitive hydrogels, which were further investigated in vitro and vivo (Figure 1(A)).

**Materials and methods**

**Materials**

Albiflorin (purity 98%) was obtained from Beijing Ounaer Bioengineering Technology Co., Ltd. (Beijing, China). Sodium alginate (SA, A0682, 100 g) was from Sigma-Aldrich Co. (St. Louis, MO). Span 80, Tween 80, Poloxamer 407 (P 407), and Poloxamer 188 (P 188) were from Sigma-Aldrich Co. (St. Louis, MO). Fluoxetine hydrochloride (Fluo) was bought from Eli Lilly and Company (Suzhou, China). Corticosterone ELISA Kit and Prostaglandin E2 (PGE2) ELISA Kit were obtained from Cayman Chemical Company (Ann Arbor, MI). All the other chemicals were of analytical grade.

**Animals**

Male SD rats (180–220 g) and male ICR mice (18–20 g) were purchased from the Vital River Laboratory Animal Technology Co., Ltd. (Beijing, China). Male SD rats and male ICR mice were fed at 24 ± 1 °C and relative humidity (56 ± 5%). Procedures of the animal experiment were consistent with the Guidelines for Care and Use of Experimental Animals. Experimental protocols were permitted by the Animal Ethics Committee of Tianjin University of Traditional Chinese Medicine (approval number: TCM-2018-038-E12; TCM-LAEC2021124). All experiments were reported under the Animal Research: Reporting In Vivo Experiments (ARRIVE) guidelines, which also was consistent with the recommendations for good publishing practice in physiology (du Sert et al., 2020).

**Preparation of albiflorin nanogels**

Albiflorin nanogels were synthesized with the reverse microemulsion method (Qi et al., 2015). In brief, Span 80 (1.05 mL) and Tween 80 (0.45 mL) were combined into liquid paraffin (150 mL) with stirring of 500 rpm (EURO-ST IKA, Staufen, Germany) for 30 min. Sodium alginate solution (0.5%, w/v) was obtained by dissolving the SA in deionized water with magnetism stirring for 6 h at 45 °C. In the following process, an amount of 45 mL 5A solution was added dropwise into the above oil phase with stirring of 1000 rpm for 60 min. Next, 15 mL ZnCl2 solution containing 0.5% of albiflorin (w/v) was added, and continuously stirred for 1 h to form uniform-sized droplets. Finally, the lower layer was obtained using centrifugation. Nanogels without albiflorin as control were also prepared by the above methods.

**Characterization of albiflorin nanogels**

Size and zeta potential of albiflorin nanogels were determined by particle size analyzer (NanoZS, Malvern Instruments, Malvern, UK), which was based on a method of phase-analysis light scattering (PALS). Morphology of nanogels was observed using transmission electron microscope (TEM) (H-7650, Hitachi, Tokyo, Japan). Differential scanning calorimetry (DSC) curves of albiflorin nanogels were obtained by DSC (Jade DSC, Perkin-Elmer Corp, Wilton, CT). FT-IR spectra of albiflorin nanogels were obtained by Thermo Nicolet iS10 FT-IR spectroscopy (Thermo Scientific, Waltham, MA).

**Drug loading rate of albiflorin nanogels**

Briefly, albiflorin was extracted form albiflorin nanogels by ultrasonic extraction and centrifugation. The supernatant was filtered using a filter with a pore size of 0.45 μm. The amount of albiflorin was determined using a 2695 HPLC system (Waters, NY) with a Kromasil 100-5 C18 column (250 × 4.6 mm, 5 μm, Sweden) at room temperature (25 °C). The mobile phase included water (65%) and methanol (35%), and the detection was conducted at 230 nm. Drug loading efficiency of albiflorin nanogels was calculated using the following formula:

\[
\text{Drug loading rate} (\%) = \frac{\text{amount of albiflorin in nanogels}}{\text{amount of nanogels}} \times 100\%
\]

**Antioxidant activities of albiflorin nanogels**

The DPPH radical scavenging assay was performed using the previous methods that had been reported (Rufino et al., 2009).
The test samples (including albiflorin solution and albiflorin nanogels, 3.6 mg/mL calculated by the weight of albiflorin in methanol or nanogels) and DPPH were dissolved in methanol as $A_0$. The methanol was used to replace DPPH as $A_0$. The mixture was added into the 96-well plate, and three parallel operations were done. The mixture was reacting in dark at 25°C for 15 min. The methanol and DPPH were mixed as $A_{\text{max}}$. OD values of all samples were measured at 517 nm. The DPPH radical scavenging rate was calculated as follows:

$$\text{DPPH scavenging rate}\% = \left(1 - \frac{A_{\text{max}} - A_0}{A_{\text{max}}} \right) \times 100\%$$

**Preparation and characteristic of albiflorin-NGSTH**

First, precooled normal saline and albiflorin nanogels with the settled formulation ratio were mixed with stirring of 800 rpm for 4–6 h at 4°C. Second, P 407 and P 188 were added into vial to form albiflorin-NGSTH at 4°C. In the end, albiflorin-NGSTH as a stock solution was used to the next study. The optimization of different ratio of P 407 and P 188 is reported in Table 1.

The optional formulation of albiflorin-NGSTH was screened by the tube inversion method. The intelligent thermostat metal bath was used to control temperature of the tubes containing stock solutions. Temperature of the
There are control group (normal saline, intranasal delivery), A total of 96 mice were randomly divided into eight groups. According to previous research methods (Deng et al., 2019), tissue distribution of albiflorin-NGSTH was performed on an incubator shaker with stirring of 55 rpm at 37°C. Antidepressant activities of albiflorin-NGSTH in mice behavioral despair tests was performed by the rhesometer (MCR 302, Anton-Paar, Graz, Austria). Range of the temperature was set at 20–40°C. Frequency and shear stress were adjusted to 1 Hz and 0.01 Pa, respectively. The stock solutions of 0.5 mL were detected by rhesometer. Finally, rheological properties of the stock solutions were analyzed.

Release behavior of albiflorin-NGSTH

In vitro release behavior of albiflorin-NGSTH was carried out in phosphate-buffered saline (PBS, pH 6.4). Briefly, the gel stock solution (2 mL) was added into a dialysis bag, and then suspended in PBS. The release behavior of albiflorin-NGSTH was performed on an incubator shaker with stirring of 55 rpm at 37°C (Darwin, 2019). At the specified time, the release samples of 2 mL were collected from the release medium. Meanwhile, 2 mL of fresh PBS was added into release medium. The release samples were detected using HPLC with a wavelength of 230 nm. Each batch was analyzed in triplicate.

In vivo tissue distribution of albiflorin-NGSTH

According to previous research methods (Deng et al., 2019), rhodamine B was used to label the nanogels. Herein, rhodamine B-labeled albiflorin-NGSTH was prepared by the rhodamine B-labeled albiflorin nanogels and poloxamer. In vivo distribution of the rats was detected by fluorescence imaging. Male SD rats (220–240 g) were given to rhodamine B-labeled albiflorin-NGSTH by intranasal delivery. Next, in vivo distribution (including the brain, liver, spleen, and kidney) was observed by the Maestro system (CRI, Woburn, MA) at 30, 60, 120, 240, 480, 720, and 1440 min.

Antidepressant activities of albiflorin-NGSTH in mice behavioral despair tests

A total of 96 mice were randomly divided into eight groups. There are control group (normal saline, intranasal delivery), fluoxetine group (positive drug, 10 mg/kg, gavage), albiflorin solution group (7 mg/kg, gavage, i.g.), albiflorin solution group (1.75 mg/kg, intravenous (i.v.) injection), albiflorin nanogel (0.35 mg/kg calculated by the weight of albiflorin in albiflorin nanogels, intranasal delivery), and albiflorin-NGSTH (0.175, 0.35, and 0.7 mg/kg calculated by the weight of albiflorin in albiflorin nanogels, intranasal delivery). According to our previous methods, the exploratory behavior and locomotor activity of the mice were investigated by behavioral equipment (Multifunctional 256,060 Series, TSE Systems, Bad Homburg vor der H öhe, Germany), which included number of rearings, total distance, and activity time of the mice. Tail suspension test (TST) of the mice was further recorded and analyzed by the above behavioral equipment (Xu et al., 2020, 2021).

Establishment of chronic unpredictable mild stress (CUMS) model

CUMS rats model was based on a previous method with slight modification (Willner et al., 1987). Experimental rats were randomly divided into control group (normal saline, intranasal delivery), CUMS group (normal saline, intranasal delivery), fluoxetine group (positive drug, 10 mg/kg, i.g.), albiflorin solution group (7 mg/kg, i.g.), albiflorin solution group (1.75 mg/kg, i.v.), and albiflorin-NGSTH (0.175, 0.35, and 0.7 mg/kg calculated by the weight of albiflorin in albiflorin nanogels, intranasal delivery). All rats underwent CUMS protocol for 5 weeks, except for the control group. Experimental rats were administrated in the last seven days of the CUMS protocol. The CUMS protocol included multiple stimuli, namely, swimming in ice water, food, and water deprivation within 24 h, 10 min of light stimulation, electrical stimulation (36 mV, 10 sec each time), 2 min of oscillation and tail clip. The weight change and sugar water preference of the rats were recorded during the CUMS protocol.

Open field test (OFT) of CUMS model

After the CUMS protocol and the administration were performed, locomotor activity of the rats was evaluated using the OFT. Behavioral characteristics (including total distances, number of rearings, activity time) were observed and recorded using animal behavioral video analysis system (Multifunctional 256060 Series, TSE Systems, Bad Homburg vor der H öhe, Germany).

Biochemical indicators and pathological changes of rats

All rats were sacrificed after the OFT. Plasma of the rats were obtained by centrifuging at 3000×g for 10 min. Corticosterone and prostaglandin E₂ (PGE₂) of the rats in the plasma were detected using enzyme-linked immunosorbent assay (ELISA). The hippocampi of the rats were obtained from the brain, and 10% neutral formalin was used to fix the hippocampi. Hematoxylin and eosin (H&E) were used to stain tissue after the fixed tissue undergo a series of treatments that included dehydration, embeddedness, and sectioning.

### Table 1. Optimization of different ratio of P 407 and P 188.

| Batches | Albiflorin nanogel (w/w, %) | P 407 (w/w, %) | P 188 (w/w, %) | Normal saline (w/w, %) |
|---------|---------------------------|---------------|----------------|-----------------------|
| 1       | 12                        | 14            | 0              | 74                    |
| 2       | 12                        | 14            | 0              | 73                    |
| 3       | 12                        | 14            | 2              | 72                    |
| 4       | 12                        | 16            | 0              | 72                    |
| 5       | 12                        | 16            | 1              | 71                    |
| 6       | 12                        | 16            | 1              | 70                    |
| 7       | 12                        | 18            | 0              | 69                    |
| 8       | 12                        | 18            | 1              | 68                    |
| 9       | 12                        | 18            | 2              | 68                    |

stock solutions were altered at an increase of 1°C/min. Gelling temperature of the stock solutions was detected using tube inversion method. The temperature that the solutions kept immobile was recorded, as the gelation temperature. Each batch was analyzed in triplicate.

The rheological analysis of the stock solutions was performed by the rhesometer (MCR 302, Anton-Paar, Graz, Austria). Range of the temperature was set at 20–40°C. Frequency and shear stress were adjusted to 1 Hz and 0.01 Pa, respectively. The stock solutions of 0.5 mL were detected by rhesometer. Finally, rheological properties of the stock solutions were analyzed.
RNA-seq of hippocampus transcriptome

Total RNA was obtained from the hippocampi of rats using Trizol reagent kit according to the manufacturer's instructions. RNA degradation and contamination were monitored on 1% agarose gels and its purity was checked using the Nano Photometer spectrophotometer (IMPLEN, Westlake Village, CA). RNA concentration was measured by Qubit RNA Assay Kit in Qubit 2.0 Flurometer (Life Technologies, Carlsbad, CA), and its integrity was determined by the RNA Nano 6000 Assay Kit of the Bioanalyzer 2100 system (Agilent Technologies, Santa Clara, CA). Sequencing libraries were generated using NEBNext Ultra RNA Library Prep Kit for Illumina (NEB, Ipswich, MA) following manufacturer’s recommendations and index codes were added to attribute sequences to each sample. At last, PCR products were purified (AMPure XP system) and library quality was assessed on the Agilent Bioanalyzer 2100 system. The cDNA libraries were sequenced on an Illumina HiSeq platform according to standard sequencing protocol. Differential expression analysis of albiflorin-NGSTH and CUMS group was performed using the DESeq2 R package (1.16.1). The resulting p values were adjusted using the Benjamini and Hochberg’s approach for controlling the false discovery rate. Genes with an adjusted p value <.05 found by DESeq2 were assigned as differentially expressed. Gene Ontology (GO) enrichment analysis of differentially expressed genes was implemented by the cluster Profiler R package, in which gene length bias was corrected. GO terms with corrected p value less than .05 were considered significantly enriched by differentially expressed genes. The cluster Profiler R package was used to test the statistical enrichment of differential expression genes in KEGG pathways.

Statistical analysis

All data were analyzed using Origin Pro 2021 (OriginLab, Northampton, MA). Data were expressed as the means ± standard error (mean ± SE). One-way analysis of variance (ANOVA) was used to analyze between multiple groups. A p < .05 or p < .01 was considered as statistically significant difference.

Results and discussion

Preparation and characterization of albiflorin nanogels

Preparation of albiflorin nanogels is shown in Figure 1(A). Results of size and zeta potential are demonstrated in Figure 1(B,C), respectively. The size distribution of nanogels was randomly measured, and the obtained data represented that albiflorin nanogels and blank nanogels were 45.6 ± 5.2 nm and 49.4 ± 0.0 nm, respectively. Compared to blank nanogels, the particle size of albiflorin nanogels was reduced, which might be due to a greater intermolecular force that albiflorin participated in the mechanism of the nanogels by hydrogen bonding. Polydispersity index (PDI) that described the particle size distribution of the nanogels was less than 0.20, which indicated that albiflorin nanogels had a narrow particle size distribution (Willner et al., 1987). In terms of zeta potential, surface charge density shows the electric potential (Sikora et al., 2016). Both blank nanogels and albiflorin nanogels were negatively charged with zeta potential of about −10.0 ± 0.3 mV and −19.8 ± 0.9 mV, respectively. Therefore, the absolute value of zeta potential was increased by addition of albiflorin, which suggested that albiflorin might improve the stability of nanogels by a stronger intermolecular force. At the same time, albiflorin nanogels displayed a pale blue color under the LED lights. TEM microphotographs presented that the nanogel was almost subspherical shape (Figure 1(D)). The experimental results demonstrated that content of albiflorin in alginate nanogels was 2.43% by HPLC analysis, which was suitable for intranasal delivery of depression therapy.

The physical status of SA, albiflorin, SA-albiflorin-ZnCl₂, blank nanogels, and albiflorin nanogels was examined by DSC (Figure 1(E)) and FTIR (Figure 1(F)). Albiflorin nanogels did not show a characteristic peak of albiflorin on the FTIR compared to free albiflorin. Results of the DSC showed that albiflorin nanogels, compared to albiflorin, did not appear an endothermic peak or exothermic peak of albiflorin. The above curves indicated that albiflorin might participate in the structure of alginate nanogels, which might include physical inclusion and hydrogen bonding. Nevertheless, because of the low drug loading rate of alginate nanogels and the weak force between the compounds, some characteristic peaks such as the hydrogen bond of alginate and albiflorin were not be found on the infrared spectrum.

Antioxidant activities of albiflorin nanogels are shown in Figure 1(G). It was observed that, compared to albiflorin solution, albiflorin nanogels had a higher DPPH scavenging rate, and presented strongly antioxidant activities on free radicals. It was reported that Zn²⁺ had been selected as a cross-linker cation because of its antioxidant properties, and its supplementation was considered as an antioxidant to protect cells (DiSilvestro, 2000; Szuster-Ciesielska et al., 2009; Varghese et al., 2009). Sodium alginate, isolated from brown algae, also exhibited important antioxidant activity (Sellimi et al., 2015). Therefore, the increase of antioxidant activities of albiflorin nanogels may be related to Zn²⁺, SA, and zinc alginate.

Preparation and characterization of albiflorin-NGSTH

The preparation process of albiflorin-NGSTH is shown in Figure 2(A). After the addition of poloxamer, albiflorin-NGSTH presented a temperature-sensitive property under the condition of low temperature and high temperature. This was because polyoxypropylene oxide, as hydrophobic units of poloxamer, produced dehydration and aggregation, so formed a dense and ordered grid. Poloxamer tightly encapsulated the albiflorin-NGSTH in the interior. Meanwhile, the polyoxyethylene oxide shell also rapidly generated gelation in vivo. Therefore, poloxamer sustained release of albiflorin
and increased the local drug concentration by self-assembly characteristic.

Initially, the optimal formulation that was adapt to intranasal delivery was screened by the tube inversion method. It was observed that with the addition of P 407, the gelation temperature decreased gradually, and P 188 was the opposite in Figure 2(B). When the ratio of P 407 and P 188 was close to 16/2, the gelling temperature (30.5°C) conforms to intranasal administration. Gelling temperature of the optional formulation was further confirmed by rheometer (Figure 2(C)). The point of intersection of storage modulus (G') and loss modulus (G'') curve was as the final gelation temperature. It was found that the dramatic increase of the G' and G'' at approximately 28°C indicated gelling temperature of albiflorin-NGSTH. Furthermore, the viscosity of the albiflorin-NGSTH also was increased significantly by the change of temperature, which indicated that albiflorin-NGSTH had mucoadhesive properties (Figure 2(D)).

In vitro drug release of albiflorin in albiflorin-NGSTH is shown in Figure 2(E). The release results showed that the accumulative release rate of albiflorin from albiflorin-NGSTH could reach near 100% in the first 12 hours in the PBS (pH 6.4). The release behavior of albiflorin-NGSTH could last for 12 hours, implying that albiflorin was well encapsulated into the nanogels. Furthermore, albiflorin-NGSTH may prolong the retention time of albiflorin in the brain, and increase the potential neuroprotection time for the focal area of brain. Meanwhile, different release mechanisms were proposed for albiflorin-NGSTH releasing in PBS. Based on the values of $R^2$ in Table 2, the release process of albiflorin-NGSTH most likely fits to the Higuchi and Ritger–Peppas equation, which suggested that the release mechanism of albiflorin-NGSTH not

Table 2. Models and equations of albiflorin releasing from albiflorin-NGSTH in PBS.

| Model          | Equation                | $R^2$  |
|----------------|-------------------------|--------|
| Zero-order     | $Q = 7.36t + 20.93$     | 0.9414 |
| First-order    | $Q = 98.01 \left(1 - e^{-0.27t}\right)$ | 0.9731 |
| Higuchi        | $Q = 29.55t^{1/2} - 0.40$ | 0.9943 |
| Ritger–Peppas  | $Q = 29.32t^{0.5}$      | 0.9943 |

Figure 2. Preparation and characterization of albiflorin-NGSTH. (A) The preparation process of albiflorin-NGSTH. (B) Gelling temperature of the formulations. (C) Rheological analysis of the optimal prescription (P 407/P 188, 16/2). (D) Viscosity of the optimal prescription. (E) In vivo release of albiflorin-NGSTH.
only presented a sustained and controlled property for albiflorin, but might be due to drug diffusion governing the drug release.

**Fluorescence distribution of albiflorin-NGSTH**

*In vivo* distribution and fluorescence intensity analysis of rhodamine B-labeled albiflorin-NGSTH in rats are shown in Figure 3(A,B), respectively. It was observed that rhodamine B-labeled albiflorin-NGSTH was intranasally administered after 30 min; the fluorescence intensity of the brain was the most obvious compared to later time. The above results showed that nanogels can fast penetrate the brain by nose-to-brain route and predominantly accumulate in the brain. Liver and kidney also showed stronger fluorescence, which indicated that most of the drugs were absorbed into the blood through the nasal mucosa and further distributed to the liver and kidneys. By 1440 min, brain of the rat presented a weak fluorescence, which indicated that albiflorin-NGSTH exhibited...
a long and sustained release for albiflorin. At the same time, fluorescence gradually disappeared at the other organs. Due to smaller size of nanogels, large quantities of albiflorin nanogels accumulated in the liver and were eventually excreted by the kidneys. Therefore, these results confirmed the rapidly antidepressant effects of albiflorin-NGSTH, and it can continuously release albiflorin.

**Antidepressant activities of albiflorin-NGSTH in mice behavioral despair tests**

In order to evaluate whether albiflorin-NGSTH effects on independent exploratory behavior and anxiety of the mice, open field test (OFT) was performed in open field apparatus. It is observed in Figure 4(A–C) that each administration group, compared to control group, had no significant difference. These results showed that albiflorin and its preparation had no effects on independent exploratory behavior and anxiety of the mice.

As shown in Figure 4(D), antidepressant activities of albiflorin-NGSTH were further evaluated by TST, a method that screened and assessed new potential antidepressant drugs (Willner, 1984). It was found that the intranasally administrated albiflorin-NGSTH was lower than that of fluoxetine solution and albiflorin solution administrated by intragastric or i.v. administration, but albiflorin-NGSTH significantly decreased immobility duration of the mice in TST and reversed behavioral despair of the mice. Compared to albiflorin nanogel, albiflorin-NGSTH showed the lower immobility duration, which indicated that self-assembled thermosensitive hydrogel system helped to increase antidepressant activities of albiflorin nanogel. These experimental results demonstrated superiority of albiflorin-NGSTH administrated by nasal cavity compared to oral administration and necessity of self-assembled thermosensitive hydrogel system on the nanogel that was used to intranasal delivery. Therefore,
antidepressant effects of albiflorin-NGSTH will be mainly studied in the further experiment.

**Pharmacodynamic study of albiflorin-NGSTH on CUMS rats**

CUMS model was widely used as the study of depressive behavior (Gronli et al., 2005; Nirmale et al., 2008). CUMS model is shown in Figure 5. In the experiment, CUMS significantly decreased the body weight of rats (Figure 6(A)), growth rate of body weight (Figure 6(B)), and sucrose preference of rats (Figure 6(C)). However, albiflorin-NGSTH reversed the above adverse effects. In the behavior test of rats, albiflorin-NGSTH increased low number of rearing (Figure 6(D)), low activity time (Figure 6(E)), and short total distance (Figure 6(F)) induced by CUMS model. However, albiflorin-NGSTH was less than the dose of the other group. These results further indicated the advantage of albiflorin-NGSTH.

Meanwhile, some studies had demonstrated that depressive patients had an abnormal of inflammation and HPA axis (Druzhkova et al., 2019), and these abnormal changes were improved by antidepressant drugs (Wang et al., 2011). In this study, PGE_2 and corticosterone as indicators of proinflammatory cytokines and HPA axis respectively were reversed using albiflorin-NGSTH. Albiflorin-NGSTH decreased corticosterone levels (Figure 6(G)) and PGE_2 levels (Figure 6(H)). Albiflorin-NGSTH, compared to the oral albiflorin solution, better reversed corticosterone and PGE_2 levels. Compared to i.v. albiflorin, intranasally administrated albiflorin-NGSTH suggested an advantage of delivery route. To sum up, albiflorin-NGSTH can increase body weight and sucrose preference of rats, improve abnormal behavior of CUMS model, and reserve high plasma levels of corticosterone and PGE_2 of CUMS model. Experimental results showed that albiflorin-NGSTH, a low dose, had good antidepressant effects on the CUMS model.

**Repairing effect of albiflorin-NGSTH on hippocampus injury of CUMS model**

The hippocampus is considered an important region for the development of depression, which was also related to the growth of learning and memory (Fasick et al., 2015). It was observed that neuronal atrophy of the hippocampus showed the dysfunction of cellular processes, which were consistent with characteristics of clinically depressed populations (Morales-Medina et al., 2017). Morphological changes of the neurons were detected using H&E staining to evaluate the antidepressant effect of preparations. Results of staining in the hippocampus are indicated in Figure 7(A,B). It was observed that there was a significant change of pyramidal cells. Compared to the CUMS model, pyramidal cells of albiflorin-NGSTH in CA1 and CA3 subregion showed a regular and tidier cell arrangement, which was close to normal cells. Although intragastric administration of fluoxetine and
albiflorin, and tail i.v. injection of albiflorin presented a similar effect in the treatment of CUMS model, but the dose of albiflorin-NGSTH was very small. Therefore, albiflorin-NGSTH improved the morphology of neuronal cells, which presented a better antidepressant effect at a low dose in the CUMS model.

Transcriptome studies of albiflorin-NGSTH on hippocampus of CUMS model

Differentially expressed genes were shown in the volcano map, which included the comparison between CUMS model and control group (Figure 8(A)), the comparison between albiflorin-NGSTH and CUMS model (Figure 8(B)), and the comparison between fluoxetine and CUMS model (Figure 8(C)). It was observed that CUMS significantly regulated the differential expressed genes of control group, and caused a depressive behavior of rats. Differentially expressed genes were prominently regulated by administration of albiflorin-NGSTH and fluoxetine. Albiflorin-NGSTH had a widely regulation for differential expressed genes according to the results of volcano map and Venn diagram (Figure 8(D)).

Gene Ontology is an international standard classification system for gene function, and differentially expressed genes were classified according to cellular component, molecular function, and biological process. KEGG is also the classic public database of pathways (Fang et al., 2020). The above methods were performed using gene function analysis and pathway analysis in the present study. The distribution of target genes in enriched GO functions in the albiflorin-NGSTH vs. CUMS model is shown in Figure 8(E). KEGG enrichment analysis was performed to screen for a significantly enriched KEGG pathway (Figure 8(F)). From the point of view of biological pathway, the specific pathway was found with significant difference in expression between groups through the enrichment analysis of differential expressed genes. Enrichment analysis of GO and KEGG showed that differentially expressed genes induced by CUMS were involved in the regulation of multiple neural pathways, including cAMP signal pathway, calcium ion signal pathway and cGMP PKG signal pathway. It was reported that cAMP signal pathway, calcium ion signal pathway and cGMP PKG signal pathway were related to the incidence of depression (Armstrong et al., 2009; Bergantin, 2020). Therefore, the results of pharmacodynamics and transcriptomics showed albiflorin-NGSTH had a good antidepressant effect on depressive disorder through the regulation of cAMP signal pathway, calcium ion signal pathway, and cGMP PKG signal pathway.

Conclusions

In this study, alginate nanogel was prepared using reverse microemulsion method. Next, alginate nanogel loaded with albiflorin was wrapped into thermosensitive hydrogel for
intranasal delivery of albiflorin. The results showed that the prepared albiflorin nanogel met the characteristics of nano-preparation and presented stronger antioxidant activities. The gelling temperature of albiflorin-NGSTH conformed to intranasal delivery. Experimental results showed that albiflorin-NGSTH can adhere to the nasal mucosa for continuous release of albiflorin and rapidly enter to the brain in the first 30 min. Results of mice behavioral despair tests showed that albiflorin-NGSTH had no effects on independent exploratory behavior and anxiety of the mice, and significantly decreased immobility duration of the mice in TST. Compared to other administration group, albiflorin-NGSTH at a low dose effectively improved the abnormal behavior and biochemical indicators of CUMS model, and showed a good antidepressant efficacy and preparation superiority. Antidepressant effects of albiflorin-NGSTH were mainly achieved by regulating oxidative phosphorylation, cAMP, calcium ion, and cGMP PKG pathway.

Author contributions
Yuan-Lu Cui and Qiang-Song Wang contributed to the study concepts and design. Dong Xu, Tao Qiao, and Yue Wang...
Disclosure statement
The authors report no conflicts of interest in this work.

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References
Armstrong GA, Rodgers CJ, Money TG, et al. (2009). Suppression of spreading depression-like events in locusts by inhibition of the NO/cGMP/PKG pathway. J Neurosci 29:8225–35.
Bergantin LB. (2020). Depression rises the risk of hypertension incidence: discussing the link through the Ca$^{2+}$/cAMP signalling. Curr Hypertens Rev 16:73–8.
Berton O, Nestler EJ. (2006). New approaches to antidepressant drug discovery: beyond monoamines. Nat Rev Neurosci 7:137–51.
Chen BY, Luo M, Liang JM, et al. (2018). Surface modification of PPG for a neutrophil-nanoparticle co-vehicle to enhance the anti-depressant effect of baicalein. Acta Pharm Sin B 8:64–73.
Chen X, Wang M, Yang X, et al. (2019). Injectable hydrogels for the sustained delivery of a HER2-targeted antibody for preventing local relapse of HER2$^{+}$ breast cancer after breast-conserving surgery. Theranostics 9:6080–98.
Chen X, Zhang J, Wu K, et al. (2020). Visualizing the in vivo evolution of an injectable and thermosensitive hydrogel using tri-modal bioimaging. Small Methods 4:2000310.
Crupero EF, Bondi ML, Pitarresi G, et al. (2011). Nanoparticulate systems for drug delivery and targeting to the central nervous system. CNS Neurosci Ther 17:670–7.
Cristea IA, Naudet F. (2019). US Food and Drug Administration approval of esketamine and brexanolone. Lancet Psychiatry 6:975–7.
Darwin BD. (2019). Alkaline treatment for preventing acidosis in the rumen culture fermenting carbohydrates: an experimental study in vitro. J Adv Vet Anim Res 6:100–7.
Deng XQ, Zhang HB, Wang GF, et al. (2019). Colon-specific microspheres loaded with puerarin reduce tumorigenesis and metastasis in colitis-associated colorectal cancer. Int J Pharm 570:118644.
DiSilvestro RA. (2000). Zinc in relation to diabetes and oxidative disease. J Nutr 130:1509S–1515S.
Druzhkova T, Pochigaeva K, Yakovlev A, et al. (2019). Acute stress treatment for preventing acidosis in the rumen culture fermenting carbohydrates: an experimental study in vitro. J Adv Vet Anim Res 6:100–7.
Deng XQ, Zhang HB, Wang GF, et al. (2019). Colon-specific microspheres loaded with puerarin reduce tumorigenesis and metastasis in colitis-associated colorectal cancer. Int J Pharm 570:118644.
DiSilvestro RA. (2000). Zinc in relation to diabetes and oxidative disease. J Nutr 130:1509S–1515S.
Druzhkova T, Pochigaeva K, Yakovlev A, et al. (2019). Acute stress response to a cognitive task in patients with major depressive disorder: potential metabolic and proinflammatory biomarkers. Metab Brain Dis 34:621–9.
du Sert NP, Hurst V, Apliuwalia A, et al. (2020). The ARRIVE guidelines 2.0: updated guidelines for reporting animal research. J Cereb Blood Flow Metab 40:1769–77.
Fang K, Xu JX, Chen XX, et al. (2020). Differential serum exosome microRNA profile in a stress-induced depression rat model. J Affect Disord 274:144–58.
Fasick V, Spengler RN, Samankan S, et al. (2015). The hippocampus and TNF: common links between chronic pain and depression. Neurosci Biobehav Rev 53:139–59.
Gronli J, Marison R, Fiske E, et al. (2005). Effects of chronic mild stress on sexual behavior, locomotor activity and consumption of sucrose and saccharine solutions. Physiol Behav 84:571–7.
Han J, Xia Y, Lin L, et al. (2018). Next-generation metabolomics in the development of new antidepressants: using albiflorin as an example. Curr Pharm Des 24:2530–40.
Hanson LR, Frey WH. (2008). Intranasal delivery bypasses the blood–brain barrier to target therapeutic agents to the central nervous system and treat neurodegenerative disease. BMC Neurosci 9:1–4.
Ho SL, Poon CY, Lin C, et al. (2015). Inhibition of β-amyloid aggregation by albiflorin, aloemodin and neohesperidin and their neuroprotective effect on primary hippocampal cells against β-amyloid induced toxicity. Curr Alzheimer Res 12:424–33.
Lindqvist D, Dhabhar FS, James SJ, et al. (2017). Oxidative stress, inflammation and treatment response in major depression. Psychoneuroendocrinology 76:197–205.
Liu CJ, Guo XL, Ruan CP, et al. (2019). An injectable thermosensitive photothermal-network hydrogel for near-infrared-triggered drug delivery and synergistic photothermal-chemotherapy. Acta Biomater 96:281–94.
Madry H, Gao L, Rey-Rico A, et al. (2020). Thermosensitive hydrogel based on PEO–PPG–PEO poloxamers for a controlled in situ release of recombinant adeno-associated viral vectors for effective gene therapy of cartilage defects. Adv Mater 32:1906508.
Mitchell AJ. (2006). Two-week delay in onset of action of antidepressants: new evidence. Br J Psychiatry 188:105–6.
Morales-Medina JC, Iannitti T, Freeman A, et al. (2017). The olfactory bulbectomized rat as a model of depression: the hippocampal pathway. Behav Brain Res 317:562–75.
Nigam K, Kaur A, Tyagi A, et al. (2019). Nose-to-brain delivery of lamotrigine-loaded PLGA nanoparticles. Drug Deliv Transl Res 9:879–90.
Nirmal J, Babu CS, Harisudhan T, et al. (2008). Evaluation of behavioural and antioxidant activity of Cytisus scoparius Link in rats exposed to chronic unpredictable mild stress. BMC Complement Altern Med 8:15–8.
Qi Y, Jiang M, Cui YL, et al. (2015). Synthesis of quercetin loaded nanoparticles based on alginate for Pb(II) adsorption in aqueous solution. Nanoscale Res Lett 10:408.
Ruan CP, Liu CJ, Hu HL, et al. (2019). NIR-II light-modulated thermosensitive hydrogel for light-triggered cisplatin release and repeatable chemo-photothermal therapy. Chem Sci 10:4699–706.
Rufino MS, Fernandes FA, Alves RE, et al. (2009). Free radical-scavenging behaviour of some north-east Brazilian fruits in a DPPH system. Food Chem 114:693–5.
Sellimi S, Younes I, Ben Ayed H, et al. (2015). Structural, physicochemical and antioxidant properties of sodium alginate isolated from a Tunisian brown seaweed. Int J Biol Macromol 72:1358–67.
Shi J, Yu L, Ding J. (2021). PEG-based thermosensitive and biodegradable hydrogels. Acta Biomater 21:00246–46.
Sinogawa K, Sharg AD, Minelli C. (2016). Size and γ-potential measurement of silica nanoparticles in serum using tunable resistive pulse sensing. Langmuir 32:2216–24.
Song J, Hou X, Hu X, et al. (2015). Not only serotonergic system, but also dopaminergic system involved in albiflorin against chronic unpredictable mild stress-induced depression-like behavior in rats. Chem Biol Interact 242:211–7.
Suh KS, Choi EM, Lee YS, et al. (2013). Protective effect of albiflorin against oxidative-stress-mediated toxicity in osteblast-like MC3T3-E1 cells. Fitoterapia 89:33–41.
Szuster-Ciesielska A, Plewka K, Daniluk J, et al. (2009). Zinc supplementation attenuates ethanol- and acetdehyde-induced liver stellate cell activation by inhibiting reactive oxygen species (ROS) production and by influencing intracellular signaling. Biochem Pharmacol 78:301–14.
TCSMP. (2012). Traditional Chinese Medicine systems pharmacology database and analysis platform. Available at: http://tcmssp.com/molecule.php [last accessed 6 Apr 2021].
Trotta V, Pavan B, Ferraro L, et al. (2018). Brain targeting of resveratrol and antioxidant properties of sodium alginate isolated from a Sargassum swartzii by nasal administration of chitosan-coated lipid microparticles. Eur J Pharm Biopharm 127:250–9.
Vanavil B, Selvaraj K, Aanandhalakshmi R, et al. (2020). Bioactive and thermoreversible sulphated polysaccharide from Sargassum wartzii with drug delivery applications. Int J Biol Macromol 153:190–200.
Varghese J, Faith M, Jacob M. (2009). Zinc prevents indomethacin-induced renal damage in rats by ameliorating oxidative stress and mitochondrial dysfunction. Eur J Pharmacol 614:114–21.

Voican CS, Corruble E, Naveau S, et al. (2014). Antidepressant-induced liver injury: a review for clinicians. Am J Psychiatry 171:404–15.

Wang QS, Gao T, Cui YL, et al. (2014). Comparative studies of paeonoflorin and albiflorin from *Paeonia lactiflora* on anti-inflammatory activities. Pharm Biol 52:1189–95.

Wang QS, Yan K, Li KD, et al. (2021). Targeting hippocampal phospholipid and tryptophan metabolism for antidepressant-like effects of albiflorin. Phytomedicine 153735.

Wang Y, Yang F, Liu YF, et al. (2011). Acetylsalicylic acid as an augmentation agent in fluoxetine treatment resistant depressive rats. Neurosci Lett 499:74–9.

Wang YL, Wang JX, Hu XX, et al. (2016). Antidepressant-like effects of albiflorin extracted from *Radix Paeoniae Alba*. J Ethnopharmacol 179: 9–15.

WHO. (2021). World Health Organization. Available at: https://www.who.int/health-topics/depression [last accessed 6 Apr 2021].

Willner P. (1984). The validity of animal models of depression. Psychopharmacology 83:1–16.

Willner P, Towell A, Sampson D, et al. (1987). Reduction of sucrose preference by chronic unpredictable mild stress, and its restoration by a tricyclic antidepressant. Psychopharmacology 93:358–64.

Wu X, Wang X, Chen X, et al. (2021). Injectable and thermosensitive hydrogels mediating a universal macromolecular contrast agent with radiopacity for noninvasive imaging of deep tissues. Bioact Mater 6: 4717–28.

Xu D, Lu Y-R, Kou N, et al. (2020). Intranasal delivery of icariin via a nanogel-thermosensitive hydrogel compound system to improve its antidepressant-like activity. Int J Pharm 586:119550.

Xu D, Qiu C, Wang Y, et al. (2021). Intranasal co-delivery of berberine and evodiamine by self-assembled thermosensitive in-situ hydrogels for improving depressive disorder. Int J Pharm 603:120667.

Yang X, Chen X, Wang Y, et al. (2020). Sustained release of lipophilic gemcitabine from an injectable polymeric hydrogel for synergistically enhancing tumor chemoradiotherapy. Chem Eng J 396:125320.

Zhang YH, Zhang YM, Yu J, et al. (2019). Boronate-crosslinked polysaccharide conjugates for pH-responsive and targeted drug delivery. Chem Commun 55:1164–7.

Zhao ZX, Fu J, Ma SR, et al. (2018). Gut-brain axis metabolic pathway regulates antidepressant efficacy of albiflorin. Theranostics 8:5945–59.