Drug Release from Gelsolin-Targeted Phase-Transition Nanoparticles Triggered by Low-Intensity Focused Ultrasound

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**Purpose:** Current strategies for tumour-induced sentinel lymph node detection and metastasis therapy have limitations. It is essential to identify and provide warnings earlier for tumour metastasis to carry out effective clinical interventions. In addition, traditional cancer chemotherapy encounters drastic limitations due to the nonspecific delivery of antitumour drugs and severe side effects. We aimed to exploit the potential of gelsolin (GSN) monoclonal antibody as a targeting agent and perfluorohexane (PFH) as a phase-transition agent to maximize the cytotoxic effect of poly(lactic-co-glycolic acid) (PLGA) nanoparticle-based drug controllable release systems for Hca-F cells.

**Methods:** We co-encapsulated PFH and doxorubicin (DOX) into PLGA nanoparticles (NPs) and further conjugated GSN monoclonal antibody onto the surface of NPs to form GSN-targeted phase transition polymer NPs (GSN-PLGA-PFH-DOX) for both imaging and therapy of tumours and metastatic lymph nodes. To promote and trigger drug release on demand, low-intensity focused ultrasound (LIFU) was applied to achieve a controllable release of the encapsulated drug.

**Results:** GSN-PLGA-PFH-DOX NPs exhibited characteristics such as a narrow size distribution and smooth surface. GSN-PLGA-PFH-DOX NPs could also specifically bind to Hca-F cells and increase the ultrasound contrast agent (UCA) image contrast intensity. GSN-PLGA-PFH-DOX NPs enable GSN-mediated targeting and biotherapeutic effects as well as LIFU-responsive drug release, resulting in synergistic cytotoxic effects in GSN-overexpressing cells in vitro.

**Conclusion:** Our work might provide a strategy for the imaging and chemotherapy of primary tumours and their metastases.

**Keywords:** poly(lactic-co-glycolic acid), gelsolin, phase transition, controllable drug release

**Introduction**

Traditional cancer chemotherapy encounters drastic limitations due to the nonspecific delivery of antitumour drugs and severe side effects.\(^1–3\) Therefore, the design and fabrication of novel drug delivery systems are promising and rapidly developing areas. Microbubbles, known to be UCAs, have played important roles in the diagnosis, differential diagnosis and treatment of benign and malignant tumours. In addition, they have been used to load therapeutic agents for efficient tumour chemotherapy.\(^4–6\) However, these gas-filled microbubbles have a limited capacity for loading therapeutic agents, short circulation time, and large micrometre size, making it difficult to achieve effective drug concentration at the tumour site, which
hampers their wide application in drug delivery. It has been reported that only particles with a diameter of <700 nm are permitted to pass through the vascular endothelial space of tumours. To address these problems, different kinds of nanoscale drug-delivery systems with selective targeting and controlled release properties have been designed and fabricated to minimize systemic cytotoxicity. The stability of nanodroplets is increased, and they can accumulate in tumour tissues by passive targeting due to their nanoscale size.

Some synthetic polymers, which are composed of one or two monomers and demonstrate controllable degradation rates, have been widely used for drug delivery applications. PLGA is a degradable polyester polymer that is randomly polymerized by two monomers, lactic acid and glycolic acid. It has been certified by the FDA in the United States and has been formally included in the United States Pharmacopoeia as a medicinal excipient. It has good biocompatibility, encapsulation and film-forming properties and ultimately metabolizes into water and carbon dioxide in vivo and is thus called a “good environmental material”. It is widely used in pharmaceuticals, medical engineering materials and modern industrial fields. PLGA NPs can protect the biological activity of drugs and enhance their tolerance to external conditions (acids, alkalis, enzymes, water, heat, etc.) and prolong the half-lives of biological drugs in the body, which has important clinical significance for long-term drug users. Meanwhile, to enhance the accuracy of drug delivery to tumours, controlled drug-delivery systems have been functionalized with a variety of tumour-specific targeting ligands—such as aptamers, peptides, antibodies and growth factors—to selectively target molecules and receptors that are overexpressed on the surface of tumour cells.

The lymphatic system plays an important role in the spread of cancer cells. Tumour metastasis to the lymphatic system is a universal sign of tumour progression and is closely related to the prognosis of patients. Once tumour cells invade the lymphatic system, they can take root and seed into other regions of the body by these invaded lymphatic nodes. Studies have revealed that GSN plays important roles in cell adhesion, migration and invasion. Differential proteomics has revealed that GSN is closely associated with the lymphatic metastasis of tumours and is mainly located in the cytoplasm and cytoskeleton of the mouse hepatocarcinoma cell line Hca-P with low metastatic potential. Meanwhile, it also appears in the cell membrane of the mouse hepatocarcinoma cell line Hca-F, which has a high metastatic potential. It is a kind of skeleton-regulating protein that regulates polymerization and depolymerization of cell actin in a Ca-dependent manner and thus affects cell morphology and motility. Abbeele et al demonstrated that the downregulation of gelsolin tends to decrease the invasion and migration of many tumour cells, such as cervical HeLa cells and MDA-MB-231 cells, in human breast cancer. Hence, it may be an important membrane antigen in the early stages of the invasion and metastasis of tumour cells and is expected to become a molecular target for early identification and prediction of lymphatic metastasis. Nevertheless, the existing imaging diagnosis mainly depends on the morphology of tumours and characteristics of blood perfusion but is unable to provide specifications of the biological behaviours of tumours, which leads to obvious limitations in the early identification of tumour metastasis. Therefore, designing a molecular targeted contrast agent for specific imaging of tumour antigens will be helpful for early detection, accurate diagnosis and precise treatment of tumour metastasis.

Despite the great efforts to design and produce novel drug delivery systems, drug efficacy in treating tumours remains limited by the relatively small amount of drug delivery and release in available systems. In addition, at present, most of the developed PLGA contrast agents are nanoscale, and their imaging ability does not seem very optimistic in in vivo experiments. Recent studies have demonstrated that PFH holds great potential in imaging-based approaches via a “liquid-to-gas” phase transition in the tumour region. In contrast to conventional UCA microbubbles, PFH droplets are generally enclosed in liposomes, PLGA, or albumin shells to enhance their stability and avoid air embolism at the beginning of administration. Furthermore, the small dimensions of PFH-containing NPs enhance their ability to permeate the internal regions of a tumour and elude capture by the mononuclear phagocyte system to further prolong their circulation in the bloodstream. Meanwhile, when PFH-containing NPs are triggered by ultrasound (US) to enter the gaseous phase, they possess significantly augmented US contrast due to their volume expansion. Therefore, to develop a controllable drug delivery system with the simultaneous ability to realize a molecular imaging diagnosis of early metastasis of tumours, we fabricated drug-encapsulated GSN-targeted phase-transition PLGA nanoparticles using the double emulsion-solvent evaporation
technique and carbodiimide method. In the NPs we produced, the phase-transition material PFH and an anticancer drug (DOX) were selected to be incorporated in the PLGA shell. In addition, we investigated the influence of different volumes of PFH on drug encapsulation efficiency. PFH in the core of NPs can undergo a liquid-gas phase transition upon activation by US energy, called acoustic droplet vaporization. When converged in tumor sites, these NPs convert into microbubbles under LIFU irradiation. These acoustic microbubbles oscillate and collapse by the ultrasound-targeted microbubble destruction (UTMD) effect.\(^5\),\(^3\),\(^2\),\(^3\) During this process, the anticancer drug encapsulated in bubbles is released in tumor cells and possesses significantly augmented US contrast.

**Experimental Section**

**Materials and Instruments**

PLGA (lactide: glycolide = 50:50, \(M_w = 12,000\) Da) was purchased from Jinan Daigang Biological Material Limited Company (Shandong, China). PFH was purchased from J&K Scientific Ltd. (Beijing, China). DOX was obtained from Beijing Yihe Biological Co. Ltd. (Beijing, China). GSN monoclonal antibodies, poly(vinyl alcohol) obtained from Beijing Yihe Biological Co. Ltd. (Beijing, China). DOX was purchased from J&K Scientific Ltd. (Beijing, China). GSN monoclonal antibodies, poly(vinyl alcohol) obtained from Beijing Yihe Biological Co. Ltd. (Beijing, China). DOX was purchased from J&K Scientific Ltd. (Beijing, China). GSN monoclonal antibodies, poly(vinyl alcohol) obtained from Beijing Yihe Biological Co. Ltd. (Beijing, China). DOX was purchased from J&K Scientific Ltd. (Beijing, China). GSN monoclonal antibodies, poly(vinyl alcohol) obtained from Beijing Yihe Biological Co. Ltd. (Beijing, China). DOX was purchased from J&K Scientific Ltd. (Beijing, China). GSN monoclonal antibodies, poly(vinyl alcohol) obtained from Beijing Yihe Biological Co. Ltd. (Beijing, China). DOX was purchased from J&K Scientific Ltd. (Beijing, China). GSN monoclonal antibodies, poly(vinyl alcohol) obtained from Beijing Yihe Biological Co. Ltd. (Beijing, China). DOX was purchased from J&K Scientific Ltd. (Beijing, China).

**Characteristics of the NPs and Determination of DOX Loading**

First, 200 \(\mu\)L of GSN-PLGA-PFH-DOX NPs was diluted in double-distilled water to 10 \(\mu\)L, and then 1 drop was deposited on a 200-mesh copper grid. When the sample was dry, the morphology of these NPs was studied using a scanning electron microscope (SEM, Hitachi S-3400N, Japan), and the structure was analysed using transmission electron microscopy (TEM, Hitachi H-7600, Japan). In addition, the size distribution and surface zeta potential of each NP type were assessed using a Malvern Zetasizer Nano ZS unit (Malvern Instrument, UK).

Drug loading and encapsulation efficiency were determined using a high-performance liquid chromatography (HPLC) method. The amount of DOX recovered from washed supernatants during preparation \(W_k\) was determined. The drug payload \(W_{NP}\) was calculated as the difference between the total amount of DOX \(W_T\) and
The drug encapsulation efficiency and loading efficiency were calculated by the following equations:

\[
\text{Encapsulation efficiency} = \frac{W_{NP}}{W_T} \times 100\%
\]

\[
\text{Loading efficiency} = \frac{W_{NP}}{W_t} \times 100\%
\]

where \(W_T\) represents the total weight of DOX used in the preparation of the NPs; \(W_{NP}\) represents the total drug amount in the NPs; and \(W_t\) represents the total weight of NPs.

**Connection of GSN Monoclonal Antibody with NPs**

Dil-labelled GSN-PLGA-PFH-DOX was collected and diluted to 2 mg/mL with double-distilled water, and then 20 μL of fluorescein isothiocyanate (FITC)-labelled goat-anti-mouse IgG antibody was added and oscillated for 2 h in an ice bath, followed by observations under a laser scanning confocal microscope (LSCM, Olympus FV1200, Japan). Considering PLGA-PFH-DOX as the control, the GSN monoclonal antibody connection rate on the surface of GSN-PLGA-PFH-DOX was detected by flow cytometry.

**In vitro Liquid-Gas Phase Transition and Responsive Drug Release Activated by LIFU**

Two millilitres of GSN-PLGA-PFH-DOX NP solution (1 mg/mL) was added to the preformed wells of an agarose gel phantom (3% agarose). Then, a LIFU transducer (1 MHz, acoustic intensity: 1.2 W/cm², duty cycle: 50%, Chongqing Medical University, P. R. China) was used to irradiate the NPs with a power of 1–8 W and a duration of 1–5 min. B-mode US imaging signals and contrast-enhanced ultrasound (CEUS) imaging signals of the post-formed microbubbles from NPs were observed in real time using a MyLab 90 ultrasound diagnostic system (Esaote, Italy). The average ultrasound intensity was determined by DFY software (Chongqing Medical University, P. R. China), and the change in size of NPs was observed under an optical microscope (OM, Olympus BX51-P, Japan).

To measure the drug release patterns, we collected two types of drug-loaded NPs, divided them into four groups and incubated them in conical flasks with 20 mL of pre-warmed PBS (120 rpm/min) at 37 °C. For the GSN-PLGA-DOX+LIFU and GSN-PLGA-PFH-DOX+LIFU groups, LIFU irradiation (8 W, 3 min) was applied to activate drug release at a predetermined time point (8 h). One millilitre of release medium was removed for measurement at 0.5, 1, 2, 3, 4, 6, 8, 10, 12, 24, 48, and 72 h. Then, the same amount of fresh buffer was added to maintain the sink condition. HPLC was used to determine the content of DOX in these samples, calculate the cumulative release rate of the drug, and draw a release curve.

**Cellular Uptake of NPs**

Hca-F cells in logarithmic phase were seeded into confocal culture dishes with 1640 complete medium (without foetal bovine serum) for 4 h. They were divided into 3 groups: Hca-F cells+ PLGA-PFH-DOX, Hca-F cells+ GSN monoclonal antibody+ GSN-PLGA-PFH-DOX (receptor blockade group), and Hca-F cells+ GSN-PLGA-PFH-DOX. First, 100 μL of GSN monoclonal antibody was added to the receptor blockade group and incubated for 20 min. After that, 100 μL of coumarin-6-labelled corresponding NP sample was added, and the cells were incubated at 37 °C and 5% CO₂ for 2 h. Subsequently, cells were fixed with 4% paraformaldehyde and stained fluorescently with DAPI after washing with PBS. Intracellular uptake of NPs was investigated under LSCM. Image-Pro Plus 6.0 software (IPP 6.0, Media Cybernetics, USA) was used to measure the green fluorescence intensity in the field of vision and count the number of cells. The fluorescence intensity of each cell was obtained by dividing the two. Each group took three fields of vision.

**Cytotoxicity Assay, Cell Apoptosis and Half Inhibition Concentration**

Hca-F cells were seeded into 96-well plates at a density of 1×10⁴ per well and cultured overnight. Free DOX at a concentration of 20 μg·mL⁻¹ and GSN-PLGA-PFH NPs, PLGA-PFH-DOX NPs, and GSN-PLGA-PFH-DOX NPs at a concentration of 300 μg·mL⁻¹ were prepared in double-free medium. To investigate the cytotoxicity of NPs, eleven groups were formed, namely, the empty group, control group, and experimental groups: GSN-PLGA-PFH NPs, PLGA-PFH-DOX NPs, GSN-PLGA-PFH-DOX NPs, DOX, control + LIFU, GSN-PLGA-PFH NPs + LIFU, PLGA-PFH-DOX NPs + LIFU, GSN-PLGA-PFH-DOX NPs + LIFU, and DOX + LIFU groups. Then, 0.1 mL of medium was added to the control group when 0.1 mL of NPs or free DOX was added to the other groups. Then, these cells were incubated for 24 h at 37 °C and 5% CO₂. For those groups with US irradiation, LIFU irradiation (8 W, 3 min) was performed after 2 h incubation.
Then, the CCK-8 method was used to detect the cell survival rate of each group.

To investigate the half inhibition concentration (IC50) of NPs, 0.1 mL of GSN-PLGA-PFH-DOX NPs at different concentrations (0.025, 0.05, 0.1, 0.25, 0.5 and 1.0 mg·mL$^{-1}$) was added to Hca-F cells when the control group was added to 0.1 mL medium. Then, these cells were incubated for 24 h at 37 °C and 5% CO$_2$. LIFU irradiation (8 W, 3 min) was performed after 2 h of incubation. Then, the CCK8 method was used to detect the OD value of each group, and the cell inhibition rate was calculated.

Cell viability rate (%) = \( \frac{(OD_1 - OD_0)}{(OD_2 - OD_0)} \times 100\%
\)

Cell inhibition rate (%) = \( \frac{(OD_2 - OD_1)}{(OD_2 - OD_0)} \times 100\%
\)

where OD$_0$ represents the OD value of the empty group; OD$_1$ represents the OD value of the experimental group; and OD$_2$ represents the OD value of the control group.

Two-colour flow cytometric analysis was used to investigate cellular apoptosis. The experiment was grouped as above, and Hca-F cells were collected at the end of the incubation. Cells were washed with PBS and stained with reagents from the Apoptosis Assay Kit (Bestbio Co. Ltd, P. R. China) following the manufacturer’s instructions and analysed using a flow cytometer (FACSVantage SE, BD Biosciences, US).

**Results and Discussion**

**Design, Synthesis and Characterization of GSN-PLGA-PFH-DOX**

The fabrication of GSN-PLGA-PFH-DOX NPs is elucidated in Figure 1. First, we constructed PLGA NPs encapsulating PFH and DOX. Then, the GSN monoclonal antibody was coupled to the surface of the PLGA-PFH-DOX NPs via the carbodiimide method, forming GSN-PLGA-PFH-DOX NPs for US imaging and chemotherapy applications. GSN receptor-specific endocytosis of NPs is also demonstrated in Figure 1. When GSN-PLGA-PFH-DOX NPs were endocytosed into Hca-F cells, LIFU was applied to trigger the liquid-gas phase transition of PFH, thus achieving an enhanced ultrasound imaging effect and on-demand drug release.

The GSN-PLGA-PFH-DOX NPs appeared as a milky-white mixed suspension. SEM imaging of these NPs (Figure 2A) showed that GSN-PLGA-PFH-DOX NPs exhibited a smooth and uniform spherical morphology. The presence of PFH and DOX in the shells of NPs was demonstrated by the enhanced contrast manifested as dark domains in Figure 2B by TEM. The size distribution of GSN-PLGA-
PFH-DOX NPs was between 200 and 300 nm (mean diameter = 285.62± 12.76 nm), and the polydispersity index was PI = 0.141±0.052. The mean diameters of pure PLGA NPs, PLGA-PFH NPs, PLGA-DOX NPs and PLGA-PFH-DOX NPs were 248.35±13.23 nm (PI= 0.133±0.053), 259.17±16.45 nm (PI= 0.136±0.027), 259.46±13.71 nm (PI=0.364±0.075) and 260.51±13.71 nm (PI=0.263±0.043), respectively (Figure 2C and D). These results suggest that the addition of PFH and DOX in the centre does not significantly affect the NP size distributions. The particle sizes of GSN-PLGA-PFH-DOX NPs compared with other NPs were all P<0.05, and the difference was statistically significant. These results indicate that the connection of GSN monoclonal antibody on the surface of the NPs is the main reason for the increase in particle size. Figure 2E shows that when the NPs carry GSN monoclonal antibody, the negative charge decreases gradually and tends to the positive potential (1.68±0.53 mV). This is similar to the results of previous studies, which showed that the GSN monoclonal antibody was successfully connected with NPs.

The DOX loading amount and encapsulation efficiency were measured by HPLC. The encapsulation efficiencies of DOX in GSN-PLGA-PFH-DOX with different volumes of PFH (25, 50, 100, 200, 350, 500 μL) were 67.56±3.21%, 66.44±4.06%, 67.92±2.55%, 68.45±3.43%, 67.16±2.66%, and 68.22±3.14%, respectively. The DOX loading efficiencies of these NPs were 6.76±0.32%, 6.64±0.41%, 6.79±0.26%, 6.85±0.34%, 6.72±0.27%, and 6.82±0.31%, respectively. From these results, we can deduce that the addition of PFH to the polymer shell does not significantly affect the DOX encapsulation efficiency or the loading efficiency (Figure 2F). Comparatively, the encapsulation and loading efficiencies of PLGA-PFH-DOX were 68.06±4.63% and 6.81±0.46%, respectively. It can be inferred that the modification of GSN onto the surface of PLGA nanoparticles does not affect the DOX encapsulation or loading efficiency in some respect. The stability of the NPs could be maintained when stored at 4 °C, and the particle size and surface potential of the NPs had no significant change when tested at room temperature within 7 days (Figure 2G and H).
Connection of GSN Monoclonal Antibody

The carbodiimide method, as a classic chemical method for connecting proteins and small peptide molecules, is currently the most popular and widely used method for the production of biochemical conjugates. This method has high target-linking efficiency and mild reaction conditions. EDC is a water-soluble crosslinking agent that allows biological coupling between carboxyl and amine groups. In this reaction, the amino group from the GSN monoclonal antibody forms an amide bond with the carboxyl group on the PLGA surface. Figure 3A showed that GSN-PLGA-PFH-DOX marked by DiI fluorescent dye emitted red fluorescence, Figure 3B showed FITC-labeled secondary antibody emitted green fluorescence (FITC-labeled secondary antibodies can bind to GSN monoclonal antibodies attached to the surface of NPs), and Figure 3C showed orange fluorescence after the two fluorescence channels were superimposed. This result shows that the GSN monoclonal antibody was successfully attached to the surface of NPs. Flow cytometry revealed that the connection rate of GSN monoclonal antibody and NPs was 98.31% (Figure 3D), which was higher than that in the control group (0.073%) (P<0.05) (Figure 3E). These results show that the GSN monoclonal antibody was successfully connected, laying the foundation for subsequent targeted experiments.

In vitro Phase Transition and Drug Release Activated by LIFU

Ultrasound-mediated drug delivery NPs can be monitored by US imaging and induced to release therapeutic agents using external US radiation after they are detected in the target tissues by diagnostic US. Liquid PFH encapsulated in NPs changed to the gas phase under LIFU irradiation. Subsequently, larger microbubbles are formed and exploded, which leads drugs to be released from drug delivery systems. We selected adequate irradiation power and duration of LIFU to induce the phase transition of PFH incorporated in NPs. Using an LIFU transducer, the US energy was focused onto the NPs. DFY software was applied to evaluate the value of the average US signal intensity. GSN-PLGA-PFH-DOX NPs had the highest average B-mode and CEUS signal (Figure 4A–C) values at US parameters of 8 W/3 min, suggesting that a large number of PLGA NPs were converted to microbubbles.
With the increase in irradiation duration, some PLGA microbubbles were destroyed and collapsed, decreasing the intensities of harmonic US images, which also laid the foundation for subsequent drug release therapy. In addition, LIFU irradiation is the focus, making the process of triggering the phase change both efficiently and safely. Therefore, a US irradiation parameter of 8 W/3 min was selected to activate the phase transition of PLGA NPs and release the loaded drug after bubble collapse. Following conversion of NPs into bubbles, tumour imaging signals could be collected, and the drug could be released after the collapse of the microbubbles. During the process of the LIFU-responsive phase transition, we observed the variations in PFH-loaded NPs every 1 min via OM. The size of the NPs increased approximately 200-fold from their original size (Figure 4D).

To investigate drug-releasing behaviour under LIFU irradiation, four groups were formed, namely, GSN-PLGA-DOX (without LIFU radiation), GSN-PLGA-DOX +LIFU radiation (8 W, 3 min), GSN-PLGA-PFH-DOX (without LIFU radiation), and GSN-PLGA-PFH-DOX +LIFU radiation (8 W, 3 min). As displayed in Figure 4E, DOX was continuously released from the NPs without LIFU radiation, but burst release of the drug was observed in the NPs after LIFU exposure. For GSN-PLGA-PFH-DOX group, drugs were released slowly from the NPs within 72 h, and no obvious burst drug-releasing profiles were observed because liquid PFH did not change to the gas phase without external stimulus. Although the mechanism of US-triggered drug release is not yet fully understood, it has been reported\(^{37}\) that the liquid-gas phase transition of PFH under US radiation causes the gaps between the shell polymer molecules to become larger, which may facilitate the escape of drug molecules from the bubbles.
In vitro Targeted Ability

The observation results by LSCM for the in vitro uptake experiment are shown in Figure 5A. More GSN-PLGA-PFH-DOX NPs were taken up by Hca-F cells, which resulted in strong cell fluorescence (1853.64±153.69 a.u.) (Figure 5B). When the target GSN on the surface of Hca-F cells was sealed, a markedly decreased amount of GSN-PLGA-PFH-DOX NPs was absorbed, which resulted in weak cell fluorescence (768.33±151.42 a.u.). Meanwhile, fewer PLGA-PFH-DOX NPs were unconnected to the GSN monoclonal antibody and were absorbed by Hca-F cells (701.63±181.53 a.u.).

Cytotoxicity Assay, Cell Apoptosis and Half Inhibition Concentration

In our study, the drug loading of GSN-PLGA-PFH-DOX was 6.85±0.34%; therefore, the amount of DOX in 300 μg/mL NPs was equivalent to that in the free DOX group (20 μg/mL). To investigate the cytotoxicity and cell apoptosis efficiency of NPs, various groups of agents were incubated with Hca-F cells for 24 h. As shown in Figure 5C and D, compared with the control group, the survival rates of Hca-F cells in the GSN-PLGA-PFH and GSN-PLGA-PFH+LIFU groups were not significantly decreased, and the cell apoptosis rates were not significantly increased, which indicated that the cytotoxicity of nondrug-loaded PLGA NPs to cells was very low. The cytotoxicity levels of the PLGA-PFH-DOX NP and GSN-PLGA-PFH-DOX NP groups were lower than that of the free DOX group. Comparatively, the cell viability rates of groups treated with PLGA-PFH-DOX NPs, GSN-PLGA-PFH-DOX NPs and free DOX were significantly decreased after LIFU irradiation. Meanwhile, the cell apoptosis rates of these three groups were significantly increased. The reason may be that the mechanical and cavitation effects of LIFU irradiation increase the permeability of the cell membrane, which increases the amount of nanoparticles or drugs entering the cells, thereby increasing the therapeutic effect. In addition, before the

![Figure 5](https://example.com/figure5.png)

Figure 5 (A) Cell uptake observed by LSCM (Coumarin-6: green, DAPI: blue) (Scale bar=50 μm). (B) Histogram of the quantitative study of the cellular uptake efficiencies of different groups (**P<0.001 vs the other two groups). (C) Cell viability rates of various therapeutic agents, *P<0.05. (D) Cell apoptosis rate of various therapeutic agents, *P<0.05. (E) Fitting curve of half inhibition concentration of NPs.
application of combined LIFU irradiation, the cytotoxicity of GSN-PLGA-PFH-DOX NPs was less than that of free DOX, indicating that the DOX contained in GSN-PLGA-PFH-DOX NPs was relatively stable without significant burst release. After combined LIFU irradiation, the cytotoxicity of GSN-PLGA-PFH-DOX NPs was significantly higher than that of free DOX (P<0.05), indicating that targeted delivery of drugs was increased under the targeted effect of GSN monoclonal antibody. Meanwhile, the blasting effect of the NPs and the release of DOX after combined LIFU irradiation further enhanced the cytotoxicity of NPs. The results of the half inhibition concentration experiment are shown in Figure 5E. When the concentration of GSN-PLGA-PFH-DOX NPs was 0.1 mg·mL⁻¹, the inhibition rate of Hca-F cells was 29.11±2.14%, and when the concentration of NPs was 0.25 mg·mL⁻¹, the cell inhibition rate increased to 65.03±2.35%. The IC50 of NPs calculated by GraphPad Prism 9 software (GraphPad Software Inc., San Diego, CA, USA) was 0.17 mg·mL⁻¹.

Conclusion
In summary, we successfully developed gelsolin-targeted and doxorubicin-loaded phase transition nanoparticles (GSN-PLGA-PFH-DOX NPs) as a targeted drug carrier and an efficient UCA for both the imaging and therapy of tumours and metastatic lymph nodes. The GSN-PLGA-PFH-DOX NPs exhibited efficient targeted capability to tumour tissues in vitro and correspondingly enhanced US imaging. Over 72 h of incubation, DOX was slowly and steadily released after the initial burst release. Furthermore, US stimulation can accelerate drug release on demand. LIFU stimulation, causing a controllable release of encapsulated drug from the GSN-PLGA-PFH-DOX NPs, may allow cytotoxic drugs to accumulate at higher levels in tumours, which can maximally kill Hca-F cells with fewer side effects and prevent tumour recurrence and metastasis. The combination of targeted imaging and anticancer drug transportation based on GSN-PLGA-PFH-DOX NPs as the contrast agent for ultrasonography could be attractive in the future development of nanomedicine and noninvasive medicine for simultaneous accurate diagnosis, drug delivery and therapy for cancers or other diseases.

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

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