X-Ray Induced Photodynamic Therapy: A Combination of Radiotherapy and Photodynamic Therapy

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Abstract

Conventional photodynamic therapy (PDT)’s clinical application is limited by depth of penetration by light. To address the issue, we have recently developed X-ray induced photodynamic therapy (X-PDT) which utilizes X-ray as an energy source to activate a PDT process. In addition to breaking the shallow tissue penetration dogma, our studies found more efficient tumor cell killing with X-PDT than with radiotherapy (RT) alone. The mechanisms behind the cytotoxicity, however, have not been elucidated. In the present study, we investigate the mechanisms of action of X-PDT on cancer cells. Our results demonstrate that X-PDT is more than just a PDT derivative but is essentially a PDT and RT combination. The two modalities target different cellular components (cell membrane and DNA, respectively), leading to enhanced therapy effects. As a result, X-PDT not only reduces short-term viability of cancer cells but also their clonogenecity in the long-run. From this perspective, X-PDT can also be viewed as a unique radiosensitizing method, and as such it affords clear advantages over RT in tumor therapy, especially for radioresistant cells. This is demonstrated not only in vitro but also in vivo with H1299 tumors that were either subcutaneously inoculated or implanted into the lung of mice. These findings and advances are of great importance to the developments of X-PDT as a novel treatment modality against cancer.

Key words: photodynamic therapy; radiotherapy; lung cancer; clonogenecity; nanoparticles.

Introduction

PDT is a relatively new and minimally invasive cancer therapy approach [1-3]. PDT utilizes photosensitizers that are activated by light in the presence of oxygen, producing reactive oxygen species (ROS) that are cytotoxic [3]. PDT can kill cancer cells directly and or damage tumor microvessels, leading to tissue ischemia [2, 4, 5]. It has been utilized in the clinic for treatments of different cancer types, including esophageal cancer, non-small cell lung cancer, bladder cancer, and head and neck cancer [6-10]. Despite of the promises, however, PDT suffers from the shallow tissue penetration of light, especially in the visible spectrum window [11]. There has been progress of developing near-infrared (NIR) photosensitizers, for instance Lumin, Motexafin lutetium, and TOOKAD (absorption peaks at 770 nm, 732 nm and 753 nm) [12, 13]. However, even in the NIR region, light can travel less than 1 cm in tissues. This restriction has largely limited the applications of PDT in the clinic.
Recently, we and others have developed a new PDT derivative called X-ray induced PDT, or X-PDT [14-17]. The goal is to break the shallow penetration restriction by using X-ray as an energy source [18]. The idea of utilizing X-ray to overcome the shallow penetration of PDT was first raised by Chen et al. in 2006 [19], but it was not until very recently that we and the Chen group demonstrated its feasibility in vivo [14-17]. In particular, we showed in a recent study that SrAl$_2$O$_4$:Eu$^{2+}$ nanoparticle (SAO:Eu, a scintillator which convert X-ray photons to visible photons) and MC540 (a photosensitizer with matching excitation wavelength) co-loaded mesoporous silica nanoparticles (MC540-SA0:Eu@mSiO$_2$) can produce singlet oxygen ($^{1}$O$_2$) under X-ray radiation, leading to efficient cancer cell death, even when the cells are beneath thick tissues [14].

It is clear that during X-PDT, not all X-ray energy, but a portion of it, is converted to visible photons to activate PDT. Despite of the studies by us and others observed enhanced treatment efficacy with X-PDT relative to radiation therapy (RT) alone at the same radiation doses [14-17], this phenomenon is intriguing, indicating that there is more to X-PDT than a mere PDT process. In the current study, we tap into the mechanisms behind X-PDT-induced cell death. Specifically, we conducted comprehensive studies to examine the impacts of X-PDT on cell viability, clonogenicity, apoptosis, necrosis, DNA damage, and membrane lipid damage. Our studies showed that X-PDT contains a RT component, and hence it is essentially a RT and PDT combination. The two modalities interplay to attack both cell membrane and DNA, leading to lethal damage that is beyond the repairs of cells. The synergy explains the better cell killing efficacy of X-PDT than RT, even for cells that are refractory to radiotherapy. In particular, we found that X-PDT can efficiently kill H1299 cells, which are radioresistant non-small cell lung cancer (NSCLC) cells. The enhanced efficacy was observed not only in vitro but also in vivo with a subcutaneous tumor model or when H1299 cells were percutaneously implanted into the lung. These findings are of great value to our understanding of X-PDT as a novel treatment modality and its further transformation for eventual clinical translation.

**Results**

**Preparation of MC540-SA0:Eu@mSiO$_2$ nanoparticles**

The MC540-SA0:Eu@mSiO$_2$ nanoparticles were prepared by following our published protocol [14]. Briefly, SrCO$_3$, Al$_2$O$_3$, Eu$_2$O$_3$ and graphite powders were mixed and heated in a tube furnace at 1450 °C for 2 h under an argon flow. The pressure was maintained at 5 Torr. The as-synthesized bulk SAO:Eu was subject to mechanical grinding, followed by sedimentation, filtration and centrifugation, to yield nanoscale nanoparticles (73.5 ± 26.9 nm, Figure 1a&b).

![Figure 1](http://www.thno.org)
X-ray diffraction (XRD) analysis found that the composition of the material was monoclinic SrAl₂O₄ (JCPDS #74-0794, Figure S1). Inductively coupled plasma (ICP) confirmed the chemical composition and found that Eu accounts for ~1% of the total mass. As described in the previous study, SrAl₂O₄:Eu can be stimulated by both Uv-vis and X-ray to emit green photoluminescence (centered at ~520 nm, Figure 1c), which is attributed to 4f⁶5d¹→4f⁷ transition of Eu²⁺ ions [15].

SrAl₂O₄:Eu is a highly hydrolytic material and is quickly reduced to constituent ions in an aqueous solution [15]. For bio-applications, we coated SrAl₂O₄:Eu nanoparticles with two layers of silica (Figure 1b). These include an inner, solid silica layer that prevents direct contact with the aqueous surroundings, and an outer, mesoporous silica layer that provides a docking place for photosensitizers [14]. Into the resulting SrAl₂O₄:Eu@mSiO₂ nanoparticles, MC540 (Figure S2), a common photosensitizer that has been investigated in both pre-clinical and clinical studies [20], was loaded. The absorbance wavelengths of MC540 well match the emission wavelengths of SAO:Eu (Figure 1c, Figure S3). Using singlet oxygen sensor green (SOSG) assays, we studied the ability of X-PDT to producing ¹O₂. It was found that X-ray alone (1-4 Gy) was not able to produce ¹O₂, either with PBS, MC540, or SrAl₂O₄:Eu nanoparticles (Figure 1d). As a comparison, when both SrAl₂O₄:Eu@mSiO₂ (0.05 mg/mL) and X-ray were applied, efficient ¹O₂ production was observed (Figure 1d).

**Impact of X-PDT on cell viability and clonogenecity**

We first investigated the impact of X-PDT on cell viability by MTT assays. Radioresistant NSCLC H1299 cells were used in the studies [21-23]. In the absence of radiation, SAO:Eu and SAO:Eu@mSiO₂ nanoparticles were not toxic to H1299 cells (Figures 1e, 2a). The low toxicity of SAO:Eu@mSiO₂ was also

![Figure 2](http://www.thno.org)
observed in our previous studies with other cell lines [24] and is not surprising because all the constituents of the nanoparticles, including MC540, SAO:Eu, and SiO$_2$, are not toxic in the dark [14]. Meanwhile, RT alone (0-5 Gy) did not induce significant cell death at 24 hours either (Figure 2a). As a comparison, when X-ray irradiation was applied after MC540-SA0:Eu@mSiO$_2$ (50 µg/mL) incubation (i.e. X-PDT), there was significant cell viability drop (Figure 2a). Specifically, when irradiation of 0.83, 1.67, 3.33, and 5 Gy was applied, the 24-hour cell viability was reduced to 31.4 ± 2.3%, 19.6 ± 4.6%, 18.6 ± 7.4%, and 17.5 ± 5.6%, respectively.

While MTT assay is adequate to measures short-term cell viability, it is suboptimal in assessing the reproductive capacity of cancer cells. For RT, however, reproductive capacity is a more relevant ending point. This is because ionizing radiation mainly targets DNA, most importantly double-strand DNA break [25]. This is often manifested not as immediate cell death but reduced clonogenicity in days or weeks [26]. To investigate, we performed clonogenic assays with H1299 cells that received RT alone (0, 1, 2.5, 5, 7.5 and 10 Gy) or RT plus MC540-SA0:Eu@mSiO$_2$ (50 µg/mL). X-PDT was much more effective than RT alone at all doses (Figure 2b). Taking 5 Gy for instance, X-PDT was able to reduce the survival fraction (SF) to 17.5%, compared to that of 42.5% for RT alone. The dose enhancement factor (DEF) was calculated to be 1.67 (Methods Section).

Impact of X-PDT on DNA and lipid membranes

To further investigate the cause of X-PDT induced cell death, we performed Apoptotic/Necrotic/Healthy assay (PromoCell, Heidelberg, Germany), again with H1299 cells. With RT only (5 Gy), cells showed a mediocre level of apoptosis at 24 h but no detectable necrosis (Figure 2c). On the contrary, when cells were treated with MC540-SA0:Eu@mSiO$_2$ (50 µg/mL) plus radiation, there manifests extensive cell necrosis (Figure 2c). This pattern resembles membrane-targeted PDT, which causes oxidative degradation of unsaturated lipids and surface proteins [27]. To confirm, we conducted lipid peroxidation assay (Lipid Peroxidation Kit, Life Technologies, Carlsbad, CA). Compared to the control (PBS only), the lipid peroxidation level was increased by 90.5 ± 23.0% after RT (5 Gy); meanwhile, X-PDT under the same irradiation dose led to an increase of 201.5 ± 34.0% (Figure 3a). Such lipid peroxidation is attributed mainly to the PDT component of X-PDT and is responsible for the short-term cell necrosis and viability drop.

Figure 3. Impacts of X-PDT on cellular compartments. (a) Lipid damage assessment, measured by lipid peroxidation assays. (b) DNA damage, assessed by single cell electrophoresis assays. (c) Western blot assays, which further confirms the impact of X-PDT on DNA and membrane lipids.
Meanwhile, X-PDT also causes DNA damage. This was confirmed by comet assays (also known as single cell gel electrophoresis assay, Figure 3b). Compared to the control, cells received RT alone (5 Gy) displayed a relatively long and intense tail, suggesting extensive DNA double-strand break [28]. Such a tail was also observed with X-PDT treated cells, but was shorter, less intense, and more discrete, likely associated with the extensive necrosis [28]. This suggests that there remains a RT component in X-PDT, which explains the reduced cell survival in clonogenic assays.

To further assess the cellular impacts of X-PDT, we analyzed expression levels of histone H2AX and Cox-2 in X-PDT treated H1299 cells with Western blot (Figure 3c). Histone H2AX plays a critical role in recruiting repair- or damage-signaling factors to the sites of DNA damage [29, 30] and is thus an indicator of RT-induced DNA breakage. Cox-2, on the other hand, is involved in lipid peroxidation, and is often up-regulated after PDT-induced membrane damage [31, 32]. We found that X-ray radiation (5 Gy) alone induced expression of H2AX but minimally affected the level of Cox-2 (Figure 3c). With X-PDT, on the other hand, both Cox-2 and H2AX expressions were increased, although the H2AX level was lower than that after RT (Figure 3c). These results corroborate with the preceding studies, again confirming that X-PDT is essentially a combination of RT and PDT. It is postulated that because the two modalities target different cellular compartments (DNA and unsaturated membrane lipids, respectively), a synergy in treatment occurs that leads to much more efficient cancer cell killing.

**Impact of tissue depth on the efficacy of X-PDT**

We next studied the impact of tissue thickness on the treatment efficacy of X-PDT in vivo. This was first studied with mouse subcutaneous tumor models established with H1299 cells. Briefly, we intratumorally injected MC540-SA0:Eu@mSiO2 nanoparticles (4.25 mg/kg) to tumor bearing animals; we then irradiated the tumors (5 Gy, with the rest of the body lead-shielded), with pork tissues of 1- or 2-cm thickness lain on top. In control group, animals were injected with PBS and received no radiation.

Tumors in the control group grew very rapidly, and either died or reached an end point (tumor diameter > 1.7 cm) within 10 days (Figure 4).
As a comparison, in all the treatment groups, tumor growth was efficiently slowed. On Day 16, the tumor suppression rates were 54.2% and 33.8%, for the animals bore with 1- and 2-cm thick pork, respectively. The results suggest that tissue thickness can still affect the efficacy of X-PDT; the impact, however, was much less severe than with PDT, which lost its efficacy beyond 1 cm thickness. Such tissue impact can likely be compensated by increasing radiation doses and is less of a concern in the clinical setting, where deep-penetrating megavoltage X-rays are used [33]. The efficiency in cancer cell killing was further confirmed with H&E analysis (Figure 4b). Compared with the control animals, where cancer cells were densely packed, X-PDT dramatically reduced cell density and disrupted connective tissues (Figure 4b). Meanwhile, we observed no signs of systematic toxicities to the surrounding tissues (Figure 4c).

Next, we investigated the X-PDT efficacy by injecting nanoparticles and cell mixture to the thorax of mice and irradiated the injection sites in vivo. Due to thick soft tissues and bones, diseases at this position are not accessible by conventional PDT [34]. Specifically, we injected MC540-SA0:Eu@mSiO2 nanoparticles (4.25 mg/kg), along with $5 \times 10^5$ firefly luciferase expressing H1299 cells (H1299-Luc), into the left lateral thorax of nude mice (Figure 5a). Radiation (5 Gy) was applied to the tumor inoculation sites, with the rest of the animal body shielded by lead. Control group animals received radiation only or PBS only. The tumor growth was then monitored in vivo by bioluminescence imaging (BLI). In PBS and RT only groups, the BLI signals were detected in the lung areas on Day 7 and continued increasing afterwards (Figure 5a). In X-PDT treated animals, on the other hand, the BLI signals were significantly suppressed. By regions of interest (ROI) analysis, on Day 12, the average BLI signals were $5.6 \times 10^5$, $2.9 \times 10^6$, and $2.6 \times 10^6$ photons/sec/cm²/sr for the X-PDT, RT, and PBS groups, respectively (Figure 5b). Ex vivo imaging confirmed the efficiency of X-PDT induced tumor suppression, finding strong residual signals in the lungs of control animals but close-to-background signals.
signals from the X-PDT group (Figure 5c,d). Moreover, we found multiple large tumors in the lungs dissected from control animals, but few detectable colonies in X-PDT treated mice (Figure 5e). Such reduced tumorigenicy by X-PDT was further confirmed by H&E staining (Figure 5f).

Meanwhile, X-PDT did not cause detectable systematic toxicities. These include no signs of side effects to normal tissues from H&E staining (Figure 6a). Also, hemodiagnosis found that the serum aspartate aminotransferase (AST) and alanine aminotransferase (ALT) levels remained unchanged over the course of the treatment, again suggesting low toxicities (Figure 6b). This is in accordance with our previous observations that i.v. injected MC540-SAO:Eu@mSiO₂ nanoparticles are degraded after the conclusion of therapy and efficiently cleared from the hosts [14].

Discussions

While initially developed as a PDT derivative, the current study suggests that X-PDT is essentially a PDT and RT combination. While a portion of X-ray energy was converted to activate PDT, ionizing radiation continues playing an important role in X-PDT. The PDT and RT components target different cellular components and the combination causes enhanced effect that is beyond the repairs of cells. This explains the greater cytotoxicity with X-PDT than with RT, especially for cells that are refractory to RT. In fact, previous studies have observed synergy between PDT and RT [35-38]. However, it was found that PDT and ionizing irradiation need to be given at the same time so as to override cell repairs [35, 36]. This requires photo-irradiation and X-ray irradiation to be applied simultaneously, which is difficult to achieve in the clinic, not to mention the shallow penetration of PDT. X-PDT, on the other hand, is an inherent PDT and RT combination, with both processes regulated by one deep-penetrating radiation. From this perspective, X-PDT represents a major advance for seamlessly integrating PDT and RT. In this sense, X-PDT should be viewed as not only a PDT derivative but also a RT derivative.

RT remains a major therapy option in clinical oncology. More than 50% of all cancer patients receive RT during their curative process [39]. One major limitation of RT is that not all cancer cells are well responsive to RT, and tumors that are originally responsive may develop resistance over the course of therapy [40]. Increasing radiation doses can, to a certain degree, address the issue, but will inevitably cause collateral damage to normal tissues [41]. Radiosensitizing agents of different types have been developed to sensitize cancer cells to RT; however, many radiosensitizers are cytotoxic agents [42]. As discussed above, X-PDT can be viewed as a RT derivative and as such a novel radiosensitizing technology. This was confirmed by our clonogenic assays, which observed a high DEF with X-PDT, even against cells that are refractory to RT. Unlike many conventional radiosensitizers, MC540-SAO:Eu@mSiO₂ nanoparticles are not toxic in the absence of radiation. Meanwhile, they are highly biodegradable, efficiently cleared out of the hosts after treatment, and causing no side effects to normal tissues (Figure 6). These properties suggest great promise of X-PDT in clinic translation to improve RT efficacy, reduce normal tissue radiation exposure, and battle with radioresistant tumors. In the present studies, nanoparticles were directly injected into tumors. In the future studies, it is worthwhile to reduce nanoparticle size, improve photosensitizer loading, and conjugating targeting ligands to the nanoparticle surface for achieve formulations that can be systematically injected to mediate tumor selective X-PDT treatment.

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**Figure 6.** Systematic toxicities of X-PDT. (a) H&E staining with normal tissues taken from X-PDT, RT, MC540-SAO:Eu@mSiO₂, and control groups. Scale bars, 100 μm. (b) Hemodiagnosis. ALT and AST level showed no significant changes before and 14 days after X-PDT treatment.
Conclusion

In summary, we investigated in this study the impacts of X-PDT on cancer cells. We found that X-PDT is not simply a PDT process; rather, it is essentially a RT and PDT combination. The combination leads to enhanced therapy effects that make the treatment much more efficient than RT alone, even when used against cells that are refractory to RT. We also show that X-PDT can be exploited to suppress tumors lain under deep tissues. The findings and advances are of great value to the developments of X-PDT as a novel treatment modality against cancer.

Materials and Methods

Nanoparticle synthesis and coating SAO:Eu was synthesized by a carbo-thermal reduction and vapor-phase deposition method that was published previously [43]. To render SAO:Eu amenable to bio-related applications, bulk SAO:Eu was ground into nanoparticles with a diameter of ~ 80 nm. The bare SAO:Eu nanoparticles were coated with a layer of solid silica by following a previously published protocol [14]. In a typical synthesis, 10 mg of SAO:Eu nanoparticles were dispersed in a mixture of 25 mL of ethanol, 2 mL of H2O, and 1 mL of ammonia (25%), and stirred for 10 min. Then, 75 μL of tetraethyl orthosilicate (TEOS) was added to solution and the mixture was stirred at room temperature for 12 hours. The resulting nanoparticles were subsequently coated with a layer of mesoporous silica. A mixture of 3-aminopropyltriethoxysilane and TEOS (5 v/v%) was used as silane precursors [24]. Solid silica coated SAO:Eu nanoparticles (10 mg) from the first step were dispersed in a mixture of 45 mL of H2O, 0.3 mL of 2 M NaOH, and 10 mg of cetyltrimethylammonium bromide (CTAB) for 30 min. After heating to 70 °C, 50 μL of TEOS and 300 μL of ethyl acetate were added, and the solution was magnetically stirred for 2 h. The raw products were collected by centrifugation and washed with ethanol for three times. The resulting nanoparticles were resuspended in PBS (Thermo Scientific) for further studies.

Characterizations of SAO:Eu and MC540-SA0: Eu@mSiO2 nanoparticles. UV-Vis absorption spectra were recorded on a Shimidzu 2450 UV-Vis spectrometer. Photoluminescence measurements were performed on a Hitachi F-7000 fluorometer. A mini-X X-ray tube (Amptek Inc.) was used as the X-ray source, and was set at 50 kV and 70 μl for all the experiments in this study. TEM samples were prepared by dripping sample solutions onto carbon-coated copper grids and evaporating the solvent. TEM images were taken on an FEI Tecnai 20 transmission electron microscope operating at 200 kV.

Cell culture H1299, a human NSCLC cell line, was used in in vitro and in vivo studies. The H1299 cells have been transfected to stably express firefly luciferase (i.e. H1299-Luc). The cells had been tested and were rodent pathogens free. H1299-Luc cells were grown in RPMI 1640 medium supplemented with 10% FBS, 100 units/mL of penicillin (ATCC) and 250 μg/mL hygromycin. The cells were maintained in a humidified, 5 % carbon dioxide (CO2) atmosphere at 37 °C.

In vitro toxicity study 10^4 H1299 cells were seeded in 96-well plates (Corning) and cultured to 90% confluency. For cytotoxicity studies, the cells were then incubated with 0, 5, 10, 25, 50 and 100 μg/mL MC540-SA0:Eu@mSiO2 nanoparticles for 24 h. Subsequently, they were washed with PBS for two times, and then were evaluated by MTT assay (Sigma Aldrich) by following the vendor’s protocols.

In vitro viability study. 10^4 H1299 cells were seeded in 96-well plates (Corning) and cultured for 24 h. The cells were then divided into 5 groups: 1) PBS group, 2) MC540-SA0:Eu@mSiO2 nanoparticles only, 3) MC540 only, 4) RT only, and 5) X-PDT. For Group 2 and 5, the final concentration of MC540-SA0:Eu@mSiO2 nanoparticles was 50 μg/mL and the incubation time was 1 hour. All the cells were incubated in the dark. Cells in Group 4 and 5 were irradiated with 50 kV X-ray at 0.83, 1.67, 3.33 and 5 Gy while all the other cells received irradiation. After the treatment, all cells were washed with PBS for two times. Cell culture medium was replenished and the incubation was maintained for 24 h (5 % CO2 and 37 °C). Cell viabilities were evaluated by MTT assays (Sigma Aldrich) following the vendor’s protocol.

Apoptosis/necrosis/healthy assay H1299-Luc cells treated with X-PDT (5 Gy), RT (5 Gy) and PBS were subjected to Apoptotic/Necrotic/Healthy assay (PromoCell GmbH, Heidelberg, Germany) following the vendor’s protocol. Briefly, FITC-annexin V (green) binds to the apoptosis marker of phosphatidyl serine, and ethidium homodimer III (red) binds to DNA under the necrotic conditions. The cells stained with annexin V antibody alone or together with ethidium homodimer III were counted as early or late stages of apoptotic cells, respectively [34]. The cells labeled with ethidium homodimer III alone were counted as
necrotic cells. In addition, membrane permeable Hoechst 33342 (blue) stains the nuclei. Healthy cells would only display blue staining. The resulted cells were then evaluated by fluorescent imaging on an Olympus IX71 fluorescent microscope (3 trials per group).

**Clonogenic assay** Mono-layered H1299-Luc cells were prepared in 6-well cell petri-dishes one day before the experiments. On the experimental day, the 6-well cell petri-dishes were randomly divided into the control group, the X-ray radiation group, and the X-PDT group. The two treatment groups were treated by X-ray (50 kv) for a dose of 1, 2.5, 5, 7.5 and 10 Gy while the control group received no irradiation. The irradiated and un-treated cells were then harvested using trypsin-EDTA and plated into 100 mm cell petri-dishes. Each petri-dish was seeded with 100 viable cells and was placed into a cell culture incubator (5% CO₂ and 37°C) for 14 days. After incubation for 14 days, the cells were fixed by formalin and stained with 0.5% Gentian Violet. Colonies containing more than 50 cells were counted and the survival fractions were calculated by comparing to the control. Dose enhancement factor (DEF) was calculated as the ratio between RT and X-PDT radiation doses at 10% survival fractions. All the experiments were repeated 3 times.

**Lipid peroxidation assay** Lipid peroxidation levels were measured using Image-iT® Lipid Peroxidation Kit (Life Technologies Corporation, Carlsbad, CA) by following the vendor’s protocol. Briefly, H1299-Luc cells seeded in 6-well petri-dishes were treated with RT or X-PDT under 5 Gy X-ray irradiation. Un-irradiated cells were used as control. The Image-iT reagent was incubated with cells for 30 min. The cells were washed three times with PBS and observed on an Olympus IX71 fluorescence microscope. Data were analyzed by ImageJ (NIH, Bethesda, Maryland, U.S.). The experiments were repeated 5 times.

**Single cell gel electrophoresis (comet assay)** Mono-layered cells were prepared in 6-well cell petri-dishes one day before the experiments. On the experimental day, the 6-well cell petri-dishes were randomly divided into the control group, the X-ray radiation group, and the X-PDT (with MC540-SAO:Eu@mSiO₂) group. Then the two treatment groups were irradiated by X-ray (50 kV) for a dose of 5 Gy while the control group received 0 dose irradiation. The irradiated and un-treated cells were then harvested. 1 × 10⁵/mL of the treated cells were immediately combined with molten LMAgarose (at 37°C) at a ratio of 1:10 (v/v). 50 μL of the solution was pipetted onto a CometSlide™ (Trevigen Inc., Gaithersburg, MD). The slide was then immersed in 4°C lysis solution for 1 hour or overnight. After that, ~850 ml 4°C 1× neutral electrophoresis buffer was added to the electrophoresis gel box and the slides were placed in a slide tray. The power supply was set at 21 volts. After 45 minutes, the slides were gently removed and immersed in DNA Precipitation Solution for 30 minutes, and then in 70% ethanol for 30 minutes at room temperature. The slides were dried and stained in SYBR® safe for 30 min in the dark. The slides were then ready for microscopic imaging.

**Western blot** Approximately 5 × 10⁵ H1299-Luc cells were seeded in a 6-well plate prior to radiation therapy. Cells were harvested 1 hours and 24 hours after the treatment. The cells were lysed in a lysis buffer. Proteins were separated by pre-cast 12% Bis-Tris NuPage™ SDS-PAGE (Life Science™), transferred to iBlot® polyvinylidene difluoride membrane (Invitrogen™, Grand Island, NY, USA), and immunoblotted using primary antibodies against COX2, H2AX, and β-actin. HRP-conjugated secondary antibody (Cell Signaling Technology, Danvers, MA, USA) was applied subsequently. The protein bands were then visualized with SuperSignal™ West Pico Chemiluminescent Substrate (Thermo Fisher Scientific) using a Fluorchem HD2 chemiluminescent imaging system (Protein Simple, Santa Clara, CA).

**Animal studies** All the animal studies conformed to the Guide for the Care and Use of Laboratory Animals published by the National Institutes of Health, USA, and a protocol approved by the Institutional Animal Care and Use Committee (IACUC), University of Georgia. Nude mice (4–6 weeks old, Harlan) with a body weight of ~20 g were used for the animal model establishment. The animals were housed in alternating lighting conditions under a 12 h dark and 12 h light regime.

**Therapy studies with subcutaneous tumor models** 15 nude 5–6 week athymic nude mice were randomly divided into three groups, 1) control group, 2) 1-cm thick pork group, and 3) 2-cm thick pork group. Animal models were established by subcutaneous injection of 5 × 10⁵ H1299-Luc tumor cells onto the hind legs of mice. When tumor sizes reached 100 nm, 100 μL PBS solution containing MC540-SAO:Eu@mSiO₂ nanoparticles (4.25 mg/kg) was intratumorally injected in Group 2 and 3. The same amount of PBS was injected into the tumors of the control group. For therapy studies, irradiation (5 Gy) was applied to tumors, with the rest of the animal body covered by lead. The control group received no irradiation.

**Therapy studies with cancer cells implanted into the lungs** 20 nude mice were randomized to

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receive the following treatments group (n = 5): 1) PBS, 2) MC540-SAObEu@mSiO2 only, 3) RT only, and 4) X-PDT (MC540-SAObEu@mSiO2 plus RT). 5 × 10^6 H1299-Luc cells were injected into the left lateral thorax of anesthetized nude mice. For Group 1 and 3, PBS solution was premixed with the cells into a final volume of 50 µL. For group 2 and 4, nanoparticles were premixed with cells into a final volume 50 µL matrigel solutions (2.5 mg MC540-SAObEu@mSiO2/mL, or 1.7 mg SAObEu/mL). Group 1 and 2 received no X-ray irradiation. Group 3 and group 4 received X-ray irradiation (5 Gy) 5 minutes after the injection. Only one therapy dose was given to each animal. The tumor growth was monitored BFI by an IVIS Lumina scanner (PerkinElmer Inc. Waltham, Massachusetts). Tumors and major organs from the euthanized animals were harvested, weighed, and cryosectioned. The tissue sections were then subjected to standard H&E staining to assess treatment outcomes and side effects (BBC Biochemical). Hemodiagnosis assay was performed by withdrawing the blood before the treatment as well as before euthanizing the mice. The concentrations of AST and ALT were determined by using commercial kits from Thermo-Scientific.

Statistical analyses Quantitative data were expressed as mean ± s.e.m. A two-tailed Student’s t-test was used for statistically comparing the treatment group and the control group. P < 0.05 was considered statistically significant.

Supplementary Material

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Competing Interests
The authors have declared that no competing interest exists.

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