Inhibition of prostate cancer growth using doxorubicin assisted by ultrasound-targeted nanobubble destruction

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Abstract: Ultrasound (US)-targeted microbubble destruction has been widely used as an effective drug-delivery system. However, nanobubbles (NBs) have better stability and stronger penetration than microbubbles, and drug delivery assisted by US-targeted NB destruction (UTND) still needs to be investigated. Our aim was to investigate the effect of doxorubicin (DOX) on the inhibition of prostate cancer growth under UTND. Contrast-enhanced US imaging of transplanted PC3 prostate cancer in mice showed that under a combination of 1 W/cm² US power and a 100 Hz intermittent pulse with a “5 seconds on, 5 seconds off” mode, NBs with an average size of (485.7±33) nm were effectively destroyed within 15 minutes in the tumor location. PC3 cells and 20 tumor-bearing mice were divided into four groups: a DOX group, a DOX + NB group, a DOX + US group, and a DOX + NB + US group. The cell growth-inhibition rate and DOX concentration of xenografts in the DOX + NB + US group were highest. Based on another control group and these four groups, another 25 tumor-bearing mice were used to observe the treatment effect of nine DOX injections under UTND. The xenografts in the DOX + NB + US group decreased more obviously and had more cellular apoptosis than other groups. Finally, electron microscopy was used to estimate the cavitation effect of NBs under US irradiation in the control group, NB group, US group, and NB + US group. The results of scanning electron microscopy showed that PC3 cells in the DOX + NB + US group had more holes and significantly increased cell-surface folds. Meanwhile, transmission electron microscopy confirmed that more lanthanum nitrate particles entered the parenchymal cells in xenografts in the NB + US group compared with the other groups. This study suggested that UTND technology could be an effective method to promote drugs to function in US-irradiated sites, and the underlying mechanism may be associated with a cavitation effect.

Keywords: ultrasound therapy, nanobubble, cavitation effect, prostate cancer

Introduction
Chemotherapy and gene therapy have become important means of tumor therapy. However, drugs/genes can only exert their therapeutic effects after reaching target tissues or target organs; otherwise, the side effects produced may be harmful to the human body. For the majority of chemotherapeutic drugs and gene drugs, identifying the proper drug/gene-delivery system to maintain high concentrations of the drug/gene in specific target tissues and exert the best therapeutic effects is difficult. Viral vectors, represented by adenoviruses and lentiviruses, have very high transduction efficiency; however, their potential infection and immunogenicity risks have not yet been effectively solved. Although liposome transfection plays an important role in vitro cell studies, its transfection efficiency is low and its price is high. Therefore, their
applications are just used in the experiments. Ultrasound (US)-targeted microbubble destruction (UTMD) technology can specifically destroy US contrast agents (UCAs) in US-irradiated sites, and play an important role in the targeted delivery of drugs/genes.\(^{1,2}\) Currently, this technology is widely studied and explored in a variety of diseases, mainly cardiovascular diseases, eye diseases, and tumors.\(^{3–7}\)

However, previous studies in this field focused mainly on micron-level UCAs; there are few studies on the recently developed nanoscale UCAs. Nanoscale UCAs are contrast agents with a particle diameter <1,000 nm, and can be used for imaging under contrast-enhanced US (CEUS) mode. Compared to microscale UCAs, nanoscale UCAs have features of strong penetration ability and excellent stability; therefore, they have been studied extensively, especially in applications in solid-tumor imaging.\(^{8,9}\) Nanoscale UCAs can be distributed more extensively in tumor tissues using enhanced permeation and retention (EPR) effects: large vascular endothelial gaps, incomplete basement membranes, and imperfect lymphatic drainage.\(^{10}\) In our previous studies, we constructed nanobubbles (NBs) carrying small interfering RNA against the androgen receptor. The NBs that carried androgen-receptor small interfering RNA were destroyed by US irradiation, and experiments on in vitro prostate cancer cells and xenograft tumors in animals were done. The results showed that this technique had good inhibitory effects on tumors.\(^{11,12}\) However, chemotherapeutic drugs assisted by US-targeted NB destruction (UTND) for the treatment of tumors and the underlying mechanism still require further investigation. Therefore, we applied lipid NBs and the classic broad-spectrum chemotherapeutic drug doxorubicin (DOX) to treat prostate cancer using UTND in this study. Next, we further studied the underlying mechanism using electron microscopy in order to provide a research basis for the inhibition of tumor growth by chemotherapeutic drugs using UTND.

**Materials and methods**

**Preparation of ultrasound NBs**

The lipid components 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC; Avanti Polar Lipids, Alabaster, AL, USA), 1,2-dipalmitoyl-sn-glycero-3-phosphatidic acid (DPPA; Corden Pharma, Liestal, Switzerland), 1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine (DPPE; Corden Pharma), 1,2-dipalmitoyl-sn-glycero-3-phosphoglycerol (DPPG; Corden Pharma), and polyethylene glycol 2000, were dissolved in a hydration solution containing glycercin and phosphate-buffered saline (PBS) at a ratio of 1:9 (v:v) and incubated overnight in a horizontal shaker at 37°C. The solution was transferred to a penicillin bottle, the air in the head of the bottle replaced with perfluoropropane gas ($C_3F_8$; Research Institute of Physical and Chemical Engineering of Nuclear Industry, Tianjin, People’s Republic of China), and the bottle shaken using the horizontal reciprocating mechanical oscillation method with an operating frequency greater than 4,500 rpm for 80 seconds using an ST-B series amalgamator (AT&M Biomaterials Co, Beijing, People’s Republic of China). The obtained suspension was centrifuged sequentially under conditions of 300×g and 300 rpm for 3 minutes to discard the lipid materials that were not encapsulated into NBs and microbubbles in the top layer.

After the separated NBs were negatively stained with a phosphotungstic acid solution, the size and morphology of the NBs were observed using transmission electron microscopy (TEM; JEM-1400; JEOL, Tokyo, Japan) and under the bright field of laser-scanning confocal microscopy (Zeiss 780; Carl Zeiss AG, Oberkochen, Germany), respectively. The particle sizes of the fabricated NBs were measured in a Malvern Zetasizer Nano ZS90 detector (Malvern Instruments, Malvern, UK) according to the principles of dynamic light scattering after they had been diluted with PBS.

**In vitro cell culture and the establishment of the tumor-xenograft animal model**

The PC3 prostate cancer cell line from the American Type Culture Collection was cultured in 1640 culture medium containing 10% fetal bovine serum and 1% penicillin and streptomycin in a 5% CO\(_2\) and 37°C incubator. During the establishment of tumor xenografts into 4- to 5-week-old BALB/c-nu nude mice (HFK Bioscience Co, Beijing, People’s Republic of China), 200 μL of tumor cells in logarithmic phase at a concentration of 2.5×10\(^6\)/mL were mixed with 200 μL of Matrigel (BD Biosciences, San Jose, CA, USA) and subcutaneously injected into nude mice. The mice were continuously housed in a clean environment. The animal experiments in this study were approved by the Institutional Animal Ethics Committee of the Third Military Medical University, and performed in accordance with the International Guiding Principles for Biomedical Research involving Animals-1985.

**In vitro cytotoxicity**

In vitro cytotoxic activity of DOX (Solarbio, Beijing, People’s Republic of China) against PC3 cells was evaluated by using the Cell Counting Kit (CCK)-8 assay (Dojindo Laboratories, Kumamoto, Japan). Briefly, cells (5×10\(^4\) cells/well) were seeded into 96-well plates and cultured for 12 hours to adhere at the bottom of the plates. Then, different concentrations
of DOX, 0–50 µM, were added and further incubated for 8 hours. After that, the medium was changed and replaced with fresh medium containing CCK-8 solution, and cells were further incubated for 1 hour. Finally, the absorbance of the sample taken from each well was measured at 450 nm in a Varioskan Flash multimode reader (Thermo Fisher Scientific, Waltham, MA, USA), on the basis of which the percentage of surviving cells at every concentration was calculated relative to the untreated one. After the optimal concentration of DOX against PC3 cells had been determined, the cells were divided into four groups: the DOX-only group, the DOX combined with NB (DOX + NB) group, DOX under US irradiation (DOX + US) group, and DOX combined with NB under US irradiation (DOX + NB + US) group. In the relevant groups, the intervention was done according to the following settings: the optimal dose of DOX as aforementioned (10 µM DOX), NBs 5×10⁷, US power 1 W/cm², and irradiation time 20 seconds in WED-100 US therapeutic equipment (Well D Medical Electronics Co Ltd, Shenzhen, People’s Republic of China). A similar procedure with CCK-8 assay was used to calculate cell viability in each treatment group. All experiments were performed in triplicate.

Measurement of DOX concentrations in all tissues and organs of nude mice

First, US power and time required for US burst were confirmed in nude mice. Injection of 5×10⁷ NBs was performed through the retro-orbital venous sinus of tumor-bearing mice. The WED-100 US therapeutic equipment was used to irradiate tumor regions from the side of tumor xenografts. A sufficient amount of coupling agent to fill the region between the probe and the tumor xenografts was used. The frequency of the therapeutic probe was 1 MHz. Irradiation of tumor regions was performed using US power of 1, 1.75, and 2.5 W/cm². To ensure reperfusion of the tumor regions after NB destruction, the built-in 100 Hz pulse of the therapeutic equipment combined with the manual “5 seconds on, 5 seconds off” mode was used. During the process of US irradiation, an L12-5 probe scanning from the upper part of xenografts was initiated in superficial CEUS mode of an iU22 US diagnostic apparatus (Philips, Amsterdam, the Netherlands), which is used for real-time monitoring of the perfusion of NBs in tumor regions under the condition of 0.12 mechanic index until no NBs were observed inside the tumor xenografts. This period of time was considered the duration of US irradiation of NBs.

Next, 50 µg/mL DOX was prepared using double-distilled water, and whole-spectrum absorbance was measured at 200–1,000 nm in a multimode reader. After the optimal measuring wavelength was determined, standard curves of DOX at a concentration range of 1–50 µg/mL were generated. A total of 20 tumor-bearing nude mice with tumor sizes of ~1 cm were randomly divided into four groups as described in the “In vitro cytotoxicity” section. The intervention was performed according to the following settings: injection dose of DOX 0.1 mg, NBs 5×10⁷, and therapeutic US condition performed in tumor-xenograft regions using the aforementioned irradiation parameters.

Finally, the nude mice were killed 6 hours after the end of the experimental treatment. Livers, implanted tumors, kidneys, hearts, lungs, and spleens of the nude mice of each group were collected and fully minced for US tissue homogenization. DOX was extracted from the tissue supernatant using chloroform–methanol extraction. Supernatants of these tissues were collected from normal tumor-bearing mice and added to a series of DOX with known concentrations for the same manipulation. Absorbance was measured, and standard curves were generated to calculate the drug concentrations in all the organs.

Treatment for tumor xenografts in nude mice

Another 25 nude mice were randomly grouped, with five animals in each group. On the basis of the previous four groups, a blank control group was added. Two CEUS images of tumors were examined with a dosage of 1×10⁷ NBs 2 days before treatment and 2 days after the end of treatment. The maximum intensities of signals were extracted using QLab8.1 offline software (Philips) for comparison. The specific treatment procedures were strictly carried out as described in the following content. When the diameter of the tumor xenograft was 5 mm, treatment was initiated. The treatment was performed nine times every other day. The US parameters in the corresponding groups were the same as the above. The amount of injected DOX was 0.05 mg, and the amount of injected NBs was 5×10⁷. The tumor volume and the mass of the nude mice on treatment days 0, 6, 12, and 18 were recorded. The tumor volume was calculated as $V = \frac{4}{3}\pi r^3$ and the tumor mass as $M = \rho V$. Mice were killed by cervical dislocation 2 days after the last treatment. Tumor tissues and myocardial tissues were collected. The tumor xenografts were weighed, and tumor density was calculated as $\rho = M/V$.

Assessment of tumor-xenograft and myocardial tissue sections

Heart and tumor xenografts of the nude mice in the five groups were washed, fixed in 4% paraformaldehyde, embedded in
paraffin, sectioned, and stained with hematoxylin–eosin (H&E). In addition, the apoptosis conditions of cells in myocardial tissue and tumor-xenograft sections were analyzed according to the instruction manual of the terminal deoxynucleotidyl transferase deoxyuridine triphosphate nick-end labeling (TUNEL) assay reagent kit (Beyotime, Shanghai, People’s Republic of China). According to the principle of the TUNEL method, terminal deoxynucleotidyl transferase functions to transfer peroxidase-modified deoxyribonucleotides to the 3′-OH terminus of DNA broken during cell apoptosis, allowing apoptosis to be observed in situ.

### Analysis of biological effects under UTND using electron microscopy

First, the effects of NBs on surface morphology under US irradiation were observed at the cell level. PC3 prostate cancer cells (1×10⁶) were inoculated onto 24-well plates. The cells were cultured overnight to allow them to attach to the surface of coverslips. The cells were washed with PBS and divided into four groups: control, NB, US, and NB + US. The number of NBs in each well was 1×10⁴. The wells were filled with PBS, and irradiation was performed for 20 seconds from the top of cell wells using the aforementioned US therapeutic probe. Next, the cells were rapidly washed with PBS, fixed in 2.5% glutaraldehyde for 2 hours, and dehydrated for replacement using different gradients of alcohol (50%, 75%, 95%, and 100%) via the retro-orbital venous sinus. The US parameters to transfer peroxidase-modified deoxyribonucleotides to the 3′-OH terminus of DNA broken during cell apoptosis, allowing apoptosis to be observed in situ.

#### Statistical analysis

Histograms and line charts were generated using GraphPad Prism 6.0 (GraphPad Software, La Jolla, CA, USA). Statistical analysis was performed using SPSS 22.0 software (IBM Corporation, Armonk, NY, USA). Data are presented as mean ± standard deviation. The drug concentrations in all the organs and the data derived from evaluating the treatment in all the groups were analyzed using a one-way analysis of variance. *P<0.05* indicated that a difference had statistical significance.

### Results

#### In vitro observation of NBs

The prepared NBs appeared to be in milky suspension. After centrifugation and stratification treatment, the obtained NBs exhibited a dark-black round region encapsulated in a light-black circular structure with homogeneous size distribution under TEM (Figure 1A). Under the bright field of laser-scanning confocal microscopy, NBs were evenly distributed and showed a round shell structure (Figure 1B). The particle size of the obtained NBs was 485.7±33 nm, and the polydispersity index was 0.026 according to a Malvern detector (Malvern Instruments; Figure 1C). The distribution range of the particle size was narrow, which was conducive to single-frequency US excitation.

#### In vitro cell viability under UTND

We first demonstrated in escalating-dose studies of DOX (0–50 µM) that the cytotoxicity of DOX reached a plateau after 10 µM of DOX had been added to PC3 cells. Through CCK-8 assay, cell viability was 43.16%±1.7% in the DOX + NB + US group, 54.15%±1.52% in the DOX + US group, 72.47%±1.7% in the DOX + NB group, and 71.9%±2.32% in the DOX group (Figure 2). Except for the fact that there was no statistical difference between the DOX + NB group and the DOX group, the differences in cell viabilities between any two groups were significant (*P<0.05*). The results indicated that US had a synergetic inhibition effect in DOX against PC3 cells, and NBs could strengthen this inhibition effect under US irradiation.

#### Distribution conditions of DOX in different groups

US monitoring with CEUS mode showed that NBs could be completely destroyed under three different levels of US power (1, 1.75, and 2.5 W/cm²) (Figure 3). However, the next pulse had already been emitted after the burst of NBs by the previous pulse in the built-in 100 Hz pulse, and thus NBs could not perfuse the xenograft region fully. Therefore, an effective
Inhibition of PCa with DOX under US-targeted NB destruction

A burst of NBs could not be achieved in the xenograft region, and the biological effects of the NBs could not be effectively exerted in the xenograft region. Therefore, for the selection of the US irradiation mode, the built-in 100 Hz in the US therapeutic equipment combined with the manual functional 5 seconds on–5 seconds off mode was used. For the selection of US power, it was shown that the US therapeutic probe became significantly hot after a certain period of high-US power input. Therefore, we selected the lowest US power in this equipment, 1 W/cm², for irradiation to exclude the interference of thermal effects produced by US. Real-time observation using the US diagnostic apparatus showed that NBs in the circulation of the body of mice could be completely destroyed within 15 minutes under those conditions of therapeutic US in the xenograft region.

DOX absorbance was measured using full-spectrum scanning. The results showed that DOX absorbance had two peaks: 227 and 485 nm. Different concentrations of DOX were used to generate curves. The correlation analysis between absorbance and DOX concentrations in the supernatant of all the relevant tissues and organs had the best linear correlation at the 485 nm wavelength (Table 1). Through the measurement of DOX absorbance in the supernatants of

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**Figure 1** Morphology and size distribution of NBs.

Notes: (A) NBs observed under TEM after negative staining; (B) NBs observed under LSCM; (C) size-distribution curve of NBs from dynamic light-scattering test in a Malvern detector.

Abbreviations: LSCM, laser-scanning confocal microscopy; NBs, nanobubbles; SD, standard deviation; TEM, transmission electron microscopy; PdI, polydispersity index.

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**Figure 2** Growth-inhibition assay (CCK-8) for PC3 cells in different treatment groups.

Note: *P*<0.05.

Abbreviations: DOX, doxorubicin; NB, nanobubble; US, ultrasound.
livers, tumor xenografts, kidneys, hearts, lungs, and spleens of nude mice in all the groups, DOX concentrations in these tissues were obtained (Figure 4).

After statistical analysis, there were no significant differences in any two DOX concentrations from kidneys, lungs, and spleens in the four groups. However, DOX concentrations from any two groups in livers, tumors and hearts showed a significant difference ($P<0.05$), except between the DOX group and the DOX + NB group. In detail, DOX concentrations in tumor-xenograft tissues were 43.71±5.03 ng/g in the DOX + NB + US group, 36.9±5.72 ng/g in the

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**Table 1** Standard curves between the absorbance and the DOX concentration in different tissue supernatants

<table>
<thead>
<tr>
<th>Solution</th>
<th>$R^2$</th>
<th>Formula</th>
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<tbody>
<tr>
<td>Methanol</td>
<td>0.992</td>
<td>$y = 0.002x + 0.046$</td>
</tr>
<tr>
<td>Liver homogenate</td>
<td>0.981</td>
<td>$y = 0.002x + 0.056$</td>
</tr>
<tr>
<td>Spleen homogenate</td>
<td>0.986</td>
<td>$y = 0.002x + 0.053$</td>
</tr>
<tr>
<td>Kidney homogenate</td>
<td>0.991</td>
<td>$y = 0.002x + 0.061$</td>
</tr>
<tr>
<td>Pulmonary homogenate</td>
<td>0.982</td>
<td>$y = 0.003x + 0.05$</td>
</tr>
<tr>
<td>Myocardial homogenate</td>
<td>0.981</td>
<td>$y = 0.002x + 0.06$</td>
</tr>
<tr>
<td>Tumor homogenate</td>
<td>0.998</td>
<td>$y = 0.003x + 0.053$</td>
</tr>
</tbody>
</table>

Note: Absorbance at 485 nm indicated by $y$; DOX concentration (µg/mL) indicated by $x$.

Abbreviation: DOX, doxorubicin.
DOX + US group, 29.48±5.76 ng/g in the DOX + NB group, and 29.58±4.82 ng/g in the DOX group. However, DOX concentrations in myocardial tissues and hepatic tissues in all groups generally showed the opposite distribution, in contrast with tumor tissues: the concentrations from myocardial tissues and hepatic tissues in the DOX group and DOX + NB groups were the highest, followed by the DOX + US group (18.21±3.63 and 117.22±18.14 ng/g, respectively), and DOX concentrations in the DOX + NB + US group were the lowest (13.49±3.14 and 97.92±11.25 ng/g, respectively). All of these results suggested that US irradiation in the tumor region could shift the DOX distribution from some specific organs to the irradiation tumor, and NBs under US irradiation could also enhance the effect. It was beyond the scope of our study as to why no differences of DOX concentrations in the four treatment groups occurred in the kidney, lung, or spleen.

Effects of DOX on the inhibition of xenograft growth under UTND technology

According to the experimental schedule (Figure 5A), five groups of mice were treated, and some indicators were observed. The results showed that the xenograft volumes, the body weights of mice, and reductions in peak intensity in CEUS mode in the DOX + NB + US group and DOX + US group were significantly different from those in the control group (P<0.05; Figure 5B–G). Tumor volumes in the DOX + NB + US group (68.16±14 mm³) were smaller than those in the DOX + US group (150.87±48.35 mm³, P<0.05), while the body weight of mice and peak intensity of US imaging between these two groups showed no significant difference (23.44±0.48 g vs 23.48±0.76 g, 14.2±1.4 dB vs 15.±0.84 dB; P>0.05). US imaging under CEUS mode in the DOX + NB + US group also showed an unfilled echo-free region in the center of the tumor xenografts, indicating necrosis inside the tumor xenografts, and the DOX + US group had the second-highest level of necrosis (Figure 5D). Xenograft density was calculated according to the mass and volume of the xenograft. The results showed that density in the DOX + NB + US group (1.06±0.03 g/cm³) and the DOX + US group (1.16±0.03 g/cm³) was significantly decreased compared to that of the control group (1.49±0.13 g/cm³), while xenograft masses from any two groups were significantly different, except between the DOX and DOX + NB groups (Figure 5H). Through these analyses, the volume and mass of xenografts were sensitive indicators in reflecting the synergetic effect of NBs on DOX against the implanted tumor under US irradiation, but the density of tumor and the body weight of mice were not.

Myocardial tissue sections from animals in the five groups showed that there were no obvious differences in the H&E sections. The results of TUNEL staining, reflecting apoptosis conditions, indicated that some cardiomyocytes underwent apoptosis in the DOX and DOX + NB groups (Figure 6A–J). The H&E staining results of xenograft tissue sections revealed obvious cellular necrosis in the DOX + US and DOX + NB + US groups, which was also confirmed by TUNEL staining (Figure 6K–T). The overall results of the myocardial tissue and xenograft sections showed a negative correlation.

Observations about effects of UTND on cells and tumor xenografts with electron microscopy

To investigate whether NBs could produce cavitation effects under US irradiation, we designed four groups: control, NB, US, and NB + US. Patterns of changes in prostate cancer cells and tumor xenografts were observed using SEM and TEM, respectively. SEM results showed that the cell surface had significant changes in the NB + US group; there were more folds and small holes. Although cells in the US group had small holes, the number of holes was significantly reduced. The surfaces of the PC3 cells were smooth, and the holes on the surface were not obvious in the NB or control groups (Figure 7A–D). TEM results showed that the microstructure of all the groups had limited changes after the prostate cancer xenografts were stained with uranyl acetate and lead nitrate for electron staining. However, dense particles formed by lanthanum could be clearly observed inside cells in the US and NB + US groups (Figure 7E–H). To confirm further the presence of lanthanum agents in cells and tissues, the
Figure 5 Changes in tumor growth in the different groups.

Notes: (A) Therapeutic and monitoring schedule of tumor xenografts. (B) Conditions of tumor-bearing mice after respective treatments. Red circles indicate regions of tumor xenografts. (C) Status of isolated tumor xenografts in all five groups after treatments. (D) CEUS imaging of maximum sections of tumor xenografts in the five groups before and after treatment. Blue circles indicate tumor-xenograft regions. (E) Changes in the tumor-xenograft volume with days of intervention measures in the five groups. (F) Changes in the body weight of tumor-bearing mice with days of intervention measures in the five groups. (G) Changes in the ultrasonic signal intensity of the maximum sections of xenografts in the five groups. (H) Mass and density of final isolated tumor xenografts in all five groups. All scale bars 2 cm.

Abbreviations: CEUS, contrast-enhanced US; DOX, doxorubicin; NB, nanobubble; US, ultrasound.
Figure 6 H&E and TUNEL staining of myocardial tissues and xenograft tissues of nude mice in all groups.

Notes: (A–E) H&E staining of myocardial tissues of all groups; (F–J) TUNEL staining of myocardial tissues of all groups; (K–O) H&E staining of xenograft tissues of all groups; and (P–T) TUNEL staining of xenograft tissues of all groups. Scale bars 50 µm.

Abbreviations: DOX, doxorubicin; H&E, hematoxylin–eosin; NB, nanobubble; TUNEL, terminal deoxynucleotidyl transferase deoxyuridine triphosphate nick end labeling; US, ultrasound.

Figure 7 Cavitation effect of US burst of NBs on cells and tumor xenografts.

Notes: (A–D) Effects of the four types of treatment on the cell surface visualized under SEM. White arrows indicate pores on the cell surface. (E–H) Condition of lanthanum nitrate particles in tumor xenografts of all treatment groups visualized under TEM after electron staining. Black arrows indicate the locations of lanthanum nitrate-particle deposition. (I–L) Observation of the status of lanthanum nitrate particles in tumor xenografts under TEM without electron staining. Black arrows indicate the locations of lanthanum nitrate-particle deposition.

Abbreviations: NBs, nanobubbles; SEM, scanning electron microscopy; TEM, transmission electron microscopy; US, ultrasonic irradiation.
distribution of lanthanum agents in tumor xenografts without electron staining was observed. The results showed that the NB + US group had the most extensive distribution of lanthanum agents, which were mostly observed in the mitochondria and endoplasmic reticulum. There were few lanthanum particles in the control and NB groups (Figure 7I–L).

**Discussion**

Prostate cancer is a malignant tumor that severely threatens the health of elderly men. Currently, there are many types of treatment methods, of which chemotherapy is an important option for patients with surgical contraindications and metastasis.¹³,¹⁴ Because DOX belongs to cell-cycle-nonspecific chemotherapeutic drugs, it has killing effects in a variety of tumors. The mechanism of action of DOX mainly involves entering cells to interact with cellular genetic materials, such as DNA and RNA, to produce killing effects on tumor cells. However, DOX has significant side effects, including myelosuppression and cardiotoxicity; therefore, its clinical application has certain restrictions, especially for elderly patients or patients with poor cardiac function. In addition, because DOX has the fluorophore feature, it is easily detected and quantitated; therefore, it has been extensively applied in many studies of tumor treatment using drug carriers, such as gold particles, magnetic particles, and microbubbles.¹⁵,¹⁶

Some researchers have reported that pure low-frequency US could promote the uptake and distribution of liposomal DOX in prostate cancer xenografts, which might be associated with the production of cavitation nuclei by fluid in xenografts, such as blood under US excitation.¹⁷ Since Gramiak et al first discovered that microbubbles could enhance US images in 1968, UCAs have received extensive attention.¹⁸ Numerous studies have confirmed that UCAs can be used not only as diagnostic agents but also as a type of effective carrier to achieve the targeted delivery of drugs under US irradiation. In UTMD technology, UCAs are used as cavitation nuclei, and begin to produce periodic contractions and expansions around the equilibrium radius to generate radiation pressure and microjets at surroundings under specific acoustic pressures, which is termed stable cavitation. With further increases in US intensity, UCAs are ruptured, and the generated high heat and high pressure can result in the production of strong shock waves, high-speed jet flows, and free radicals; this effect is termed inertial cavitation. Cavitation is an important basis of UTMD technology, and can increase tissue and vascular endothelial gaps and membrane permeability to increase the entry of drugs into parenchymal cells at the US-irradiation location to exert their functions. Cavitation is typically applied in studies of the opening of the blood–brain barrier.¹⁹–²¹ A large number of studies have reported that UTMD technology promoted the release of many substances inside UCAs or carried by the outer shell, such as chemotherapeutic drugs and genes, to kill tumor cells at a short distance, thus effectively treating tumors.²²–²⁴

Other researchers have prepared UCAs using degradable polymer outer shells to encapsulate liquid droplets, which were injected into animals and entered tissue gaps through tumor endothelial gaps. These NBs coalesced into highly echogenic microbubbles to release encapsulated drugs under US irradiation.²⁵ In the aforementioned studies on the production of cavitation by US irradiation combined with UCA, they were mainly at the micron level. Our previous studies confirmed that NBs could enter tumor-tissue gaps through the EPR effect of tumor tissues to exert important functions in the targeted diagnosis of tumors. However, further investigation is still required to determine whether cavitation can be produced or promoted under US irradiation to promote the function of drugs/genes on tumors at a short distance. Based on these considerations, we examined whether the features of lipid NBs, including their strong penetration ability and more extensive distribution in tumor tissues compared with microbubbles, could be combined with the traditional chemotherapeutic drug DOX to investigate their effects on prostate cancer and underlying mechanism under US irradiation. The results might provide a basis for further in-depth studies of the use of US NBs as a drug/gene-delivery system.

The preparation of a drug or gene carrying UCAs involves two main methods. One method is directly to adhere drugs or genes to the outer membrane of UCAs (adhesion method), and the other method is to consolidate drugs or genes inside the membrane (integration method).²⁶ Although current UCAs have many different types of outer shells (including proteins, lipid materials, and polymers), UCAs with lipid shells have excellent acoustic features and safety; therefore, this type of UCA has the most extensive applications in clinical settings and laboratories. Studies of US therapy have shown that NBs composed mainly of polymers also have excellent functions in the encapsulation of drugs, for example, it has been reported that through the construction of polymer NBs, the encapsulation of a sufficient amount of DOX was achieved, which had significant inhibitory effects on tumors.²⁷ However, physical conditions during their preparation are complex and difficult to control. In addition, the destruction of polymer UCAs requires high-intensity US-power output, which has damaging effects on tissues on the acoustic channel.

Although UCAs with lipid shells have excellent safety, the lipid shell is thin, and the particle size of UCAs is small,
such that the amount of drug carried by the UCAs is limited. To achieve therapeutic effects, the amount of a lipid UCA is usually ten- to 100-fold that in diagnosis, which not only greatly increases the cost but also has safety issues when applying large doses. Our prepared NBs also had the same issue. Therefore, to investigate the effects of NBs on the promotion of DOX to inhibit prostate cancer growth under US irradiation and the underlying mechanism, this study used the method of injecting a mixture of DOX and NBs to ensure the proper DOX concentration. The effects on the inhibition of prostate cancer cells and xenografts under UTND were studied, and the related mechanism was further analyzed. The results showed that the inhibition rate of cell growth and DOX concentration in xenografts were highest in the DOX + NB + US group, and gradually reduced in the order of the DOX + US, DOX + NB, and DOX groups; however, the DOX concentrations in myocardial tissues showed the opposite trend. In addition, the reduction in xenograft volume after nine-time interventions was the most significant in the DOX + NB + US group. TUNEL staining results of myocardial tissues and tumor xenografts further showed that the xenograft tissues in the DOX + NB + US and DOX + US groups had cellular necrosis, which was most significant in the former group. Myocardial tissues had only a few apoptotic cells, indicating that NBs could increase the killing effects of DOX in irradiated regions (xenograft location) under UTND technology.

Under the condition that the total amount of injected DOX was the same, UTND could change the distribution of DOX and reduce the distribution of DOX in other unirradiated tissues and organs, thus reducing the toxic effects of DOX on the heart. SEM results showed that there were more holes in prostate cancer cells in the DOX + NB + US group, and cell-surface folds were significantly increased. In the observation of tumor xenografts using TEM, lanthanum nitrate with a molecular weight of 433 and a mean particle size of ~40 Å was used as an electron tracer of ultrastructure (when the permeability of the cell membrane was normal, lanthanum nitrate was present only in extracellular gaps and could not enter cells). The results showed that UTND could increase the uptake of lanthanum nitrate by tumor xenografts, indicating that the UTND technology was conducive to increasing the permeability of tumor-cell membranes. This method was effective for promoting drugs to act on specific sites, and could effectively increase the concentration of drugs in tumor regions. The increase in holes on the cell surface and the increase in permeability both confirmed that UTND technology could produce cavitation effects and provides evidence for relevant studies on the promotion of tumor treatment by genes or drugs.

The injection method for the mixture of drug and NBs used in this study had the following features, which are worthy of further in-depth studies and clinical translation: 1) Because lipid microbubbles have been extensively applied in the clinic, NBs using lipids as the outer shell have excellent safety features in human applications; 2) although DOX and NBs were injected into the vein as a mixture, our results confirmed that NBs altered the drug distribution under US irradiation to reduce the toxic effects of DOX on the heart, which effectively reduced damaging effects on other normal tissues at the same time as increased treatment effects compared to using DOX alone; 3) the analysis of the related mechanism showed that US irradiation on NBs produced cavitation effects similar to microbubbles to promote the permeability of tumor cells, which would be helpful for drugs or genes to produce tumor-killing effects in specific regions of US irradiation; 4) we selected 1 W/cm² US power in this experiment to perform 15 minutes of 100 Hz combined with the 5 seconds on–5 seconds off irradiation mode on tumor regions. On the basis of ensuring full perfusion and burst of all the NBs in the body, the thermal effects of the therapeutic US were reduced to the greatest extent possible, which was conducive for treatment targeting internal solid tumors and did not damage normal tissues on the acoustic channel at the same time. Therefore, the distribution of cytotoxic drugs to tumors through the circulation is increased, and nontarget-organ damage is reduced.

However, our study also had the following limitations. Although the low-intensity US therapeutic equipment used in this study can be easily and rapidly adjusted, the available parameters were limited and the irradiation time was controlled manually, together with built-in parameters in the apparatus. Also, we did not include measurements of such parameters as cavitation-threshold values or resonance frequency of our fabricated lipid NBs, which should be resolved in further study.

**Conclusion**

This study confirmed that UTND can produce cavitation effects to promote the effects of DOX on the inhibition of prostate cancer growth. In addition, under a specific drug concentration in the body, this technique can change the distribution of drugs in the body to reduce toxic side effects on normal tissues. This study not only provides experimental methods and evidence for the delivery of chemotherapeutic drugs by NBs under a low-intensity US burst but also lays the foundation for future relevant studies on tumor treatment using NBs as drug/gene-delivery carriers.
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Disclosure
The authors report no conflicts of interest in this work.

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