Hypoxia-degradable and long-circulating zwitterionic phosphorylcholine-based nanogel for enhanced tumor drug delivery

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**Abstract**
Tumor microenvironment has been widely utilized for advanced drug delivery in recent years, among which hypoxia-responsive drug delivery systems have become the research hotspot. Although hypoxia-responsive micelles or polymersomes have been successfully developed, a type of hypoxia-degradable nanogel has rarely been reported and the advantages of hypoxia-degradable nanogel over other kinds of degradable nanogels in tumor drug delivery remain unclear. Herein, we reported the synthesis of a novel hypoxia-responsive crosslinker and the fabrication of a hypoxia-degradable zwitterionic poly(phosphorylcholine)-based (\textsuperscript{1}H)PMPC nanogel for tumor drug delivery. The obtained \textsuperscript{1}HPMPC nanogel showed ultra-long blood circulation and desirable immune compatibility, which leads to high and long-lasting accumulation in tumor tissue. Furthermore, \textsuperscript{1}HPMPC nanogel could rapidly degrade into oligomers of low molecule weight owing to the degradation of azo bond in hypoxic environment, which leads to the effective release of the loaded drug. Impressively, \textsuperscript{1}HPMPC nanogel showed superior tumor inhibition effect both in vitro and in vivo compared to the reduction-responsive phosphorylcholine-based nanogel, owing to the more complete drug release. Overall, the drug-loaded \textsuperscript{1}HPMPC nanogel exhibits a
1. Introduction

Nanogel is a kind of hydrophilic three-dimensional net structure of nanoscale dimension which has been widely used as nanocarrier for tumor drug delivery. In the fabrication of nanogel, the insertion of degradable crosslinker endows the nanogel with controlled drug release in tumor microenvironment including acidic, redox and specific enzymatic conditions. For example, nanogel fabricated by crosslinker containing disulfide bonds could achieve the accelerated drug release triggered by the high concentration of glutathione in tumor cytoplasm. Furthermore, nanogel composed of ketal or acetal bonds-based crosslinker could release the loaded drug rapidly in tumor acidic environment. Despite the great progress of degradable nanogel as drug carriers, there is no report of a type of hypoxia-degradable nanogel and the advantages of hypoxia-degradable nanogel over other kinds of degradable nanogels in tumor drug delivery remain unclear.

Hypoxia featuring low oxygen concentration is a symbol of nearly all the solid tumors, which usually changes the balance of redox states in tumor tissue and leads to the enhanced reductive stress. Benefiting from the significant difference of bio-reductive enzymes between normal and tumor tissue, various hypoxia-responsive nanocarriers based on 2-nitroimidazole, nitro benzyl derivatives or azobenzene derivatives have been developed. Owing to the cleavable property of azobenzene bond in hypoxic surroundings, azobenzene derivatives have been extensively developed as building blocks in the construction of hypoxia-responsive nanocarriers. For example, Kulkarni et al. synthesized a hypoxia-responsive polymersome composed of poly(lactic acid)-azobenzene-poly(ethylene glycol), which released the loaded drug rapidly in hypoxic tumor spheroid. Yuan et al. prepared biodegradable silica nanocarriers composed of azobenzene bond which achieved specific delivery of the loaded native antibodies in tumor hypoxic environment. Recently, Yang et al. fabricated a hypoxia-degradable albumin-based nanosystem for enhanced tumor penetration and photodynamic-chemo combined therapy. Although plenty of nanocarriers based on azobenzene derivatives have been developed, most of them were focused on micelles, polymersomes or nanocomplexes, and developing a hypoxia-degradable nanogel is still in great demand.

Zwitterionic polymers have emerged as a new class of anti-fouling materials which have been utilized to fabricate nanocarriers for tumor drug delivery. It has been reported that nanocarriers based on zwitterionic polymer exhibit long blood circulation and negligible accelerated blood clearance (ABC) phenomenon. In the family of zwitterionic polymers, phosphorylcholine, carboxybetaine and sulfobetaine polymers are the most widely applied, among which phosphorylcholine polymers have attracted much attention because of the phosphorylcholine head groups similar to the phospholipids in cell membrane. Recently, phosphorylcholine polymers were used to fabricate nanogels with crosslinkers composed of disulfide bond which could achieve the controlled drug release in tumor cytoplasm. However, the super-hydrophilicity feature of phosphorylcholine polymers decreased the interaction with the hydrophobic cell membrane, which led to suboptimal anti-tumor effect. Therefore, it is significant to develop a kind of novel phosphorylcholine polymer nanogel which could be degraded regardless of the cell membrane barrier, thereby enhancing the anti-tumor effect of the drug-loaded nanogel ultimately.

In this work, we synthesized a novel hypoxic-cleavable crosslinker composed of azobenzene bonds and fabricate hypoxic-degradable zwitterionic poly(phosphorylcholine)-based (HPMPC) nanogel for tumor drug delivery, as shown in Fig. 1. The degradability and drug release profiles of the HPMPC nanogel was explored in hypoxic conditions. Furthermore, the blood circulation, biodistribution and anti-tumor effect of drug-loaded HPMPC nanogel were explored in vitro and in vivo. Moreover, the anti-tumor effect between HPMPC nanogel and reduction-degradable poly(phosphorylcholine)-based (HPMPC) nanogel was compared and analyzed. This work may offer new materials and understanding of hypoxic-degradable nanogel in tumor drug delivery.

2. Methods

2.1. Materials

2-Methacryloyloxyethyl phosphorylcholine (MPC, 97%), N,N'-bis(acryloyl)cystamine (BAC, 97%), N-(3-dimethylaminopropyl)-N-ethylcarbodiimide hydrochloride crystalline (EDC, 99%), 4-dimethylaminopyridine (DMAP, 99%), azobenzene-4,4'-dicarboxylic acid (97%), 2-aminoethyl methacrylate hydrochloride (98%), doxorubicin hydrochloride (DOX, 97%), reduced glutathione (GH, 99%), 2,2-azobisisobutyronitrile (AIBN, 99%), cyanine5 maleimide (Cy5-Mal, 97%), N,N-dimethylformamide (DMF, 99%), tetrahydrofuran (THF, 99%), dichloromethane (DCM, 99%) and acetone (AN, 99%) were purchased from Sigma—Aldrich (Shanghai, China). Dulbecco’s modified Eagle’s medium (DMEM), rat liver microsomes and nicotinamide adenine dinucleotide phosphate (NADPH, 97%) were purchased from CHI Scientific (Jiangyin, China). 4’-6-Diamidino-2-phenylindole (DAPI), the immunoglobulin M (IgM) enzyme-linked immunosorbent assay (ELISA) Kit, the immunoglobulin M (IgG) ELISA Kit and 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) were obtained from Beyotime (Shanghai, China).

2.2. Cell lines and animals

HepG2 cell line and 293T cell line were provided by Cell Bank of Shanghai, Chinese Academy of Science (CAS, Shanghai,
China). BALB/c nude mice (aged 5–6 weeks) were purchased from the Institute of Laboratory Animal Science of Jinan University (Guangzhou, China). All the animal experiments were performed according to the protocol of the Institutional Animal Care and Treatment Committee of Jinan University (Guangzhou, China).

2.3. Synthesis of hypoxic-degradable crosslinker

Azobenzene-4,4'-dicarboxylic acid (11.4 mmol), 2-aminoethyl methacrylate hydrochloride (27.3 mmol), EDC (27.3 mmol) and DMAP (2.85 mmol) were dissolved in 100 mL of DMF with the removal of air by nitrogen (Scheme S1). The system was stirred at 25 °C with the speed of 200 rpm for 48 h (Liyida, Suzhou, China). Then, the system was concentrated to 40 mL by rotary evaporation and separated into 4 centrifuge tubes of 50 mL. Each centrifuge tube was added with 40 mL of pure water and centrifuged with the removal of the liquid supernatant. The washing process was repeated for 3 times. The precipitate was dissolved into THF and dried with sodium sulfate. After the purification of silica gel column chromatography (DCM:THF = 19:1), 1.2 g of brown red powder was obtained. The final yield was calculated as 21.4%. 1H NMR (400 MHZ, DMSO-d6) 1.88 (6H, −C−C=−CH3), 3.64 (4H, −CH2−NH−), 4.23 (4H, −O−CH2−), 5.63 (2H, =CH), 6.05 (2H, =CH), 7.92 (4H, Ar), 8.11 (4H, Ar), 8.91 (2H, Ar).

2.4. Preparation of the nanogels

MPC (0.542 mmol), hypoxic-degradable crosslinker (0.162 mmol) and AIBN (0.024 mmol) were dispersed into 40 mL of AN and sonicated for 3 min at room temperature. The system was heated to 100 °C at nitrogen atmosphere with the stirring speed of 200 rpm (Liyida, Suzhou, China). The reaction was carried out for 1 h and cooled to room temperature. The system was centrifuged at 12,000 rpm and washed with pure water for 3 times (Beckman Coulter, Inc., Brea, CA, USA). Finally, the HPMPC nanogel with brown red color was acquired and stored for further use. RPMPC nanogel was obtained using BAC as the crosslinker and the detailed recipe and process were the same as HPMPC nanogel.

2.5. Modification of the nanogels with fluorescent dye

To modify the HPMPC nanogel with fluorescent dye, 5 mg of BAC was added in the preparation of the nanogel. 20 mg of HPMPC or RPMPC nanogels were dispersed in 10 mL of PBS solutions (pH = 7.4) and added with 0.2 mg of TCEP. The reaction was performed for 24 h and added with 2 mg of Cy5-Mal (dissolved in 0.5 mL of DMSO). The reaction was carried out for 12 h under a nitrogen atmosphere to obtain the Cy5-labeled nanogels.

2.6. Stability of the nanogels

HPMPC or RPMPC nanogels (0.5 mg/mL) were dispersed in cell culture medium (DMEM) for 7 days. The hydrodynamic sizes of the nanogels were detected at different time points (0, 1, 2, 3, 4, 5, 6 and 7 days) by dynamic light scattering (DLS, Malvern Nano-ZS90, Malvern, UK).

2.7. Degradability of the nanogels

The turbidity change of the HPMPC nanogel was detected by DLS in 2 mL of PBS with the addition of 100 μL of rat liver microsomes and 100 μmol/L NADPH. Air was bubbled into the

Figure 1  Schematic illustration of the hypoxia-degradable poly(phosphorylcholine)-based (HPMPC) nanogel for tumor drug delivery. HPMPC nanogel shows ultra-long blood circulation without causing the accelerated blood clearance phenomenon, which leads to the enhanced accumulation in tumor tissue. HPMPC nanogel holds the loaded drug firmly in blood or normal tissue whereas releases the drug effectively in tumor hypoxic environment, leading in improved therapeutic effect and negligible side effects.
solutions to create normoxic conditions. For hypoxic conditions, 5% or 1% of oxygen was bubbled into the systems. As a control, the turbidity change of the \(^\text{dPMPC}\) nanogel was detected in 2 mL of PBS with 0, 5 and 10 mmol/L of GSH. The nanogels were incubated in PBS 7.4 (0.01 mol/L) at 37 °C, and the turbidity was detected at different intervals. The relative turbidity was calculated as the ratio of turbidity at each time point to the initial turbidity. GPC (HP Agilent series 1100, Santa Clara, CA, USA) was applied to measure the molecular weight of the degraded product of \(^\text{dPMPC}\) or \(^\text{dPMPC}\) nanogels.

2.8. Drug loading and releasing profiles

The amount of 3 mg of DOX was dissolved in 3 mL of deionized water with the pH of 8.0. 10 mg of \(^\text{dPMPC}\) or \(^\text{dPMPC}\) nanogels were added in the above solutions and vibrated at 37 °C for 24 h. Then, the nanogels were separated by centrifugation with the rotational speed of 12,000 rpm and washed 3 times with pure water to remove the unabsorbed DOX. The supernatant was detected by UV–Visible spectrometry (Perkin–Elmer, Lambda750, Waltham, MA, USA) at the excitation wavelength of 488 nm to calculate the concentration of DOX. DOX loading content and DOX encapsulation efficiency were calculated using the following Eqs. (1) and (2):

1. \[
\text{DOX loading content} = \frac{\text{DOX loaded in nanogel}}{\text{Total weight of DOX}} \times 100
\]

2. \[
\text{DOX encapsulation efficiency} = \frac{\text{DOX encapsulated in nanogel}}{\text{DOX loaded in nanogel}} \times 100
\]

To explore the DOX release behaviors, 2 mL of DOX-loaded \(^\text{dPMPC}\) nanogels (\(^\text{dPMPC}@\text{DOX}\)) with 0.1 mg of DOX was encapsulated in a 12,000 Da dialysis bag and incubated in 100 mL of phosphate buffer saline (pH 7.4) with 5 mL of rat liver microsomes and 100 mmol/L NADPH. Air was bubbled into the solutions to create normoxic conditions. For hypoxic conditions, 1% of oxygen was bubbled into the systems. As a control, the drug release profiles of DOX-loaded \(^\text{dPMPC}\) (\(^\text{dPMPC}@\text{DOX}\)) nanogel was also performed with 0 or 10 mmol/L of GSH. At different time points, 2 mL of solution were extracted and added with 2 mL of initial solution. The cumulative amount of DOX released from the nanogel was detected by UV–Visible spectrometry (Perkin–Elmer, Lambda750).

2.9. Cellular uptake behavior

The amount of \(\sim 1 \times 10^6\) HepG2 cells per dish were seeded in confocal dishes (Corning, New York, NY, USA) and cultivated in a cell culture medium for 24 h. Afterwards, cells were incubated with Cy5-modified \(^\text{dPMPC}@\text{DOX}\) or \(^\text{dPMPC}@\text{DOX}\) at equal amounts of DOX for 1, 2, and 4 h in hypoxic environment. Hypoxia environment or normoxic environment were generated in the cells by cultivating in Biospherix C21 hypoxic chamber supplemented with 1% of oxygen or air, respectively. Cells were rinsed and stained with DAPI and observed with a Nikon laser confocal microscope (C2plus, Nikon, Tokyo, Japan). The cellular uptake of \(^\text{dPMPC}@\text{DOX}\) or \(^\text{dPMPC}@\text{DOX}\) was performed with flow cytometry (Olympus IX71, Tokyo, Japan). HepG2 cells were seeded in 6-well plates at a density of \(1 \times 10^5\) cells per well and incubated for 24 h. Then, cells were cultivated with \(^\text{dPMPC}@\text{DOX}\) or \(^\text{dPMPC}@\text{DOX}\) at equal amounts of DOX for 4 h in hypoxic environment. After that, cells were washed several times with PBS and analyzed by flow cytometry (Olympus IX71, Tokyo, Japan).

2.10. Cytocompatibility and cytotoxicity assay

HepG2 or 293T cells were seeded in 96-well plates and cultured for 24 h. Then, cells were cultivated with blank \(^\text{dPMPC}\) or \(^\text{dPMPC}\) nanogels (0, 50, 100, 200, 500 and 1000 μg/mL) for 24 h. Then, 20 μL of MTT solution (5 mg/mL) was added into the cells and cultured for 4 h. The supernatant was eliminated and 150 μL of DMSO as added into each well which was measured by an automated microplate spectrophotometer (Epoch2, BioTek Instruments, Santa Clara, CA, USA) at 490 nm. The cytotoxicity of \(^\text{dPMPC}@\text{DOX}\) or \(^\text{dPMPC}@\text{DOX}\) against HepG2 cells were also explored by the MTT assay for 24 h in normoxic or hypoxic environment (1% of oxygen). The concentration of DOX was determined as 0, 0.1, 0.5, 1.0, and 5.0 μg/mL.

2.11. Blood circulation profiles and immune compatibility

Fifteen nude female BALB/c mice (~20 g) were separated into three groups with the intravenous administration of free Cy5, Cy5-modified \(^\text{dPMPC}\) and Cy5-modified \(^\text{dPMPC}\) nanogel (500 μg/mL, 100 μL), respectively. Orbital venous blood (50 μL) of each mouse was acquired at different time intervals post injection (0.25, 0.5, 1, 2, 4, 6, 12, 24 and 48 h) \(^{37}\). In Day 5, the blood of each mouse was obtained for the detection of IgM by ELISA. In Day 7, the mice of each group received a second injection as the same dose of the first injection. In the Day 12, the blood of each mouse was obtained for the test of IgG by ELISA. All of the blood samples were treated with three IU of heparin sodium in PBS and centrifuged at 1200 rpm for 10 min (Beckman Coulter, Inc.). The fluorescent intensity of each sample was measured by a fluorescence lifetime spectrometer (QM-40, Horiba Jobin Yvon, Paris, France).

2.12. In vivo fluorescence imaging and biodistribution

Fifteen HepG2 tumor-bearing nude mice were separated into three groups \((n = 5)\) and intravenously injected with 100 μL of free Cy5, Cy5-modified \(^\text{dPMPC}\) and Cy5-modified \(^\text{dPMPC}\) nanogel (500 μg/mL, 100 μL). After the intravenous injection, all the mice were taken the picture by in vivo fluorescence imaging with an optical small animal imaging system (Bruker MI, in vivo Xtreme, Karlsruhe, Germany) at different time intervals (1, 2, 4, 8, 12, 24, 48 and 72 h). All mice were euthanized and the main organs including heart, liver, spleen, lung, kidney and tumor were collected and imaged by the small animal imaging system post-injection for 24 h. In addition, tumor tissues were stained by DAPI and observed by confocal laser scanning microscopy (C2plus, Nikon, Tokyo, Japan).

At 24- and 48-h post-injection of free Cy5, Cy5-modified \(^\text{dPMPC}\) and Cy5-modified \(^\text{dPMPC}\) nanogel (500 μg/mL, 100 μL), HepG2 tumor-bearing mice in each group \((n = 5)\) were euthanized with the main organs (heart, liver, spleen, lung and kidney) and tumor extracted. The fluorescence intensities of all the samples were measured to determine the distribution of nanogels in each organs.

2.13. In vivo anti-tumor efficacy

Twenty HepG2 tumor-bearing nude mice (~20 g) were randomly separated into four groups \((n = 5)\). After the tumor volume of arrives ~100 mm\(^3\), four groups of mice were received with PBS,
free DOX, HPMPC@DOX and HPMPC@DOX on Days 0 and 6 for each injection at a DOX dose of 2 mg/kg body weight, respectively. The tumor size and body weight of the mice were tested every other day. On Day 14, all of the mice were euthanized and tumors were dissected and weighed. Histological analysis was performed on major organs by hematoxylin-eosin (HE) staining.

2.14. Blood biochemistry and blood routine test

To explore the biosafety of blank nanogels, 15 female BALB/c mice (~20 g) were randomly divided into three groups as control, RPMPC and HPMPC. Ten of the mice were intravenously injected with 100 μL of RPMPC or HPMPC nanogels (2 mg/mL), respectively. Post-injection of 24 h, blood samples were collected from each mouse for biochemical and routine blood testing.

2.15. Statistical analysis

All data were showed as mean ± standard deviation (SD). Unpaired Student’s t-test was conducted for the comparison of two testing groups and a probability (P) less than 0.05 was regarded as statistical significance.

3. Results and discussions

3.1. Synthesis and characterization of hypoxic-degradable crosslinker and nanogel

Hypoxia changes the tumor microenvironment through the over-expressed HIF-1α, which leads to the cancer progression and metastasis. It has been reported that the azobenzene group could be cleaved with the elevated levels of reducing enzymes in hypoxic tumor microenvironment. Furthermore, hypoxic environment exhibits in liver cancer model which was proved by the previous literatures. In this work, azobenzene-contained crosslinker was synthesized by the amide reaction between azobenzene-4,4'-dicarboxylic acid and 2-aminoethyl methacrylate hydrochloride. The obtained crosslinker showed a brown red color with the yield of 21.4%. The structure of the crosslinker was confirmed by the 1H NMR and 13C NMR, respectively (Fig. 2A and B). In addition, mass spectrum showed that the molecular weight of the crosslinker was regarded as 493.20 g/mol, which agreed well with the theoretical molecular weight (492.20 g/mol, Supporting Information Fig. S1). Furthermore, the FT-IR spectra of the hypoxic-degradable crosslinker revealed that there were clear peaks at 1452 and 1558 cm⁻¹, which belongs to the stretching vibration of N=N groups and benzene groups in the crosslinker, respectively. Lastly, the band at 3302 cm⁻¹ was ascribed to the stretching vibration of N—H in the crosslinker. The above data demonstrated that the azobenzene-contained crosslinker was successfully synthesized.

Zwitterionic phosphorylcholine polymer has been demonstrated as a class of biocompatible and anti-fouling material, which was widely used in biomedical applications. Herein, HPMPC nanogel was fabricated by the copolymerization of MPC and azobenzene-contained crosslinker with reflux precipitation polymerization. As a control, RPMPC nanogel was synthesized by the copolymerization of MPC and BAC with reduction-cleavable disulfide bonds. From transmission electron microscopy (TEM), it was found that HPMPC nanogel showed uniform spherical shape with the mean diameter of 115 nm while HPMPC nanogel exhibits the mean diameter of 115 nm while HPMPC nanogel exhibits the
diameter of 118 nm (Fig. 2C and D). In the meantime, the hydrodynamic size of HPMP was 158 nm while RPMPC nanogel displayed the similar size of 163 nm (Fig. 2E). The hydrodynamic size of HPMP nanogel was larger than its TEM diameter owing to the swollen of the nanogel in aqueous solution. Moreover, the zeta potentials of both HPMP and RPMPC nanogels were nearly neutral due to the equal positive and negative charge (Supporting Information Fig. S3). In addition, the hydrodynamic sizes of HPMP and RPMPC remain nearly unchanged after incubation in cell culture medium for seven days, which indicated the favorable stability of the nanogels (Supporting Information Fig. S4).

3.2. Degradation and drug releasing profiles of the nanogels

To explore the hypoxia-degradability of the HPMP nanogel, the relative turbidity of the nanogel solutions were measured by DLS with air flow, 5% and 1% of oxygen. It was showed that the relative turbidity of HPMP nanogel remains nearly 100%, which indicated the high stability of HPMP nanogel in normoxic conditions. In 5% of oxygen, the relative turbidity of HPMP nanogel drops to 87.0% after 2 h, which demonstrated the partial degradation of the nanogel (Fig. 3A). Furthermore, the relative turbidity of HPMP nanogel reduced to 5.8% after 2 h in 1% of oxygen, indicating the sufficient degradation of HPMP nanogel in hypoxic condition. The degradation behaviors of RPMPC nanogel in different concentrations of GSH were explored. The results showed that RPMPC nanogel could be completely degraded after 50 min in 10 mmol/L GSH, which exhibited a faster degradation speed compared with HPMP nanogel in hypoxic condition (Fig. 3B). Moreover, the degradation products of HPMP and RPMPC nanogels were collected and analyzed by GPC. It was found that the molecule weight of HPMP nanogel after degradation was calculated as 2100 g/mol, which was close to that of RPMPC nanogel after degradation (Fig. 3C). The above results demonstrated that HPMP nanogel could be efficiently degraded into oligomers with low molecule weight in hypoxic condition.

DOX was selected as the model drug to test the drug loading and releasing properties of the HPMP nanogel. It revealed that the DOX loading and encapsulation efficiency of HPMP nanogel were calculated as 7.7% and 25.6%, respectively, which were close to those of RPMPC nanogel (7.4% and 24.7%, respectively). Furthermore, the drug-releasing profiles of HPMP@DOX were tested in normoxic and hypoxic conditions, respectively. It was found that HPMP@DOX holds the loaded DOX firmly with only 8.9% of leakage in normoxic condition after 48 h (Fig. 3D). However, HPMP@DOX showed a rapid DOX release in hypoxic condition and 86.3% of the loaded DOX were released, which was attributed to the degradation of the azobenzene bond in hypoxic condition. RPMPC@DOX exhibited an even faster DOX release profile in reduction environment (10 mmol/L GSH) with 94.2% of loaded DOX released after 48 h, which was ascribed to the faster degradation speed of RPMPC in 10 mmol/L GSH compared to HPMP in 1% of oxygen.

3.3. Cellular uptake and cytotoxicity of the nanogels

Hydrophilicity is a basic requirement for nanocarriers in tumor drug delivery, which could increase the stability of nanocarriers in vivo and reduce the protein corona on the surfaces. However, tumor cell membranes are usually hydrophobic, which decrease the cellular uptake of the hydrophilic nanocarriers into
**Figure 4**  Confocal laser scanning microscopy observation of DOX loaded \(^{1}H\)PMPC (A) and \(^{8}P\)MPC (B) in hypoxic environment. Flow cytometry analysis (C) and quantitative analysis (D) of the cellular uptake of \(^{1}H\)PMPC and \(^{8}P\)MPC with different treatment. (E) Cytotoxicity of DOX loaded \(^{1}H\)PMPC and \(^{8}P\)MPC to HepG2 cells with air flow or 1% of oxygen. Scale bar = 50 μm **P < 0.01 and ***P < 0.001.

**Figure 5**  *In vivo* pharmacokinetics of free Cy5, \(^{8}P\)MPC and \(^{1}H\)PMPC nanogels after the 1st intravenous injection (A) and (B) the 2nd intravenous injection in BALB/c mice. (C) The detection of immunoglobulin M (IgM) levels *in vivo* after the 1st injection. (D) The detection of immunoglobulin G (IgG) levels *in vivo* after the 2nd injection. The distribution of \(^{8}P\)MPC and \(^{1}H\)PMPC nanogels in the main organs and tumors at (E) 24 h and (F) 48 h post-injection. *n.s.*: no significant difference.
the tumor cells\textsuperscript{47,48}. Compared with other stimuli signals in tumor cells such as GSH, hypoxic conditions exist both in intracellular and extracellular environments in tumor tissue, which may be beneficial for the sufficient drug releasing of \textsuperscript{8}PMPC@DOX. The HepG2 cellular uptake of the \textsuperscript{8}PMPC@DOX was firstly evaluated by the CLSM observation. Fig. 4A revealed that red fluorescent signals of DOX, which were mainly focused on the cell nucleus, were increasingly enhanced with the extension of incubation time. On the contrary, negligible fluorescent signals were observed incubation of \textsuperscript{8}PMPC@DOX in normoxic environment, owing to the nondegradable property of \textsuperscript{8}PMPC in the air (Supporting Information Fig. S5). Given that \textsuperscript{8}PMPC@DOX with the hydrodynamic size larger than the pore diameters of cell nucleus, it could be concluded that \textsuperscript{8}PMPC@DOX was effectively degraded in hypoxic condition \textit{in vitro}. Meanwhile, Fig. 4B showed that \textsuperscript{8}PMPC@DOX exhibits significant weaker fluorescent signals compared to those of \textsuperscript{10}PMPC@DOX, which was attributed to the suboptimal cellular uptake and insufficient drug release of \textsuperscript{8}PMPC@DOX, since \textsuperscript{8}PMPC could only be degraded after the cellular uptake. Flow cytometry was further employed to quantitatively explore the cellular uptake of \textsuperscript{10}PMPC@DOX and \textsuperscript{8}PMPC@DOX (Fig. 4C). Fig. 4D showed that the DOX internalization of \textsuperscript{10}PMPC@DOX was significantly higher than that of \textsuperscript{8}PMPC@DOX, which agreed well with the CLSM results. Therefore, it could be concluded that hypoxia-degradable nanogel showed superior drug releasing ability regardless of the barrier of cell membranes compared to the reduction-degradable analogue.

The cytotoxicity of blank \textsuperscript{10}PMPC and \textsuperscript{8}PMPC nanogels were explored from the concentration of \textit{0–1000 \textmu g/mL}. It was found that both the \textsuperscript{10}PMPC and \textsuperscript{8}PMPC nanogels showed negligible toxicity to HepG2 cells and 293T cells at the high concentration of 1000 \textmu g/mL, which indicated the favorable biocompatibility of the nanogels (Supporting Information Figs. S6 and S7). Furthermore, the inhibition effects of \textsuperscript{10}PMPC@DOX and \textsuperscript{8}PMPC@DOX to HepG2 tumor cells were explored at hypoxic or normoxic conditions. Fig. 4E revealed that \textsuperscript{10}PMPC@DOX exhibits significantly stronger inhibition effect to tumor cells at hypoxic condition compared to normoxic condition, owing to the rapid drug release at hypoxic condition. More interestingly, \textsuperscript{10}PMPC@DOX showed superior tumor-killing effect than \textsuperscript{8}PMPC@DOX at hypoxic condition which was attributed to the complete drug releasing ability of \textsuperscript{10}PMPC@DOX regardless of

![Figure 6](image-url)
the cellular uptake. Therefore, the above results demonstrated that hypoxic-degradable \textsuperscript{1}PMPC nanogel displayed better anti-tumor effect \emph{in vitro} compared to the reduction-degradable analogue.

3.4. Pharmacokinetics, immune response and biodistribution of the nanogels

The blood circulation behaviors of the nanogels were explored by detecting the fluorescent signal intensity in blood after intravenous injection of free Cy5, Cy5-modified \textsuperscript{1}PMPC and Cy5-modified \textsuperscript{1}PMPC. Fig. 5A showed that both \textsuperscript{1}PMPC and \textsuperscript{1}PMPC nanogels exhibit ultra-long blood circulation compared to free Cy5, which was consistent well with the previous reports\textsuperscript{49,50}. Furthermore, the blood retention of \textsuperscript{1}PMPC (\(t_{1/2} = 45.7\) h) and \textsuperscript{1}PMPC (\(t_{1/2} = 44.2\) h) were very close after intravenous injection due to the similar hydrodynamic sizes and surface properties of the nanogels. In addition, the pharmacokinetics profiles of \textsuperscript{1}PMPC and \textsuperscript{1}PMPC nanogels were nearly unchanged after the second injection, demonstrating that the repeated injection of \textsuperscript{1}PMPC and \textsuperscript{1}PMPC nanogels would not induce the ABC phenomenon (Fig. 5B). The immune response of mice was evaluated by detecting the IgM and IgG level in blood after intravenous injection of \textsuperscript{1}PMPC or \textsuperscript{1}PMPC nanogels. Fig. 5C displayed that the administration of \textsuperscript{1}PMPC or \textsuperscript{1}PMPC nanogels did not induce the significant rise of IgM level in blood. Similarly, Fig. 5D showed that both \textsuperscript{1}PMPC and \textsuperscript{1}PMPC nanogels did not induced the rise of IgG level, which indicated the desirable immune compatibility of the nanogels. The biodistribution of the nanogels were explored by detecting the fluorescent signal intensity in the major organs (heart, liver, spleen, lung and kidney) and tumor. Fig. 5E revealed that both \textsuperscript{1}PMPC and \textsuperscript{1}PMPC nanogels could effectively accumulate in tumor tissue post-injection for 24 h due to the well-known enhanced permeation and retention effect. Furthermore, both \textsuperscript{1}PMPC and \textsuperscript{1}PMPC nanogels still remained high tumor accumulation post-injection for 48 h which was ascribed to the long blood circulation of the nanogels (Fig. 5F).

3.5. Near-infrared fluorescence imaging of the nanogels

To intuitively observe the tumor accumulation of the nanogels, near-infrared fluorescence imaging (NIFR) was carried out at varied time points post-injection of free Cy5, \textsuperscript{1}PMPC and \textsuperscript{1}PMPC. Fig. 6A showed that negligible fluorescent signals appeared at tumor tissue in the group of free Cy5 post-injection for 72 h, which demonstrated that free Cy5 could not effectively accumulate at tumor tissue. Meanwhile, bright red fluorescent signals were observed at tumor tissue post-injection of \textsuperscript{1}PMPC and \textsuperscript{1}PMPC nanogels, demonstrating the high and long-lasting accumulation of the nanogels at tumor tissue. Furthermore, the tumor accumulation of \textsuperscript{1}PMPC and \textsuperscript{1}PMPC nanogels did not

![Figure 7](image)

**Figure 7** Anti-tumor effect of drug-loaded nanogels. (A) Tumor growth curves, (B) tumor weights and (C) tumor photograph of the mice after injection with PBS, free DOX, \textsuperscript{1}PMPC@DOX and \textsuperscript{1}PMPC@DOX. (D) Weight variation of HepG2 tumor-bearing mice after injection with PBS, free DOX, \textsuperscript{1}PMPC@DOX and \textsuperscript{1}PMPC@DOX. *\(P < 0.05\), **\(P < 0.01\) and ***\(P < 0.001\).
present significant difference, which agreed well with the biodistribution experiments (Fig. 6B). Furthermore, mice were euthanized after injection for 24 h with major organs (heart, liver, spleen, lung, kidneys, and tumor) imaged by fluorescence imaging of live animals. Most of the fluorescent signals accumulated in tumors in $^4{\text{PMPC}}$ or $^4{\text{PMPC}}$ groups, demonstrating high tumor accumulation of the nanogels (Fig. S8). Moreover, CLSM observation of tumor slices post-injection for 24 h showed that strong red fluorescent signals were observed in the groups of $^8{\text{PMPC}}$ or $^8{\text{PMPC}}$, whereas negligible red fluorescent signal was found in the group of free Cy5, which indicated the specific tumor accumulation ability of the nanogels (Fig. 6C).

3.6. Anti-tumor effect and biosafety of the nanogels

To further evaluate the anti-tumor effect of DOX-loaded nanogels, HepG2 tumor-bearing mice were treated with PBS, free DOX, $^8{\text{PMPC}}@\text{DOX}$, and $^8{\text{PMPC}}@\text{DOX}$, respectively. Fig. 7A showed that the PBS group had rapid tumor growth, whereas the free DOX group exhibited a moderate tumor inhibition effect with the tumor volume of 586.1 mm$^3$ after 14 days. Furthermore, $^4{\text{PMPC}}@\text{DOX}$ displayed a significant tumor inhibition effect, which was ascribed to the high tumor accumulation and reduction degradability of the samples. Moreover, the strongest tumor growth inhibition effect was observed in the group of $^4{\text{PMPC}}@\text{DOX}$. The superior anti-tumor effect of $^4{\text{PMPC}}@\text{DOX}$ than $^8{\text{PMPC}}@\text{DOX}$ was attributed to the more sufficient DOX release of $^4{\text{PMPC}}@\text{DOX}$ in tumor tissue than $^8{\text{PMPC}}@\text{DOX}$. As mentioned above, $^8{\text{PMPC}}@\text{DOX}$ could only release the loaded DOX after the cellular uptake by tumor cells with limited efficiency, which impaired the overall anti-tumor effect of $^8{\text{PMPC}}@\text{DOX}$. As to $^4{\text{PMPC}}@\text{DOX}$, DOX could be effectively released from the nanogels regardless of the cell membrane barriers, leading to better anti-tumor effect. After the treatment of 14 days, all the mice were sacrificed; tumors were dissected, weighed, and imaged (Fig. 7C). The tumor weights for the PBS, free DOX, $^8{\text{PMPC}}@\text{DOX}$, and $^4{\text{PMPC}}@\text{DOX}$ were $0.83 \pm 0.055$, $0.52 \pm 0.048$, $0.11 \pm 0.028$ and $0.052 \pm 0.016$ g, respectively (Fig. 7B). To explore the biosafety of the treated groups in vivo, the treated mice were weighed every other day. As shown in Fig. 7D, only the free DOX group induced significant body weight loss, owing to the severe cardiotoxicity of free DOX to normal cells, which indicated that the encapsulation of DOX into zwitterionic phosphorylcholine nanogels reduces the toxicity of free DOX.

H&E staining of major organs (heart, liver, spleen, lung and kidney) was further performed to evaluate the possible side effects of the nanogels. Fig. 8D uncovered that no significant tissue damage was observed in the $^8{\text{PMPC}}@\text{DOX}$ and $^4{\text{PMPC}}@\text{DOX}$ groups. Slight myocardial fiber rupture was observed in free DOX.
group, maybe owing to the cardiotoxicity of DOX. Moreover, the biosafety of the nanogels was evaluated by the blood biochemistry and routine blood tests. Fig. S8A–C revealed that no significant difference was found among the mice in the PBS, HPMC and HPMPC group. Therefore, the above results demonstrated that HPMPC nanogel showed favorable biocompatibility and biosafety in vivo.

4. Conclusions

In summary, a type of hypoxia-degradable poly(phosphorylcholine)-based nanogel was rationally designed for tumor hypoxia-triggered drug release. The obtained HPMPC nanogel displayed desirable long blood circulation and negligible immune response, which led to the enhanced tumor accumulation. Impressively, the HPMPC nanogel could be effectively degraded and release the loaded drug in tumor tissue regardless of the cell membrane barrier which was encountered by the reduction-degradable analogue. Therefore, drug-loaded HPMPC nanogel presented superior tumor inhibition effect both in vitro and in vivo compared to the drug-loaded HPMPC nanogel. Lastly, HPMPC nanogel showed favorable biocompatibility and biosafety in vivo. Therefore, this work provided a novel hypoxia- or pH-sensitive crosslinker which could be employed to synthesize hypoxia-degradable nanogels and offered new understanding of hypoxia-degradable nanogels in tumor drug delivery.

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Author contributions

Ligong Lu, Shaojun Peng and Shun Shen designed the research. Shaojun Peng and Boshu Ouyang carried out the experiments and performed data analysis. Yongjie Xin and Meixiao Zhan participated part of the experiments. Wei Zhao provided experimental drugs and quality control. Shaojun Peng wrote the manuscript. Shun Shen revised the manuscript. All of the authors have read and approved the final manuscript.

Conflicts of interest

The authors have no conflicts of interest to declare.

Appendix A. Supporting information

Supporting data to this article can be found online at https://doi.org/10.1016/j.apsb.2020.08.012.

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