Targeted Ultrasound Nanobubbles Therapy for Prostate Cancer via Immuno-Sonodynamic Effect

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Background: Prostate cancer (PCa) poses a significant global health threat. Immunotherapy has emerged as a novel strategy to augment the inhibition of tumor proliferation. However, the sole use of anti-PD-L1 Ab for PCa has not yielded improvements, mirroring outcomes observed in other tumor types.

Methods: This study employed the thin film hydration method to develop lipid nanobubbles (NBs) encapsulating chlorin e6 (Ce6) and anti-PD-L1 Ab (Ce6@aPD-L1 NBs). Our experimental approach included cellular assays and mouse immunization, providing a comprehensive evaluation of Ce6@aPD-L1 NBs’ impact.

Results: The Ce6@aPD-L1 NBs effectively induced reactive oxygen species generation, leading to tumor cells death. In mice, they demonstrated a remarkable enhancement of immune responses compared to control groups. These immune responses encompassed immunogenic cell death induced by sonodynamic therapy and PD-1/PD-L1 blockade, activating dendritic cells maturation and effectively stimulating CD8+T cells.

Conclusion: Ce6@aPD-L1 NBs facilitate tumor-targeted delivery, activating anti-tumor effects through direct sonodynamic therapy action and immune system reactivation in the tumor microenvironment. Ce6@aPD-L1 NBs exhibit substantial potential for achieving synergistic anti-cancer effects in PCa.

Keywords: Ce6, nanobubbles, ultrasound, sonodynamic therapy, immunotherapy

Introduction

Prostate cancer (PCa) stands as a hormonally driven malignancy and ranks as the second most frequently diagnosed cancer in men globally.1 Androgen deprivation therapy emerges as a crucial treatment for patients with locally advanced and metastatic PCa, yet almost all patients eventually develop androgen-independent PCa.2 Another therapeutic avenue is chemotherapy, with docetaxel (DTX) being the established first-line cytotoxic drug proven to extend overall survival.3,4 However, the utilization of DTX-based chemotherapy may face limitations due to drug resistance and severe side effects, encompassing anaphylaxis, myelosuppression, gastrointestinal toxicity, and neurotoxicity.5

Since patients no longer amenable to surgical cure and exhibiting poor responsiveness to medical therapy, impeding tumor growth becomes pivotal in mitigating disease progression. Tumor immunosuppression, intricately linked with the tumor microenvironment (TME), expedites tumor growth.6 Consequently, immunotherapy emerges as a novel treatment strategy to augment the inhibition of tumor proliferation. Immune checkpoints, including programmed cell death 1 (PDCD1/PD-1) and programmed cell death 1 ligand 1 (PDCD1LG1/PD-L1), play pivotal roles in mediating tumor immunosuppression.7 Leveraging immune checkpoint blockade (ICB), anti-PD-L1 Ab are progressively employed to prevent tumor cells from evading the immune system, presenting a promising and viable strategy for tumor eradication.8 However, PD-L1 expression is not exclusive to tumor cells but also found in normal tissues like vascular endothelial cells, hepatocytes, and mesenchymal stem cells.9 Thus, conventional intravenous injections of free anti-PD-L1 Ab may lead to non-target interactions with normal tissues. Such non-targeted combinations not only result in low concentrations...
in the tumor but also give rise to immune-related adverse effects, including endocrine diseases, skin toxicity, and gastrointestinal symptoms.\textsuperscript{10,11}

For achieving high concentrations in the tumor while ensuring the safe use of anti-PD-L1 Ab, the development of a tumor-targeting anti-PD-L1 Ab delivery system is imperative. Microbubbles (MBs) have the capability to transport drugs or genes to tumor, and ultrasound (US)-targeted MB destruction (UTMD) enhances the therapeutic efficacy.\textsuperscript{12} However, the size of MBs, typically ranging from 1–4 µm, limits their penetration into tumor tissues. On the other hand, nanobubbles (NBs) with a size of 100–500 nm can passively accumulate in tumor tissues through the enhanced permeability and retention (EPR) effect. NBs loaded with anti-PD-L1 Ab can bind to tumor cells, facilitating targeted aggregation of NBs within the tumor. The cavitation effect generated by NBs under US irradiation further promotes the release of anti-PD-L1 Ab.\textsuperscript{13}

While anti-PD-L1 Ab have been employed in clinical experiments for treating patients with PCa, the observed improvements have been lower in proportion and level compared to other tumor types.\textsuperscript{14} Consequently, the objective of our research was to identify methods to enhance treatment effectiveness. Leveraging the characteristics of NBs and UTMD, we aimed to achieve sonodynamic therapy (SDT) through US irradiation. SDT has the potential to induce immunogenic cell death (ICD) of tumor cells, reinitiating an anti-tumor immune response. Additionally, SDT inflicts damage on tumor cells under US irradiation with sonosensitisers.\textsuperscript{15} US irradiation enhances cell membrane permeability, thereby contributing to the efficient cellular uptake of drugs.\textsuperscript{16}

Chloroene (Ce6) a stable degradation product of chlorophyll-a (the main component of green plants), possesses a four-pyrrole ring structure that readily absorbs light. Recognized as a top-tier photosensitiser and sonosensitiser in nature, Ce6 offers the advantages of being non-toxic, exhibiting good specificity, and rapid metabolism.\textsuperscript{17} Furthermore, Ce6 is a representative and promising sonosensitizer,\textsuperscript{18} which has been used to treat solid tumors in the animal study.\textsuperscript{19} The free form of Ce6, when intravenously injected into the body, fails to selectively concentrate in lesions. Therefore, nanotechnology modifications are essential to improve the tissue selectivity of free Ce6.\textsuperscript{20} Previous researches have confirmed that Ce6 could be assembled well onto the surface of NBs, and effectively promoted the generation of reactive oxygen species (ROS) under US irradiation.\textsuperscript{21}

The initial step in constructing targeted NBs is the effective conjugation of Ce6 and anti-PD-L1 Ab. Avidin-biotin serves as the preferred conjugation mode in research due to several advantages. Notably, the avidin-biotin affinity surpasses that of antigen-antibody interactions.\textsuperscript{22} Furthermore, avidin’s four binding sites facilitate increased loading of anti-PD-L1 Ab.\textsuperscript{23} Additionally, considering that free Ce6 is insoluble in water but soluble in lipids, and since NBs share the same composition, this conjugation approach aligns well with the characteristics of the components. Following this rationale, we constructed tumor-targeted NBs encapsulating anti-PD-L1 Ab and Ce6 (Ce6@aPD-L1 NBs) to validate their anti-tumor effects through in vitro and in vivo experiments.

Upon reaching the tumor via passive and active targeting modalities, Ce6@aPD-L1 NBs effectively inhibit tumor growth. This inhibition is attributed not only to the direct action of SDT but also to the reactivation of the immune system, enhancing immune responses within the TME. The synergistic immunotherapy involves ICD induced by SDT, leading to the translocation of calreticulin (CRT) to the surfaces of tumor cells. This process activates dendritic cells (DCs) and stimulates CD8+T cells, thereby enhancing the release of various tumor immune factors. In summary, these promising Ce6@aPD-L1 NBs exhibit significant potential for achieving synergistic anti-cancer effects against PCa (Figure 1).

Materials and Methods

Materials

Biotin-polyethylene glycol distearoylphosphatidylethanolamine2000 (Biotin-DSPE-PEG2000) and disaturated phosphatidyl choline (DPPC) were acquired from Melopheg. Ce6 was obtained from Macklin, 3β[N-(N’, N’-dimethylaminoethane)-carbamoyl] cholesterol (DC-chol) from Glpbio, streptavidin (SA) from Solarbio, and biotin-anti-PD-L1 Ab from Thermo Fisher Scientific.

Trichloromethane, PBS, C₃F₈, and glycerol were sourced from Servicebio (China).
Preparation of Ce6@aPD-L1 NBs
Ce6@aPD-L1 NBs were synthesized through a thin film hydration method. Initially, a mixture of 31 mg contained DPPC: biotin-DSPE-PEG2000: DC-chol: Ce6 (5:2:0.5:0.25) was dissolved in a glass-bulb tube with trichloromethane. The liquid was evaporated using a rotary evaporator, creating a thin film on the tube wall. Next, 10 mL of glycerol: PBS (1:9) was added, and the tube was shaken in a constant-temperature bath at 45°C for 1 h. Subsequently, 500 µL of the liquid was injected into a penicillin bottle with C₃F₈, subjected to rapid shake by amalgamator with 1200 rpm for 60s, and then stewed for 5 min to obtain biotin-Ce6-NB suspension. SA: suspension (3:50) was co-incubated at 37°C for 1 h, and biotin-anti-PD-L1 Ab, twice as much as SA, was added to incubate in the refrigerator at 4°C overnight, resulting in the successful preparation of Ce6@aPD-L1 NBs.

Test for Physical and Chemical Properties of NBs
The size, zeta potential, and particle dispersion index (PDI) of NBs were measured using a Zetasizer. Nano ZS at 25°C (Malvern, UK). The morphology and distribution of different NBs were observed using optical microscopy, and the red fluorescence of Ce6 was detected using fluorescence microscopy (FSMS). The inspections were conducted at 0, 24, 48, and 96 h, respectively.

Ce6 and Anti-PD-L1 Ab Loading Test of Ce6@aPD-L1 NBs
Following the synthesis of Ce6@aPD-L1 NBs, a fluorescein isothiocyanate (FITC)-conjugated biotin-anti-PD-L1 antibody was introduced to the NBs and incubated for 1 h at room temperature.
The connectivity of the anti-PD-L1 Ab was assessed using flow cytometry with excitation and emission wavelengths of 488 and 525 nm, respectively. Similarly, Ce6 was analysed using flow cytometry at excitation and emission wavelengths of 638 and 660 nm, respectively.

**Cell Culture**

The human PCa cell line PC3, obtained from ProCell (China), was cultured in F12K medium containing 10% fetal bovine serum (FBS, Gibco) and 1% penicillin/streptomycin under a humidified atmosphere of 5% CO₂ at 37°C.

**Cellular Uptake of Ce6**

PC3 cells in logarithmic growth phase were inoculated into 12 hole plates, and the cell density of each hole was about 30%. When the cells were in good condition, they were divided into three groups: Ce6@aPD-L1 NBs, Ce6 NBs and free Ce6, each group with three holes. The same content of Ce6 was added to the hole plate and the PC3 cells were treated with ultrasonic irradiation (2 MHz, 1.5 W/cm², 20% duty cycle) for 5 min. Then they were placed in 37 °C 5% CO₂ constant temperature incubator for 12 h. The cells were washed with PBS for three times and stained with diamidino phenylindole (DAPI). Last, the aggregation of Ce6 in cells was observed under FSMS and cellular uptake of Ce6 detected by flow cytometry.

**Detection of ROS Generation**

PC3 cells in the logarithmic growth phase were plated into 12-well plates, with each well reaching approximately 30% cell density. Four groups were established: NBs, free Ce6, Ce6 NBs, and Ce6@aPD-L1 NBs, each containing three wells. Subsequently, the cells were placed in a 37 °C 5% CO₂ constant temperature incubator. After 12 h of culturing, each well received 200μL of drugs and was subjected to sonication (2 MHz, 1.5 W/cm², 20% duty cycle) for 5 min. All groups were then exposed to a mixture of DMEM and DCFH-DA at a volume ratio of 1:1000, followed by an additional 30 min incubation at 37 °C. After washing with PBS, fluorescence intensity was measured using a fluorescence microscope (excitation wavelength, 488 nm; emission wavelength, 525 nm). Finally, flow cytometry was employed to determine the number of cells expressing ROS.

**Animal Model**

BALB/C-Nude male mice were procured from Beijing Vital River Laboratory Animal Technology Co., Ltd., and all procedures involving animals adhered to the guidelines of the Institutional Animal Care and Use Committee of Wuhan University. A subcutaneous implantation of 1×10⁶ PC3 cells in 100 μL of F12K medium without FBS was performed at the dorsal side of the left front leg of 4- to 6-week-old mice to establish the animal tumor model.

**Biodistribution and Pharmacokinetics of Ce6 in vivo**

When the tumor diameter reached 5 ± 2 mm, mice (n = 3) were intravenously injected with free Ce6, Ce6 NBs, and Ce6@aPD-L1 NBs (250 μL each) via the tail vein. At 0.5, 1, 6, 12, 24, and 48 h post-injection, the fluorescence signals of the mice were monitored using an in vivo fluorescence imaging instrument (AniView Phoenix, BLT) with an excitation wavelength of 620 nm and an emission wavelength of 670 nm. Tumors and major organs were excised at the time of the strongest fluorescence signals after injection and examined separately using the fluorescence imaging instrument.

**Synergistic Anti-Cancer Effect in vivo**

Tumor-bearing mice were randomly allocated into five groups and treated with PBS (control group), Free Ce6 + SDT, Ce6 NBs + SDT, aPD-L1 NBs + SDT, and Ce6@aPD-L1 NBs + SDT. The drug doses for each mouse were 250μL per injection at intervals of 2 days for a total of five times.

When the tumor diameter reached 5 ± 2 mm, mice underwent ultrasonic insonation (2MHz, 1.5 W/cm², 20% duty cycle, 10 min) at 6 h post-administration. Survival rates and body weights of the mice were recorded and plotted against...
time. Additionally, tumor volumes were calculated using the formula: \( \text{Volume} = 0.5 \times L \times W^2 \), where \( L \) and \( W \) represent the length and width of the tumor, respectively.

**Biochemistry Index and Histology Analysis**

On day 15 post the initial treatment, the tumor was excised for haematoxylin-eosin staining (HE) to observe alterations in the tumor tissue. The TUNEL immunofluorescence kit was employed to identify apoptotic cells in the tumor tissues. Additionally, real-time fluorescence quantitative polymerase chain reaction (RT-PCR) was conducted to assess the expression of B-cell lymphoma-2 (Bcl-2), Bcl-2 associated x (Bax), interferon-\( \gamma \) (IFN-\( \gamma \)), cluster of differentiation 80 (CD80), CD86, transforming growth factor-\( \beta \) (TGF-\( \beta \)), and PD-L1. Immunofluorescence staining was utilized to observe the translocation of CRT and the infiltration of CD8+ T cells into the tumor tissue.

**Biosafety Analysis**

Following the completion of the treatment, liver functions, including ALT, AST, and TBIL, were assessed. The major organs of the mice were further evaluated by HE staining to observe changes in the tumor tissue structure. Before HE staining, the liver was weighed, and the liver index weight (liver / body weight \( \times 100\% \)) was calculated for all groups.

**Statistical Analysis**

Data are presented as the mean \( \pm \) standard deviation (SD). Statistical analyses were performed conducted using ANOVA with a Tukey’s post hoc test. A significance level of \( P < 0.05 \) was considered to be statistically significant.

**Results**

**Physical and Chemical Properties of Ce6@aPD-L1 NBs**

Under an optical microscope, Ce6@aPD-L1 NBs were observed to be spherical, uniform, with no apparent aggregation or clustering. The loading contents of Ce6 and anti-PD-L1 Ab on the Ce6@aPD-L1 NBs were 86.1% and 73.8%, respectively. Within 24 h, all test indices of Ce6@aPD-L1 remained stable However, after 48 h, changes were noted in the diameter, PDI, Zeta potential, Ce6 and anti-PD-L1 Ab loading rate (\( P<0.05 \)) (Figure 2).

**Cellular Uptake of Ce6 and SDT Assessments in vitro**

After 12 h of culture with drugs, PC3 cells exhibited blue fluorescence in their nuclei and red fluorescence in the cytoplasm when stained with DAPI, confirming the uptake of different Ce6 formulations by the cells. Cellular uptakes of Ce6@aPD-L1 and Ce6 NBs were higher than that of Free Ce6, as observed through FSMS and flow cytometry. Cells treated with NBs showed minimal ROS generation, while ROS levels were detectable in cells treated with different Ce6 formulations, with the Free Ce6 group exhibiting the lowest ROS production (Figures 3 and 4).

**Biodistribution of Ce6 in vivo**

After intravenous administration for 12 h, tumors and major organs were excised and monitored using an in vivo fluorescence imaging instrument. Results showed that Free Ce6, Ce6 NBs, and Ce6@aPD-L1 NBs accumulated at tumor sites, with Ce6@aPD-L1 NBs demonstrating more efficient tumor accumulation than the others at all time points. Fluorescence signals in the liver and kidney were also detected, with the Ce6 signal being lower than that in the tumor (Figure 5).

**Synergistic Anticancer Effect of Immunotherapy and SDT**

Except for the control group, all other groups exhibited varying degrees of therapeutic effects, with tumor growth inhibited in mice treated with Ce6@aPD-L1 NBs + SDT. On day 15 after the first treatment, the tumor volume in the Ce6@aPD-L1 NBs + SDT group was 216 \( \pm \) 89 mm\(^3\). In comparison, tumors grew to 803 \( \pm \) 106, 749 \( \pm \) 137, 337 \( \pm \) 126, and 383 \( \pm \) 108 mm\(^3\) in the control, Free Ce6 + SDT, Ce6 NBs + SDT, and aPD-L1 NBs + SDT groups, respectively (Figure 6A and B).
Consistent with the tumor growth inhibition results, Ce6@aPD-L1 NBs + SDT most effectively prolonged the survival rate of mice (Figure 6C). The rate of body weight gain in the control and Free Ce6 + SDT groups decreased gradually, whereas that in the other groups increased (Figure 6D).

Tumor tissues from animals treated with PBS were hypercellular. In contrast, the cancer cell density in tumor tissues from animals receiving Ce6 NBs + SDT, aPD-L1 NBs + SDT, and Ce6@aPD-L1 NBs + SDT decreased. Moreover, the TUNEL assay showed that the tumor tissue of the Ce6@aPD-L1 NBs + SDT group exhibited the most significant apoptosis (Figure 7A). Compared with the control group, the BAX expression level was up-regulated, and the group of Ce6@aPD-L1 NBs + SDT showed the most significant increase (P < 0.01). Conversely, BCL2 was downregulated, with the group of Ce6@aPD-L1 NBs + SDT also showing the most significant decrease (P < 0.01) (Figure 7B and C).

Immunofluorescence staining demonstrated that compared to the rare tumor infiltration of CD8+ T cells in the group, tumor infiltration of CD8+ T cells increased with treatment using other drugs. Particularly, Ce6@aPD-L1 NBs + SDT promoted tumor infiltration of CD8+ T cells (Figure 8A). During aPD-L1 NBs + SDT and Ce6@aPD-L1 NBs + SDT, the CD80, CD86, and IFN-γ expression levels were significantly up-regulated (P < 0.01), while the TGF-β expression level was significantly down-regulated (P < 0.01). Similarly, the PD-L1 expression level was downregulated in the aPD-L1 NBs + SDT and Ce6@aPD-L1 NBs + SDT groups (P < 0.05) (Figure 8B).

Immunofluorescent staining revealed that the CRT fluorescence signal in the tumor tissue was marginal in the control and aPD-L1 NBs + SDT groups. In comparison, the Ce6 NBs + SDT and Ce6@aPD-L1 NBs + SDT groups exhibited the most significant increase (Figure 9).
Biosafety Analysis of Therapy

According to the results of HE staining, no apparent histological damage was observed in the heart, liver, spleen, lungs, and kidneys in all groups. Additionally, levels of ALT, AST, and TBIL were within normal range ($P > 0.05$). Likewise, there were no significant differences of the liver index among the groups ($P > 0.05$) (Figure 10).

Discussion

Cancer-targeted therapy, focusing on treating specific carcinoma sites at the cellular level, has garnered increasing attention from researchers. Immunotherapy, a promising approach to targeted therapy, can be used alone or in combination with chemotherapy, radiotherapy, or surgery. The anti-PD-L1 Ab, a representative type of ICB, has been widely applied in experimental research and clinical treatment. Due to the complex composition of TME, a sole application of ICB may not effectively activate the immune system. Therefore, we developed SDT, a non-invasive therapeutic modality combining low-intensity US and sonosensitisers.

The simultaneous achievement of immunotherapy and SDT anti-cancer effects is crucial. Ce6, a representative and promising sonosensitiser, is a stable degradation product of chlorophyll a and contains four pyrrole ring structures. Ce6 is non-toxic, exhibits good specificity and fast metabolism, and is not easily soluble in water. Therefore, it is necessary to modify free Ce6 using nanotechnology to improve its tissue selectivity, lipophilicity, and ability to bind to NBs with...
lipid components on the surface. In this study, Ce6@aPD-L1 NBs were successfully synthesized according to a specific route. Firstly, the Ce6 loading rate of NBs was 86.1 ± 5.3% and 84.6 ± 4.1% at 0 and 24 h, respectively, illustrating that Ce6@aPD-L1 NBs could carry sufficient quantities of Ce6 to ensure the effectiveness of SDT. Secondly, Ce6@aPD-L1 NBs exhibited much more efficient accumulation in the PC3 tumors of mice than free Ce6 and Ce6 NBs, and also more than in the liver and kidneys, confirming the SDT effect.

The particle size of the Ce6@aPD-L1 NBs was uniform at the nanoscale, with no obvious aggregation or clustering. The diameter, PDI, Zeta potential, Ce6, and anti-PD-L1 Ab loading rate remained nearly constant within 24 h, whereas the test indices of the NBs changed after 48 h. Within the first 24 h, the particle size of NBs was approximately 500 nm, enabling them to pass through the endothelial gap of tumor blood vessels, allowing the NBs to enter the tumor tissue

Figure 4 Generation of ROS in PC3 cells. After cellular uptake of Ce6 and ultrasonic irradiation, the generation of ROS in cells were detected by FAIMS (A) and flow cytometry (B). Scale bar = 20µm.

Figure 5 Biodistribution of Ce6 in vivo. After injecting drugs into the PCa-bearing mice for 12 hours, the tumors and major organs were detected with different intensity signal.
through EPR, thus, passively targeting tumor cells. The anti-PD-L1 Ab on Ce6@aPD-L1 NBs was successfully detected by FSMS and flow cytometry, confirming its successful connection to the NBs and its ability to actively target tumor cells. Moreover, the positively charged surface of NBs facilitated their entry into negatively charged tumor tissue.

When PC3 cells that had internalised Ce6 were exposed to US, the production of ROS in the cells was assessed through FSMS and flow cytometry. ROS generation primarily depends on the synergistic interactions of US, a non-toxic photosensitiser, and molecular oxygen, leading to cellular toxicity and apoptosis. Furthermore, following incubation with different drugs, cells from the Ce6@aPD-L1 NBs and Ce6 NBs groups exhibited higher Ce6 uptake compared to free Ce6. Additionally, these groups showed an increased generation of ROS with SDT. Based on these findings, a nanotechnology-assisted combination may enhance SDT efficiency, offering a potentially safer therapeutic option for PCa.

SDT can result in extensive tumor cell death, which includes the direct cytotoxicity of ROS and the cavitation effect of SDT, which Bilmin has summarized the sono-mechanical changes caused by the oscillations and fast movement of NBs. During tumor progression, cellular immunity is suppressed, and reactivating immunity may contribute to tumor growth inhibition, so the regulation of the TME is crucial for inhibiting tumor cell growth. SDT can induce ICD in tumor cells, triggering an anti-tumor immune response, and the translocation of CRT to the cell surface is a characteristic feature of ICD. In our study, immunofluorescent staining revealed that the CRT fluorescence signal was not observed in the control and aPD-L1 NBs + SDT groups, while abundant CRT was detected in the Ce6 NBs + SDT and Ce6@aPD-L1 NBs + SDT groups, indicating a successful triggering of the ICD reaction.

Recent studies have suggested that SDT may stimulate the release of certain tumor immune factors, including CD80, CD86, IFN-γ, TGF-β, IL2, and IL10. PCR results demonstrated that the expression levels of CD80 and CD86—markers related to the maturation of DCs—increased significantly in the tumor tissue of the Ce6 NBs, aPD-L1 NBs, and Ce6@aPD-L1 NBs group. SDT can induce the upregulation of DCs and enhance anti-tumor immune effects, associated with T lymphocyte infiltration in tumor tissue, exhibited increased expression. In contrast, TGF-β, an immune suppressive factor, displayed downregulated expression, promoting anti-tumor immune effects. The PD-1/PD-L1 signalling pathway inhibits T cell activation and may lead to apoptosis, influencing tumor immune escape. Anti-PD-L1 Ab can specifically bind to the PD-L1 of tumor cells, reactivating and proliferating T cells to enhance anti-tumor immunity.
The results indicated a significant decrease in the expression of PD-L1 in the aPD-L1 NBs and Ce6@aPD-L1 NBs, accompanied by evident infiltration of CD8+ T cells into the tumor tissue, suggesting the reactivation of anti-tumor immunity.

After treating mice with PCa, various groups exhibited varying therapeutic effects, with tumor growth inhibition observed in mice treated with Ce6@aPD-L1 NBs + SDT, except for the control group. On day 15 after the initial treatment, the tumor volume in the Ce6@aPD-L1 NBs + SDT group was 216 ± 89 mm³, while tumors in the control group grew to 803 ± 106 mm³. Consistent with the tumor growth inhibition results, Ce6@aPD-L1 NBs + SDT significantly extended the survival of mice. The rate of body weight growth in the control and Free Ce6 + SDT groups decreased gradually, whereas the other groups exhibited an upward trend, possibly due to the inhibition of tumor growth with less impact on the mice’s overall growth. And we observed that the tumor tissues from animals treated with PBS were hypercellular. In contrast, the cancer cell density in tumor tissues from Ce6 NBs + SDT, aPD-L1 NBs + SDT, and Ce6@aPD-L1 NBs + SDT mice clearly declined. Moreover, the TUNEL assay revealed abundant apoptosis in the tumor tissue of the Ce6@aPD-L1 NBs + SDT group. Compared with the control group, the BAX expression level was upregulated, with the most significant upregulation observed in Ce6@aPD-L1 NBs + SDT. Conversely, BCL2 down-regulation in the Ce6@aPD-L1 NBs + SDT group was the most significant.

NBs, as a type of liposomes, have been developed as novel ultrasound contrast agents to improve diagnostic accuracy and reliability, and they have smaller particle size, more stable performance, and longer circulation time than MBs, which leads to more accumulation in the tumor area. Through binding drugs to the shells or specific ligands, NBs can serve as a multifunctional platform to deliver drugs, increase local drug concentration, control drug release, and achieve treatment of cancer. Recently, nanotechnology development has implemented carrying multiple medications, and NBs...
loaded with Ce6 make SDT potentially a safe and feasible therapeutic means for various tumors. On the other hand, SDT was reported to induce acute inflammation which promotes lymphocytes infiltration in tumor, thus triggering an anti-tumor immunity.
Conclusion
In summary, we successfully developed effective NBs co-loaded with anti-PD-L1 Ab and Ce6 for treating PCa tumors. These NBs can effectively inhibit tumor growth not only due to the direct action of SDT but also because of the reactivation of the immune system, enhancing immune responses in the TME. Synergetic immunotherapy may involve ICD induced by SDT and ICB of PD-1/PD-L1, including the translocation of CRT to the tumor cell surface, release of tumor immune factors, activated maturation of DCs, and effective stimulation of CD8+ T cells. The promising Ce6@aPD-L1 NBs holds great potentials for achieving synergetic anti-cancer effect for PCa, but also can be applied in the treatment of other solid tumors.

Ethics Approval
This experiment was performed according to the Animal Ethics Committee of Renmin Hospital of Wuhan University (IACUC Issue No.20201103). In this study, we adhered to the ARRIVE guidelines and strictly followed the Regulations of the People’s Republic of China on the Administration of Laboratory Animals and its guiding principles for animal experiments. Our approach fully respected the lives of laboratory animals and aimed to minimize their suffering.
Acknowledgments
The authors greatly appreciated the support from the Central Laboratory and Animal Experiment Center of Renmin hospital of Wuhan University. This work was sponsored by the National Natural Science Foundation of China [Grant numbers 82271999].

Author Contributions
All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

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

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The authors report no conflicts of interest in this work.