Rational design of a nitroreductase-activatable two-photon fluorescent probe for hypoxia imaging in cell and in vivo

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1. Introduction

Hypoxia has been considered as an inherent feature of solid tumor, whereby the oxygen (O₂) concentration ranges from 4.0 % to 0 %. [1] Tumor hypoxia has been known as a pathophysiological consequence of deteriorating microenvironment, and it is mainly derived from the rapid and exaggerated growth of tumor cells [2]. Hypoxia can lead to structures, functions and diffusion conditions of tumors disturbed, and it plays significant roles in tumor invasion, metastasis and drug resistance [3]. Researchers report that hypoxia causes many tumor-associated genes overexpress, especially hypoxia inducible factors 1α (HIF-1α) and HIF-mediated genes [4,5,48]. Within the hypoxic tumor microenvironment, HIF-1α promotes angiogenesis, metabolic adaptation, and other key aspects of tumor progression, which results in deteriorating tumor hypoxic microenvironment and abnormal microvessels. [5] These a series of behaviors limit the perfusion of cytotoxic chemotherapeutic drugs, and further reduce the efficacy of chemotherapeutic drugs as well as lead to tumor resistance [5]. Hyperbaric oxygen (HBO) can alleviate tumor hypoxia by increasing oxygen partial pressure, and it has been used in many studies for seeking to reverse hypoxic conditions of tumor [6]. HBO can significantly increase intracellular oxygen partial pressure to overcome hypoxia-induced chemotherapy and radiotherapy resistance, as well as improve drug treatment efficacy. [7-9] Thus HBO-assistant chemotherapy may be a promising strategy for malignant tumor treatment. And powerful tools are needed urgently for assessing the therapeutic effect of malignant tumors.

Nitroreductase (NTR) is an endogenous enzyme that overexpressed in hypoxic tumors, and the concentration of NTR is directly concerned to the degree of hypoxia. [10] NTR is seen as an indicator of highly invasive diseases in various hypoxic tumors and NTR overexpression plays important roles in tumor invasion, progression and angiogenesis [11]. Further studies show the hypoxic status of solid tumors is closely related to the process of tumor progression [12]. Therefore, NTR

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ABSTRACT

Tumor resistance is a huge challenge for tumor treatment, which may lead to tumor treatment failure. Relieving tumor resistance and improving efficacy are long-term goals for tumor treatment. Nitroreductase (NTR) is an endogenous enzyme that highly expressed in hypoxia tumors. Herein, we develop a NTR-activatable fluorescent probe, TP-NO₂, for NTR detection during tumor treatment. TP-NO₂ with simple synthesis steps and high yield can qualitatively and quantitatively detect NTR in a wide range (0 – 20 μg/mL) with a detection limit of 26 ng/mL. The probe has been successfully applied for NTR detection in cells under simulated hypoxia conditions. Hyperbaric oxygen (HBO) can alleviate tumor hypoxia and reduce NTR concentration. We pre-evaluate the effect of tumor treatment by NTR imaging. And the results demonstrate that HBO-assistant chemotherapy can effectively treat tumor cells. We further apply TP-NO₂ for NTR detection in A549 and cis-di-chlorodiamineplatinum(II) (DDP)-resistant A549 (A549/DDP) xenograft nude mice, and we choose NTR as an indicator to pre-evaluate the treatment efficacy of malignant tumors. With the adjuvant therapy of HBO, chemotherapeutic drugs gemcitabine and carboplatin can effectively treat A549 and A549/DDP xenograft. These applications provide us a novel perspective for NTR imaging.
can be selected as a biomarker and the hypoxia degree of solid tumor can be assessed by evaluating the NTR level [13]. NTR is a kind of flavin-dependent enzyme, which can reduce nitroaromatic compounds and nitroheterocyclic derivatives to primary amines through ping-pong mechanism with two electrons transferring, and nicotinamide adenine dinucleotide hydrogen (NADH) is used as the electron donor [14]. Inspired by this reaction, designing potential molecular tools for tumor hypoxia detection are needed urgently.

Traditional hypoxia detection methods include positron emission tomography technique, pO2 electrodes, magnetic resonance imaging and multispectral optoacoustic tomography (MSOT) imaging. [15–19] However, these methods usually require complex sample pretreatment, expensive consumption, destruction of tissues and cells. NTR-activatable fluorescent probe technology has received great attentions for its advantages such as less invasiveness, highly selectivity, rapidly response, highly spatial and temporal resolution, real-time and in situ detection. [20,46,47] Recently, two-photon fluorescent probes using two near-infrared (NIR) photons as the excitation source can achieve higher spatial and temporal resolution, deeper tissue imaging depth and longer observation time [21,51,52]. However, there are fewer two-photon fluorescent probes for NTR detection available so far, and the developed two-photon fluorescent probes usually have short fluorescence emission [21–25]. Thus powerful two-photon fluorescent probes with longer fluorescence emission wavelength are required urgently for NTR detection in tumor.

Here, we designed a NTR-activatable fluorescent probe TP-NO2 with long fluorescence emission for NTR detection. TP-NO2 was designed based on a two-photon fluorophore TP-OH and a responsive group p-nitrobenzene. Our probe had highly selectivity and sensitivity for NTR detection. Using TP-NO2, we can explore the relationship between NTR and multispectral optoacoustic tomography (MSOT) imaging. [15] For NTR detection, many one-photon fluorescent probes are currently rare. Herein, we designed a two-photon fluorescent probe for NTR detection bearing nude mice until the tumor volumes typically reached to about 200 mm³. All experiments were performed in accordance with the guidelines established by the Committee of Animal Research Policy of Binzhou Medical University.

2. Experimental section

2.1. Cell culture

Human lung carcinoma (A549) cells and human neuroblastoma (SH-SY5Y) cells were purchased from the Committee on Type Culture Collection of Chinese Academy of Sciences (Shanghai, China). A549 cells were cultured with DMEM supplemented with 10 % FBS. SH-SY5Y cells were cultured with RPMI-1640 supplemented with 10 % FBS. All cells were incubated at 37 °C under a humidified atmosphere containing 5 % CO2. The cells were passaged by scraping and seeding on 20 mm Petri-dishes according to the instructions from the manufacturer. A multi gas incubator (Sanyo) was used for producing 1–21 % O2 concentration by means of N2 substitution. And 0.1 % O2 concentration was generated with an AnaeroPac™ (Mitsubishi Gas Chemical Campany, Co. Inc., Japan).

2.2. Confocal imaging

Cells were plated on Petri-dishes (Φ = 20 mm) and allowed to adhere for 24 h before imaging. The probe was added to the culture plates which were filled with 1 mL fresh complete medium. One-photon fluorescent images were acquired on an Olympus Fluoview FV1000 confocal laser-scanning microscope (Japan). The collection wavelength was λex = 530 - 630 nm (λem = 440 nm). The two-photon fluorescent images were collected using a two-photon confocal microscope (Zeiss LSM 880). The collection wavelength was λex = 530–630 nm (λem = 840 nm).

2.3. Establishment of the A549 and A549/DDP transplanted tumor nude mice

5-week-old specific pathogen free nude mice were provided by Changzhou Caverns Lab Animal Co. Ltd. The mice were housed in individual ventilated cages, and a SPF laboratory diet and water were acquired freely. Mice were group-housed on a 12:12 light-dark cycle. 2 × 10⁶ cells were suspended in media and implanted subcutaneously into nude mice. A549 and A549/DDP xenografts were established in nude mice until the tumor volumes typically reached to about 200 mm³. All experiments were performed in accordance with the guidelines established by the Committee of Animal Research Policy of Binzhou Medical University.

2.4. Mice imaging in vivo

A Bruker In-vivo Imaging System was employed to imaging tumor bearing nude mice. The excitation and emission wavelengths were chosen as 465 nm and 560 nm. The mice were anesthetized prior to injection and during imaging. After in vivo imaging, the organs (lung, heart, liver, kidney and spleen) and tumors were excised to perform ex vivo imaging.

2.5. Synthesis of compounds

Compounds were synthesized according to the general procedure and characterization details were described in the Supporting Information.

Synthesis of compound TP-NO2. Compound TP-OH (0.0342 g, 0.1 mmol) and K₂CO₃ (0.0691 g, 0.5 mmol) were dissolved in DMF (30 mL) for 30 min, then 4-nitrobenzyl bromide (0.0432 g, 0.2 mmol) was added overnight. After concentrated, the obtained crude product was purified by silica column chromatography (200–300 mesh) to afford an orange product (0.73 g, 73.5 %). ¹H NMR (500 MHz, DMSO-D₆) δ (ppm): 0.82-0.93 (m, 3 H), 1.29 (s, 6 H), 3.84—4.02 (d, 2 H), 4.44—4.70 (s, 2 H), 7.03–7.15 (s, 1 H), 7.37–7.45 (m, 1 H), 7.48–7.56 (d, 2 H), 7.57–7.70 (m, 2 H), 7.70–7.77 (t, 2 H), 7.77–7.97 (m, 4 H), 7.98–8.09 (s, 1 H), 8.35–8.46 (s, 3 H). LC-MS (API-ES): m/z C₃₁H₂₉N₂O₃⁺ Calcd 477.57, found [M⁺] 477.43.

3. Results and discussion

3.1. Design and synthesis of TP-NO2

For NTR detection, many one-photon fluorescent probes have been developed. [4,10,26,27] However, two-photon fluorescent probes are currently rare. Herein, we designed a two-photon fluorescent probe for NTR detection. We selected two-photon fluorophore TP-OH as fluorescence reporter [28]. After fluorescence reporting unit was selected, we linked p-nitrobenzene, an classical identification unit for NTR detection, to the fluorophore through an ether linkage (Scheme 1). The p-nitrobenzene group was known to decrease fluorescence emission via a donor excited photo-induced electron transfer (d-PET) process. As expected, probe TP-NO2 showed no fluorescence emission due to p-nitrobenzene quenched the fluorescence of TP-NO2. After reaction with NTR, p-nitrobenzene unit was reduced to aminobenzene group with NADH as a cofactor. Then, TP-NO2 experienced 1, 6-rearrangement elimination reaction to give a turn-on fluorescence signal at 880 nm, which ascribed to the PET suppression mechanism. And the reaction mechanism was confirmed using HPLC (Figure S13). Compared to the
developed two-photon fluorescent probe, probe TP-NO2 showed a large two-photon action cross section (S) value of 126 GM at 840 nm (Figure S15), good selectivity and sensitivity with a low detection limit of 26 ng/ml, high molar absorption coefficient, high yield as well as long fluorescence emission wavelength that can avoid autofluorescence from organisms (Table S1). [21–25] The detailed synthesized steps were shown in Scheme S1.

3.2. Spectral properties of probe TP-NO2 towards NTR

The spectral properties of TP-NO2 were detected in 10 mM HEPES buffer (simulated physiological conditions, pH 7.4) with 0.5 mM NADH as cofactor. Our probe showed sensitive spectral responses towards NTR (0–20 μg/ml) (Fig. 1a). TP-NO2 showed faint absorption in the absence of NTR. Upon NTR was added, increased absorption spectrum can be observed at 450 nm (ε = 8.2 × 10^3, φ = 0.001). Then, the fluorescence spectrums were detected (Fig. 1b). TP-NO2 exhibited no fluorescence emission in the absence of NTR. Strong fluorescence emission at 580 nm can be observed upon NTR was added (ε = 5.7 × 10^4, φ = 0.049). There was a good linear relationship between the fluorescence intensity and NTR concentration (0–20 μg/ml). The linear regression equation was $F_{\text{ex/em}}(450/580\text{nm}) = 7.916 × 10^4 \times [\text{NTR}] + 4.330 × 10^4$ with $r^2 = 0.9925$ (Figure S1b). The limit of detection was determined to be 26 ng/ml (3σ/ε) indicated that TP-NO2 had a high sensitivity for NTR detection.

The reaction kinetic was a critical parameter for applications of the probe. Then the reaction kinetic of TP-NO2 towards NTR was elucidated in the presence of NADH. As shown in Fig. 1c, the reaction rate increased with the concentration of TP-NO2 increased. In addition, the kinetic parameters of the NTR-catalyzed cleavage reaction were also detected. Fig. 1c showed a Lineweaver-Burk double-reciprocal plot of $1/V$ (V was the initial reaction rate) versus the reciprocal of the probe TP-NO2 concentration. The equation was described as: $V = V_{\max} [\text{probe}] / (K_m + [\text{probe}])$, where V was the reaction rate, [probe] was the probe concentration, and $K_m$ was the Michaelis constant. According to the equation, the calculated values of $K_m$ and $V_{\max}$ were determined to be 46.82 μM and 0.13 μM/S respectively.

3.3. NTR detection in hypoxia cells

Encouraged by the good spectral response of TP-NO2 towards NTR in Fig. 1, we further performed NTR detection in living cells. Before applied for cells imaging, MTT assays were performed for evaluating the cytotoxicity of TP-NO2. As shown in Figure S3, the high cells viability illustrated that our probe displayed low cytotoxicity towards living cells. Then, the fluorescence imaging of NTR in living cells was performed by laser scanning confocal microscope. As shown in Fig. 2, we selected A549 cells and SH-SY5Y cells as cell models for NTR imaging, and the cells were divided into six groups with different oxygen concentration (21 %, 10 %, 5 %, 3 %, 1 % and 0.1 %) for 6 h. All the cells were incubated with TP-NO2 (10 μM) for 25 min before imaging. The detailed cell processing processes were shown in experimental section. As shown in Fig. 2a, our probe showed faint fluorescence under normoxic condition (21 % O2), which indicated that the level of NTR was low under normal condition. The fluorescence intensities increased gradually with the concentration of O2 decreased, which implied that the level of NTR increased. Thus, the degree of hypoxia can be evaluated by NTR detection. [10] Flow cytometry analysis has been considered to be a high-throughput assay technology, and it has been widely used for analysis of large number of samples. Then, we performed flow cytometry analysis to confirm the fluorescence imaging results (Fig. 2d). The results from flow cytometry analysis were consistent with that from laser confocal imaging, and the fluorescence intensity increased with the O2 concentration decreased. Corresponding fluorescence intensity analysis of Fig. 2a and d were shown in Fig. 2f and h. Dicoumarin was a well-known inhibitor for NTR. [4,48] Then we incubated cells with 100 μM dicoumarin to inhibit NTR for 2 h. As shown in Fig. 2b, cells showed extremely weak fluorescence signals after incubated with dicoumarin, which further confirmed the selectivity of TP-NO2 for NTR. And we also captured faint fluorescence signals using flow cytometry (Fig. 2e). The fluorescence intensity analysis of Fig. 2b and e were shown in Fig. 2g and i. These results indicated that our probe TP-NO2 can be applied for hypoxia detection in cells via endogenous NTR detection. We also performed NTR detection using two-photon confocal microscope in Fig. 2c, and the corresponding fluorescence intensity analysis was shown in Fig. 2j and 2k. Using two-photon confocal microscope, we can capture bright fluorescence in hypoxia conditions. After treated with dicoumarin, the fluorescence signal was decreased. Thus, the results from one-photon confocal microscope and two-photon confocal microscope were consistent well.

3.4. NTR detection during the process of tumor cell treatment

Hyoxia is a typical feature of tumors. It has been reported that tumor treatment while relieving tumor hypoxia may improve the efficacy of radiotherapy, chemotherapy and immunotherapy [29]. Then we...
attempted to investigate the auxiliary treatment effect of HBO during chemotherapy. NTR is an endogenous enzyme and closely related to oxygen concentration. The results of cell experiments indicated that the hypoxia degree of tumor cells can be evaluated by NTR detection \[4,30,48\]. Since our probe TP-NO2 had been used for NTR detection successfully under different O2 concentrations, we attempted to evaluate the NTR concentration fluctuations during the process of tumor treatment. A549 and SH-SY5Y cells were selected as cell models and the cells in Fig. 3a were divided into five groups. The detailed cell processing processes were shown in Supporting Information. The cells in the control group were manipulated under normoxic conditions, and they showed weak fluorescence signals. Cells in the next group were treated under hypoxia (0.1 % O2) for 6 h, and intense fluorescence signals were recorded. Next, we incubated cells under HBO (60 % O2) for 6 h, and decreased fluorescence intensity can be observed. Gemcitabine and carboplatin have been widely used as chemotherapeutic drug for tumor treatment \[31,32,49\]. Carboplatin can cross-link to DNA and inhibit replication and transcription of tumor cells through forming
Fig. 3. NTR detection during the process of tumor treatment. (a) Laser confocal microscope imaging. (b) Flow cytometry analysis. (c) Ca$^{2+}$ concentration analysis by Fluo 4-AM. (d) $\Delta\Psi_m$ analysis by JC-1. (e) TEM observation (Scale bars: 1 μm). (f) Western blotting analysis of HIF 1α, Hsp70, cytochrome c, Bax, Bcl-2, pro-caspase 3 and cleaved-PARP. β-Actin was used as loading control. (g) Relative fluorescence quantitative analysis for (a). (h) Fluorescence quantitative analysis for (b). (i) Quantitative analysis for (f). (j) Signal transduction pathways. The experiments data were shown as mean (± s.d.) (n = 7).
intra- or interstrand adducts [32,49]. Gemcitabine can promote cell apoptosis [31]. Then we selected gemcitabine and carboplatin (G/C) as the chemotherapeutic drugs for tumor cell therapy. The cells in the chemotherapy group were incubated with G/C for 1 h [32,49]. And the changes in fluorescence signal were not obvious. Studies had shown that hypoxia can increase tumor resistance and increasing oxygen concentration can improve tumor treatment efficacy [33]. Thus, we maintained tumor cells at HBO condition for 6 h and then the cells were incubated with G/C for 1 h. We can capture achy fluorescence signals in the cells, which indicated that the intracellular NTR concentration decreased with O$_2$ concentration increased. The results demonstrated that we can evaluate the treatment effect through NTR detection during HBO-assistant chemotherapy. We further performed fluorescence imaging in A549 cells using two-photon confocal microscope. And the results from two-photon confocal microscope were consistent well with that from one-photon confocal microscope. Corresponding quantitative analysis was shown in Fig. 3g. We also performed flow cytometry analysis to confirm the fluorescence imaging results (Fig. 3b). Corresponding quantitative analysis was shown in Fig. 3h, and the results from laser confocal microscope imaging and flow cytometry analysis were consistent well. These results demonstrated that TP-NO$_2$ can be used for oxygen assessment during HBO-assistant chemotherapy and further evaluating the treatment efficacy.

Next, we evaluated the therapy efficacy of HBO-assistant chemotherapy through traditional cell damage detection. Calcium ion (Ca$^{2+}$) is one of the most important second messengers. Cell damage will cause Ca$^{2+}$ stores in the mitochondria flow out into the cytoplasm and accumulate [34,35]. We initially assessed cell damage using Fluo 4-AM (Fig. 3c). Ca$^{2+}$ concentration was highest in cells that treated with HBO-assistant chemotherapy, which demonstrated that the cell damage was most severe in this group. The fluorescence quantitative analyses of Ca$^{2+}$ concentrations were shown in Figure 5f. Mitochondrial Ca$^{2+}$-abnormal efflux can cause ΔΨ$_{m}$ collapse and directly lead to cell apoptosis. Then the ΔΨ$_{m}$ was detected by JC-1 to confirm the cell damage (Fig. 5d). And the results were consistent with the results from Ca$^{2+}$ detection. We further observed the cell morphology by transmission electron microscopy (TEM) to evaluate the cell damage. As shown in Fig. 3e, we observed continuous membrane structures and regular organelle structures in the control group. However, the organelle structures had different degrees of abnormalities in different treatment groups. We can observe the membrane structures gradually turned to vague, mitochondrial were swollen, mitochondrial ridge became blurred and the number of vacuoles increased in the experimental groups. It was worth noting that the cell damage was the most severe in the HBO-assistant chemotherapy group, indicating that the HBO-assistant chemotherapy group had the best therapeutic effect. We further performed western blot analysis to explore the signal transduction pathway in the process of tumor cell treatment for evaluating cell damage (Fig. 3f). HIF-1α is a high expression factor in tumor cells, which has been considered as an important target for tumor treatment [36,37]. We analyzed the expression levels of HIF-1α in A549 cells. We found that the expression of HIF-1α in control group was low, while its expression in hypoxia treatment group was the highest. Heat shock protein 70 (Hsp70) is a stress protein which can protect cells from adverse injure. The expression of Hsp70 was increased with the cell damage worsened, and the expression of Hsp70 under hypoxia was much higher than that under normoxia. Cytochrome c, a positively charged protein which acts as a hydrogen carrier in the electronic transmission chain, is an apoptotic mediator capable of activating apoptotic executor cleaved-caspase 3. Once the mitochondrial membrane potential collapse, it can be released from the mitochondria into cytoplasm [38]. As shown in Fig. 3f, the expression of cytochrome c was increased with the cell damage increase. Cytochrome c was also an effector for downstream signal path: Bax, Bcl-2 and ploy(ADP-ribose) polymerase (PARP) and caspase 3 [39,40]. The proteins Bax, Bcl-2, PARP and caspase 3 were detected, and expression of cleaved-caspase 3 and cleaved-PARP as well as the ratio of Bax and Bcl-2 were increased with the increased cell damage. Corresponding protein quantification analysis was shown in Fig. 3i. The results from traditional cell damage detection were consistent with that from efficacy assessment of TP-NO$_2$. Thus we can pre-evaluate the therapeutic effect of HBO-assistant chemotherapy by NTR detection using TP-NO$_2$.

Here, we emphasized that HBO-assistant chemotherapy can indeed help to kill tumor cells and improve the treatment effect. The concentration of NTR was decreased in the group that treated with HBO-assistant chemotherapy. The expression of protective protein Hsp70 was increased to protect cells from damage. The large amount of Ca$^{2+}$ would outflow from the damaged mitochondria to the cytoplasm, and led to the ΔΨ$_{m}$ collapse. Then the cytochrome c was released to cytoplasm from damaged mitochondria, which would further up-regulate the ratio of Bax/Bcl-2 and activate the apoptotic executor cleaved-caspase 3 as well as led to the PARP in nucleus cleaved. And the protein signal transduction mechanism during the process of tumor treatment was shown in Fig. 3j. The cell damages in the group that treated with HBO-assistant chemotherapy were much severer than that treated only HBO or chemotherapy. Thus we supposed that the combination therapy of HBO-assistant chemotherapy might be a powerful strategy for tumor treatment.

3.5. NTR detection in A549 and A549/DDP cell line during tumor treatment

cis-Dichlorodiamineplatinum(II) (DDP) is a commonly adopted chemotherapy drug and studies have shown that long-term drug treatment leads to DDP-resistance [41,42]. Based on the results that HBO can improve the treatment effect of tumor cells, we conducted additional experiments to explore whether HBO-assistant chemotherapy can effectively treat tumor cells. We used the A549 cell line and DDP-resistant A549 cell line (A549/DDP) as the cell models. The A549/DDP cells were obtained as described in Supporting Information. Then the cell model, A549/DDP cell line, was confirmed using MTT experiments. As shown in Figure 5f, A549/DDP cell line had a higher IC$_{50}$ value (20.04 µg/mL) than A549 cell line (1.58 µg/mL). We then performed NTR detection during the process of tumor treatment with G/C and HBO. As shown in Fig. 4a, we observed visible fluorescence signals in the control group whether in A549 or A549/DDP cells. Then, the cells were treated with chemotherapeutic drugs G/C and HBO, and faint fluorescence signals were captured. According to these results, we preliminarily deduced that HBO-assistant chemotherapy can effectively treat A549/DDP cells, and the therapeutic effect of A549 cell line was better than A549/DDP cell line. Corresponding fluorescence quantitative analysis was shown in Fig. 4f. The results from confocal imaging were further confirmed by flow cytometry analysis, and the results were shown in Fig. 4b. Corresponding fluorescence quantitative analysis was shown in Fig. 4g. The results from confocal imaging and flow cytometry analysis were consistent well.

Then the cell damages of A549 and A549/DDP cells in different groups were detected. First of all, the degrees of cell damage were detected using Annexin V/7-AAD Apoptosis Detection Kit (Fig. 4e). The cell damage was most serious in the group that treated with HBO-assistant chemotherapy. Thus HBO-assistant chemotherapy can effectively treat tumor cells. Comparing A549 cell line and A549/DDP cell line, the cell damage was much severer in A549 cell line than A549/DDP cell line. Then the ΔΨ$_{m}$ was further detected using JC-1 (Fig. 4d). The ΔΨ$_{m}$ was lowest in the group that treated with HBO-associated chemotherapy. In addition, the fluctuations of the Ca$^{2+}$ were also detected by Fluo-4 AM using flow cytometry (Fig. 4e). The Ca$^{2+}$ concentration was highest in the cells that treated with HBO-assistant chemotherapy whether in A549 or A549/DDP cells. The fluorescence quantitative analysis of Ca$^{2+}$ was shown in Fig. 4h. The experimental results from cell damage detection were consistent well, which confirmed the hypothesis that HBO-assistant chemotherapy can effectively improve the treatment effect.
treat A549 and A549/DDP cells. Furthermore, during the process of tumor treatment with HBO-assistant chemotherapy, the concentration of NTR was decreased and can reflect the therapeutic effect. These applications made our probe a promising tool for clinical surgical pre-assessment of tumor treatment.

3.6. Therapy efficacy of HBO-assistant chemotherapy in A549 xenografts mice

Encouraged by the results that HBO-assistant chemotherapy can effectively induce tumor cell apoptosis and can effectively treat tumor cells, we then evaluated the therapy efficacy of HBO-assistant chemotherapy in A549 xenografts mice. A549 xenografts were established in nude mice until tumor volume typically reached approximately to 200 mm³ [43]. Next, tumor-bearing nude mice were divided into two groups. The mice in HBO-assistant chemotherapy group were exposed to 80% oxygen every day (five times per week) for 60 min and given G/C biweekly for 28 days (Fig. 5 details seen SI) [32,44,49,50]. Then the NTR concentrations were detected using TP-NO2, and the fluorescence signal acquisition was continued for 30 min. TP-NO2 (1 μM, 50 μL in 1:99 DMSO/saline v/v) was intratumorally injected for NTR detection. As shown in Fig. 5a, bright fluorescence signals were observed in the control group, which indicated that NTR concentration in the control group was high. The concentration of NTR in the group that treated with HBO-assistant chemotherapy was decreased dramatically. Then the organs (including tumor, heart, liver, spleen, lung and kidney) were obtained for fluorescence imaging. As shown in Fig. 5d, we only observed fluorescent signals in tumor tissues. The corresponding fluorescence quantitative analysis of Fig. 5a and d were shown in Fig. 5e and f. The fluorescent signals in the tumor area can last for at least 30 min, and there was no fluorescent signal in the non-tumor area. The fluorescence changes in the above three operations were also offered as movies (Video 1–2 in SI). Thus TP-NO2 can be better used for intraoperative imaging to guide tumor resection after attaching the targeting group. And we can initially evaluate the efficacy of tumor treatment by NTR imaging.

Then we evaluated the therapeutic effect of HBO-assistant chemotherapy through hematoxylin and eosin (H&E) staining and terminal deoxynucleotidyl transferase-mediated deoxyuridine triphosphate nick end labeling (TUNEL) staining. As shown in Fig. 5b, we directly observed cell morphology using H&E. We observed that most cells in the control group had irregular nuclei, which showed the structure of severe malignancy. From the morphological observation, we also observed damaged malignant tumor in the treatment group. TUNEL staining was an indicator of cellular apoptosis, which can point out the cleaved DNA in the nucleus. [45] As shown in Fig. 5c, we observed a small number of cells with nuclei stained in the control group. However, the number of cells with nuclei stained was increased in the HBO-
Tumor hypoxia, the main feature of solid tumors, will lead tumor cells to resist chemotherapy drugs by regulating the expression of multiple genes and directly reduce the therapeutic effect of tumors, which further causes a huge challenge for tumor treatment. The HBO-assistant chemotherapy may be a promising strategy for relieving tumor hypoxia and improving the efficacy of G/C. Therefore, we can assess the tumor treatment effect during HBO-assistant chemotherapy through NTR detection.

3.7. Therapy efficacy of HBO-assistant chemotherapy in A549/DDP xenografts mice

Tumor hypoxia, the main feature of solid tumors, will lead tumor cells to resist chemotherapy drugs by regulating the expression of multiple genes and directly reduce the therapeutic effect of tumors, which further causes a huge challenge for tumor treatment. The HBO-assistant chemotherapy may be a promising strategy for relieving tumor resistance. We further evaluated the treatment effect of HBO-assistant chemotherapy on the mice models that bearing A549/DDP xenograft by NTR detection (Fig. 6). The mice in Fig. 6a were divided into two groups: control group and HBO-assistant chemotherapy group. As shown in Fig. 6a, d, e and f, we can obtain fluorescence signals only in the tumors. The results of fluorescence imaging indicated that HBO can decrease NTR concentration, and the fluorescence intensity was lowest in the group that treated with HBO-assistant chemotherapy. The fluorescence changes in the above two operations were also offered as movies (Video 3 - 4 in SI). These results suggested that HBO-assistant chemotherapy can effectively treat tumor, which might be a key point to alleviating tumor resistance. And we can assess the effect of tumor treatment using NTR detection.

We also performed H&E and TUNEL staining for assessing tumor treatment efficacy by observing tissue damage directly. As shown in Fig. 6b, there was lots of irregular and abnormal nucleus in the control group, which represented severe malignant tumor tissue. We observed damaged tissue in the HBO-assistant chemotherapy group, which suggested good therapeutic effects of HBO-assistant chemotherapy on A549/DDP bearing nude mice. The TUNEL staining in Fig. 6c illustrated that DNA was cleaved during the process of tumor treatment. Thus HBO-assistant chemotherapy can effectively treat tumor. We can more intuitively observe the effect of tumor treatment by directly detecting tumor volume and tumor weight (Fig. 5g and h). The mice weights were also recorded and the results were shown in Figure S11. These results demonstrated that the treatment effect in the HBO-assistant chemotherapy group was satisfying, and synergistic therapies can effectively inhibit the growth of tumor lesions. Thus HBO-assistant chemotherapy is a promising strategy for tumor treatment.

4. Conclusion

We have designed and synthesized a NTR-activatable fluorescent probe, TP-NO2, for NTR detection. TP-NO2 with simple synthesis steps and high yield can quantitatively detect NTR in a wide range (0–20 μg/mL) with a detection limit of 26 ng/mL. The fluorescent probe, TP-NO2, had long emission wavelength and deep tissue imaging depth. The probe was applied for NTR detection in A549 and A549/DDP cells successfully. Furthermore, HBO can greatly increase intracellular oxygen concentration and decrease the concentration of NTR. Then we applied TP-NO2 for NTR detection in A549 and A549/DDP xenografts mice. The results demonstrated that HBO-assistant chemotherapy can effectively treat tumors. HBO-assisted chemotherapy can significantly reduce the NTR concentration, and TP-NO2 can be used as a promising tool for clinical surgery pre-evaluation during HBO-assistant chemotherapy. We believe that the next probe can be better for intraoperative imaging after attaching tumor-targeting group in the future work. And the strategy we proposed provides a novel perspective in accurate diagnosis and efficacy evaluation of tumor treatment.
Declaration of Competing Interest

The authors declare no competing financial interest.

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Appendix A. Supplementary data

Supplementary material related to this article can be found in the online version, at doi:https://doi.org/10.1016/j.snb.2020.127755.

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