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# An Activatable Unimolecular Phototheranostic Agent for Synergistic Chemo-Photodynamic Therapy

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Precise control over spatiotemporal release of cancer therapeutics remains a significant challenge in developing effective combination therapies. Herein, we report a hydrogen peroxide  $(H_2O_2)$ -activatable unimolecular platform (**ICy-Cb**) that integrates cancer imaging and synchronized dual-modal therapy. An iodized hemicyanine photosensitizer and chlorambucil are integrated into a single molecular structure that selectively reacts to the high concentration of  $H_2O_2$  present in the tumor microenvironment. **ICy-Cb** initially exhibits no fluorescence and is therapeutically inert due to the suppression of its intramolecular charge transfer (ICT) effect. Upon exposure to tumor-associated  $H_2O_2$ , **ICy-Cb** simultaneously releases both therapeutic components, demonstrating exceptional tumor selectivity (5 to 8 folds higher activation in cancer cells), deep penetration in 3D tumor models (180 µm), and persistent tumor accumulation in vivo. Most importantly, this synchronized chemo-photodynamic agent achieves superior tumor growth inhibition (92%) compared to either monotherapy. This theranostic agent represents a significant advancement for precision cancer treatment by enabling spatiotemporally controlled combination therapy within the tumor microenvironment.

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#### Introduction

Cancer is a global malignant disease caused by disorderly cellular division and mutational growth.<sup>1,2</sup> Among cancer treatment modalities, chemotherapy remains predominant due to its systemic effectiveness in targeting metastatic cancer cells throughout the body, proven track record against various malignancies (especially blood cancers).<sup>3</sup> However, traditional chemotherapy, which relies solely on anticancer drugs, faces limitations such as low specificity to tumor cells and undesirable side effects, often resulting in suboptimal therapeutic outcomes.<sup>4,5</sup> Recently, photodynamic therapy (PDT), which exhibits high spatiotemporal selectivity, minimal systemic side effects, and noninvasiveness, has emerged as a promising alternative approach in oncological treatment.<sup>6,7</sup> During PDT, the photosensitizer (PS) sensitizes oxygen under light irradiation to produce highly oxidative reactive oxygen species (ROS), which cause tumor cell apoptosis or necrosis.<sup>8-12</sup> Among various photosensitizers, hemicyanine dyes represent ideal candidates for phototheranostic applications due to their excellent properties, including NIR absorption/emission profiles ( $\lambda_{max} \ge$ 680 nm), large molar extinction coefficients, and modifiable photochemical sites that enable deep tissue penetration and effective singlet oxygen generation.<sup>13-17</sup> However, the hypoxic microenvironment within deeper tumor regions greatly

diminishes the generation of ROS, leading to a reduction in the therapeutic effectiveness of PDT.<sup>18</sup> With the continuous evolution of cancer cells, the therapeutic efficacy of singlemodality treatments for malignant tumors has become increasingly limited.<sup>19</sup> As a contrast, combined therapy can capitalize on the complementarity of various treatment methods to achieve synergistic advantages, ultimately optimizing the overall therapeutic outcome.<sup>20-22</sup> For instance, chemotherapy damages cellular structures and modifies the permeability of cell membranes, thereby facilitating the uptake of PS and concurrently enhancing the sensitivity of cancer cells to oxidative stress. Conversely, the ROS generated by PDT can disrupt both the drug efflux pump and the cellular defense mechanisms, thus increasing drug retention and mitigating drug resistance of cancer cells.<sup>23-25</sup> The combination of PDT and chemotherapy creates a synergistic effect, where chemotherapy intensifies the effectiveness of PDT, and PDT, in turn, boosts the results of chemotherapy.

However, conventional combined therapy often employs a "cocktail" approach, which leads to significant challenges including inconsistent biodistribution. heterogeneous metabolism profiles, asynchronous activation at target sites, and unpredictable pharmacokinetic interactions.26-28 Additionally, most therapeutic agents are in an "always-on" state, resulting in a series of side effects such as high toxicity to normal cells, weak tumor specificity, and high phototoxicity, ultimately reducing their therapeutic efficacy against tumors.<sup>13,29-31</sup> In contrast, therapeutic agents capable of being activated by a specific biological target of interest display "offon" characteristics, thereby significantly reduce side effects while improving both the biosafety and therapeutic efficacy.<sup>32-</sup> <sup>35</sup> Hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>), as a critical reactive oxygen

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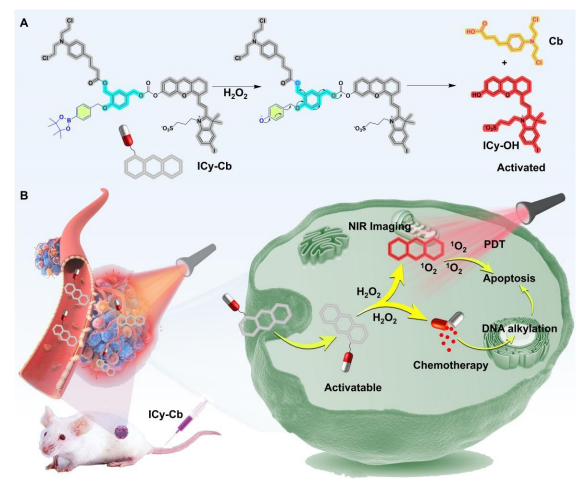
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species substantially overexpressed in various tumor types with significantly higher concentrations in tumor cells (5  $\mu$ M to 1.0 mM) compared to normal cells (less than 0.7  $\mu$ M), provides an ideal biomarker for developing tumor-specific theranostic agents.<sup>36-39</sup> Ideal H<sub>2</sub>O<sub>2</sub>-responsive systems should possess three key attributes: specific activation by tumor microenvironment H<sub>2</sub>O<sub>2</sub>; simultaneous release of multiple therapeutic components for synergistic effects; and integration of both imaging and therapeutic capabilities for real-time monitoring of drug distribution and efficacy. However, most current H<sub>2</sub>O<sub>2</sub>-responsive systems achieve only single treatment modality or require complex nanocarriers, which increase the complexity of preparation and potentially introduce new safety concerns.

In this work, we present a novel  $H_2O_2$ -activatable unimolecular phototheranostic agent, **ICy-Cb**, for cancer visualization and synchronized combination therapy (**Scheme 1**). In the architecture of this designed system, both a photosensitizer and chemotherapeutic agent are strategically incorporated within a single molecular framework that specifically responds to elevated  $H_2O_2$  levels in the tumor microenvironment. **ICy-Cb** is initially non-fluorescent and therapeutically inactive as its intramolecular charge transfer (ICT) is inhibited by the pinacol borate moiety that cages the hemicyanine dye. Only in the presence of tumor-associated

H<sub>2</sub>O<sub>2</sub>, ICy-Cb undergoes site-specific oxidation and releases the iodized hemicyanine photosensitizer (ICy-OH)1and3chPoFamb807cfl (Cb) through self-immolative fragmentation, enabling PDT and chemotherapy respectively. ICy-Cb demonstrates remarkable selectivity, providing high-contrast differentiation between normal cells and multiple cancer cell lines (4T1, MCF-7, HeLa) with 5-8 folds higher activation in cancer cells. Threedimensional tumor models revealed exceptional penetration capabilities (up to 180  $\mu m$  depth), facilitating effective treatment throughout the tumor mass. In vivo imaging confirmed selective accumulation in tumor tissues with persistent fluorescence signals lasting 24 hours, enabling precise tumor visualization. Most significantly, when compared to single-modality approaches (chemotherapy or photodynamic therapy alone), ICy-Cb-mediated synchronized combination therapy demonstrated dramatically enhanced therapeutic efficacy, achieving remarkable tumor growth inhibition (92% in vivo). By integrating dual therapeutic modalities within a single H<sub>2</sub>O<sub>2</sub>-responsive molecular platform, ICy-Cb addresses a fundamental challenge in cancer treatment: achieving precise spatiotemporal control over combination therapy while minimizing systemic toxicity, thereby representing a promising approach for translational applications in cancer theranostic.



Scheme 1. Schematic diagram of (A) molecular structure and H<sub>2</sub>O<sub>2</sub>-activated (B) in vivo anti-tumor mechanism of ICy-Cb.

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#### **Results and discussion**

#### Design and synthesis of ICy-Cb.

The strategic design and synthesis of ICy-Cb were accomplished following the synthetic route depicted in Schemes 1 and S1, with the aim of creating a dual-modality theranostic agent. An iodized hemicyanine dye (ICy-OH) was integrated into the system as an activatable PS for PDT.<sup>13</sup> Chlorambucil (Cb), a clinically validated chemotherapeutic agent widely utilized in cancer treatment, exerts its primary pharmacological action by inducing cross-linking of the DNA double helix structure in cancer cells, thereby disrupting DNA replication and extension processes, ultimately resulting in cancer cell death and tumor suppression.<sup>40-42</sup> Notably, when the carboxyl moiety of butyric acid forms an ester bond with a hydroxyl group, the cytotoxicity of **Cb** is significantly attenuated due to inhibited oxidation at the  $\beta$ -position of the butyrate side chain, making it an ideal candidate for our prodrug system.<sup>22</sup> The rationally designed molecule (ICy-Cb) was engineered to undergo activation specifically by elevated levels of  $H_2O_2$  in the tumor microenvironment, facilitating the simultaneous release of both the PS and chemotherapeutic agent for concurrent PDT and chemotherapy, respectively (Fig. 1A). To further validate this strategy, we synthesized a control molecule, ICy, by conjugating ICy-OH with pinacyl 4-phenylborate benzyl alcohol, which selectively releases the photosensitizer ICy-OH in the presence of  $H_2O_2$  (Scheme S2). All intermediates and final compounds (ICy-Cb and ICy) were thoroughly characterized via <sup>1</sup>H NMR, <sup>13</sup>C NMR, MS, and HRMS analyses (see Supporting Information), confirming their structural integrity and purity essential for subsequent biological evaluations.

#### Sensing property of probe ICy-Cb.

Upon successful synthesis and characterization of the novel compound ICy-Cb, we thoroughly investigated its sensitivity and specificity toward H<sub>2</sub>O<sub>2</sub>. The absorption spectrum of ICy-Cb, illustrated in Fig. 1B, displays a prominent absorption peak centered at 600 nm, while exhibiting negligible fluorescent emission (Fig. 1C), indicating the effective quenching of fluorescence in its native state. Upon exposure to H<sub>2</sub>O<sub>2</sub>, a remarkable spectral transformation occurs: the absorption peak at 600 nm diminishes concurrently with the emergence of a new band at 670 nm (Fig. 1B), signaling the structural conversion of the molecule. Sequential addition of increasing  $H_2O_2$  concentrations to the ICy-Cb solution resulted in progressive enhancement of the emission band at 720 nm (under 640 nm excitation), reaching maximum intensity after 60 minutes (Fig. 1C and 1E). Notably, the fluorescence intensity at 720 nm demonstrated a strong linear correlation with  $H_2O_2$ concentration, yielding a calculated detection limit of 0.3  $\mu$ M/mL (Fig. 1D). We postulate that the significant spectral shift observed can be attributed to the H<sub>2</sub>O<sub>2</sub>-mediated conversion of ICy-Cb to ICy-OH through a specific molecular mechanism. The activation process proceeds through several discrete sequential steps: initially, the pinacol borate moiety undergoes oxidation

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by H<sub>2</sub>O<sub>2</sub> to form a hydroxyl intermediate. Subsequently, electronic rearrangement within this activated species friggers the cleavage of the benzyl C-O bonds via 4- or 6-electron reduction pathways, facilitating the simultaneous release of both Cb and ICy-OH. Concurrently, the cleaved intermediates react with water molecules to spontaneously generate phydroxybenzyl alcohol. This decapping process effectively restores ICT within the hemicyanine structure, thereby inducing pronounced fluorescence enhancement in the near-infrared region, which is highly advantageous for in vivo imaging applications. To validate our proposed mechanism with chemical precision, we conducted comprehensive analysis of the reaction products using Electrospray Ionization Mass Spectrometry (ESI-MS). As illustrated in Fig. S1, the reaction mixture containing probe ICy-Cb and H<sub>2</sub>O<sub>2</sub> revealed two distinctive mass peaks at m/z = 617.0711 and m/z = 304.0811, which correspond precisely to the theoretical molecular masses of the expected products ICy-OH ([M]<sup>+</sup> = 617.0733) and Cb ([M+H]<sup>+</sup> = 304.0866), providing direct evidence for our proposed reaction pathway and product formation.

For more rigorous confirmation, we employed High-Performance Liquid Chromatography (HPLC) analysis, which provided additional quantitative evidence for the simultaneous liberation of ICy-OH and Cb. Following a 2-hour reaction period with H<sub>2</sub>O<sub>2</sub>, the chromatographic peak associated with the parent probe ICy-Cb (retention time 17 min) was virtually eliminated (Fig. 1F). Concurrently, two new distinctive peaks emerged at retention times of 6.5 min and 4.5 min, aligning precisely with the retention times of authentic standards of ICy-OH and Cb, respectively (Fig. 1F). The H<sub>2</sub>O<sub>2</sub>-catalyzed cleavage kinetics of ICy-Cb were quantitatively monitored through timecourse HPLC analysis. This investigation demonstrated the simultaneous accumulation of Cb and ICy-OH in a consistent 1:1 stoichiometric ratio, confirming their synchronized release during the oxidative process (Fig. 1G-H). The reaction reached equilibrium after 90 minutes, with calculated release efficiencies of 86.7% and 85.3% for Cb and ICy-OH, respectively. These findings conclusively demonstrate the efficient H<sub>2</sub>O<sub>2</sub>mediated dissociation of ICy-Cb into its constituent therapeutic components, Cb and ICy-OH.

To ensure the clinical applicability of our probe, we conducted a comprehensive specificity analysis to evaluate the selectivity of ICy-Cb toward H<sub>2</sub>O<sub>2</sub> among various potential biological interferents. The probe was incubated with a diverse array of biologically relevant reactive species including nitric oxide (NO), peroxynitrite (ONOO-), hydroxyl radical (•OH), singlet oxygen (<sup>1</sup>O<sub>2</sub>), superoxide (O<sup>2-</sup>), peroxyl radical (ROO•), tert-butoxyl radical (t-BuO•), and hypochlorous acid (HClO). As demonstrated in Fig. 1I, upon excitation at 640 nm, only the ICy-**Cb**+H<sub>2</sub>O<sub>2</sub> system exhibited significant fluorescence enhancement at 720 nm, while all other analytes induced negligible changes in fluorescence intensity, even after extended incubation periods of 90 minutes. These results unequivocally confirm the remarkable selectivity of ICy-Cb for H<sub>2</sub>O<sub>2</sub> activation, a critical requirement for applications in complex biological systems.

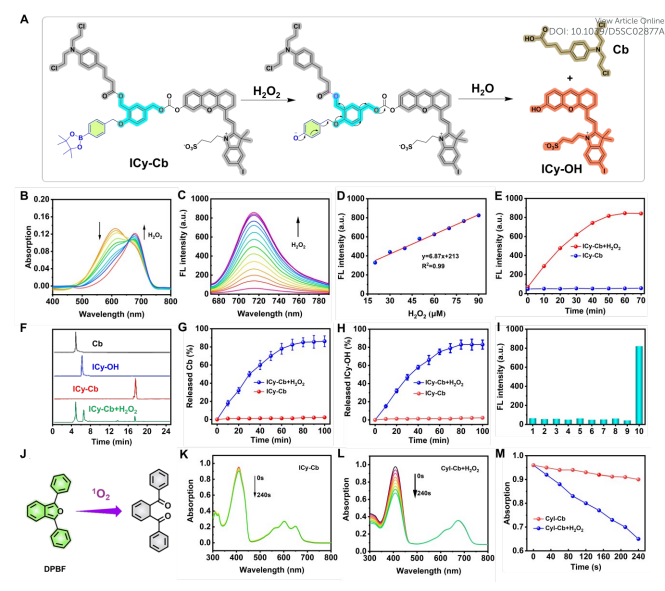
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**Fig. 1**. (A) Schematic illustration of H<sub>2</sub>O<sub>2</sub>-mediated fluorescence turn-on and drug release of **ICy-Cb**. (B) UV-vis and (C) Fluorescence spectra absorbance changes of **ICy-Cb** (5 μM) in PBS buffer (10.0 mM, pH 7.4, containing 30% DMSO), after introduction of H<sub>2</sub>O<sub>2</sub> (0.0–100.0 μM). (D) The linear relationship between I<sub>720 nm</sub> and the concentration of H<sub>2</sub>O<sub>2</sub> (20.0–90.0 μM). (E) Time-dependent fluorescence of **ICy-Cb** with the addition of H<sub>2</sub>O<sub>2</sub> (100.0 μM) in PBS buffer (10.0 mM, pH 7.4, containing 30% DMSO). (F) HPLC chromatographic analysis was performed on several samples: pure **Cb** (black), **ICy-OH** (blue), **ICy-Cb** (red), and a mixture (green) of **ICy-Cb** and H<sub>2</sub>O<sub>2</sub>. Release efficiency of (G) **Cb** and (H) **ICy-OH** from **ICy-Cb**. (I) Fluorescence measurements of **ICy-Cb** (5.0 μM) were recorded in response to H<sub>2</sub>O<sub>2</sub> (100.0 μM) and various other substances (100.0 μM): (1) Blank, (2) NO, (3) ONOO<sup>-</sup>, (4) •OH, (5) <sup>1</sup>O<sub>2</sub>, (6) O<sub>2</sub><sup>-</sup>, (7) ROO<sup>+</sup>, (8) *t*-BuO<sup>+</sup>, (9) HCIO and (10) H<sub>2</sub>O<sub>2</sub>. All measurements were taken 90 minutes after adding the stimulants at room temperature, using an excitation wavelength of 670 nm and an emission wavelength of 720 nm. (J) Schematic illustration of DPBF detecting <sup>1</sup>O<sub>2</sub>. Photodegradation curves of DPBF in the presence of **ICy-Cb** (K) and **ICy-Cb** +H<sub>2</sub>O<sub>2</sub> (L) were obtained under 660 nm light irradiation for varying durations (0-240 s). (M) DPBF degradation (monitored at 415 nm) induced by **ICy-Cb** and **ICy-CH** under irradiation 660 nm irradiation.

To evaluate the ROS generation capacity of our compounds following  $H_2O_2$  activation, we performed comprehensive photochemical analyses based on the promising observation that **ICy-Cb** released **ICy-OH** with restored fluorescence upon  $H_2O_2$  treatment. When liberated **ICy-OH** absorbs irradiation,

molecules are elevated to an excited singlet state, with a portion of this energy being emitted as fluorescence-valuable for tumor imaging and drug tracking applications. The remaining excited molecules undergo intersystem crossing (ISC) to reach the triplet state, subsequently transferring energy to This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence

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molecular oxygen to generate singlet oxygen  $({}^{1}O_{2})$ , thereby enabling PDT. The 1,3-diphenlisobenzofuran (DPBF), as a <sup>1</sup>O<sub>2</sub> capturing agent (Fig. 1J), was used to study the <sup>1</sup>O<sub>2</sub> generation capacity of ICy-Cb, ICy-Cb +H<sub>2</sub>O<sub>2</sub> and ICy-OH by the absorption changes of DPBF at 415 nm. As seen in Fig. 1K-M, the absorbance of DPBF at 415 nm decreased dramatically in the presence of ICy-Cb+H<sub>2</sub>O<sub>2</sub> under 660 nm (10 mW/cm<sup>2</sup>) light irradiation with a degradation curve closely matching that of ICy-OH (Fig. S4). However, under the same conditions, no significant change in the absorption of DPBF in ICy-Cb solution can be observed (Fig. 1L). This is also illustrated by the data in Fig. 1M. These results indicate that ICy-Cb is an activatable PS whose PDT activity can be enhanced by H<sub>2</sub>O<sub>2</sub>. The singlet oxygen quantum yield ( $\Phi\Delta$ ) was calculated to be 0.39% for ICy**Cb** and 2.46% for **ICy-OH**, demonstrating that the  $H_2Q_2$ activated product ICy-OH displays significantly leanance 270\$ production capacity (approximately 6.3-fold increase).

Subsequently, we examined the spectral characteristics of the control probe ICy in response to H<sub>2</sub>O<sub>2</sub> (Fig. S2-3). This probe displayed a primary absorption maximum at 605 nm with essentially no fluorescence emission at 720 nm in its native state. Upon exposure to H2O2, the probe exhibited a pronounced bathochromic shift in absorption to 670 nm, accompanied by significant enhancement of red fluorescence emission at 720 nm ( $\lambda_{ex}$  = 670 nm). This spectral behavior closely paralleled the fluorescence response pattern observed with probe ICy-Cb, further validating our molecular design rationale.

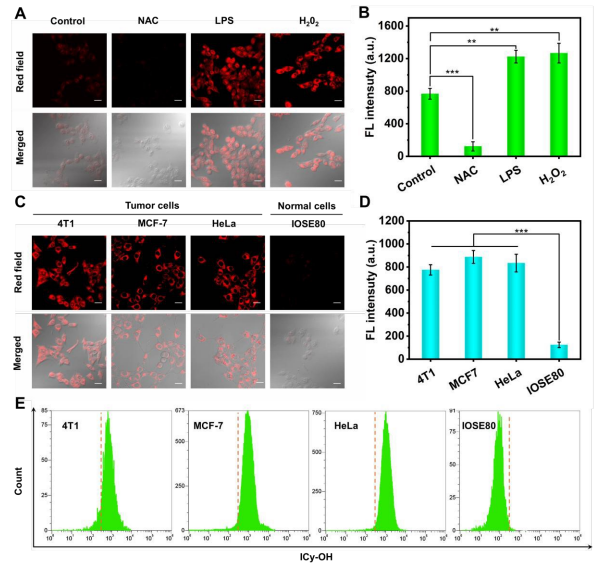


Fig. 2. (A) 4T1 cells pretreated with NAC (500 µM), LPS (1 µg/mL) and H<sub>2</sub>O<sub>2</sub> (100 µM) for 30 min, then incubated with 10 µM ICy-Cb for 60 min. ( $\lambda_{ex}$  = 640 nm,  $\lambda_{em}$  = 670–750 nm) (B) Relative fluorescence intensity from (A). (C) Fluorescence images of tumor cells (4T1, MCF-7, HeLa) and normal cells (IOSE80) incubated with ICy-Cb (10 µM), respectively. (D) Relative fluorescence intensity from (C). (E) Data of flow cytometry after cells (4T1, MCF-7, HeLa and IOSE80) were stained by 10 μM ICy-Cb for 60 min. Scale bar = 10 μm.

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#### Fluorescence imaging of H<sub>2</sub>O<sub>2</sub> in living cells.

To assess the cellular applicability of ICy-Cb given its robust response to H<sub>2</sub>O<sub>2</sub> and potent photodynamic properties, we conducted comprehensive cellular-level H<sub>2</sub>O<sub>2</sub> imaging studies. Initially, we investigated the cellular uptake kinetics of ICy-Cb in 4T1 breast cancer cells. As illustrated in Fig. S5-6, ICy-Cb exhibited detectable fluorescence within 40 minutes of incubation, with signal intensity progressively increasing over time, demonstrating efficient internalization and activation within cancer cells. To evaluate the probe's H<sub>2</sub>O<sub>2</sub>-responsive imaging capabilities, 4T1 cells loaded with ICy-Cb were subjected to various experimental conditions (Fig. 2A-B). In the control group (Fig. 2A), cells treated solely with ICy-Cb exhibited weak fluorescence. However, following treatment with lipopolysaccharide (LPS), which stimulates endogenous H<sub>2</sub>O<sub>2</sub> production,43 a significant enhancement in fluorescence intensity was observed. Similarly, direct addition of H<sub>2</sub>O<sub>2</sub> resulted in significant fluorescence amplification. Confirmatory negative control experiments employing N-acetyl-cysteine (NAC),<sup>44</sup> a well-established reactive oxygen species scavenger, verified that the observed fluorescence modulation was specifically attributable to intracellular H<sub>2</sub>O<sub>2</sub> levels.

To further investigate the subcellular organelle localization, various commercial trackers, including ER-Tracker Green, Mito-Tracker Green, Lyso-Tracker Green and DiO, as well as **ICy-Cb** (10  $\mu$ M), were applied to 4T1 cells for 60 min respectively (Fig. S7). The distribution of **ICy-Cb** in 4T1 overlaps most with ER-Tracker Green, with a correlation coefficient of 0.89. In contrast, the correlation coefficients between **ICy-Cb** and Mito Tracker Green, Lyso-Tracker Green or DiO were found to be 0.75, 0.55 and 0.27 respectively (Fig. S7). Therefore, this molecule is primarily located in the endoplasmic reticulum.

Having characterized the cellular uptake and localization properties, we proceeded to evaluate ICy-Cb's ability to simultaneously detect differential H<sub>2</sub>O<sub>2</sub> levels across various cell types. We performed comparative imaging studies on normal ovarian epithelial cells (IOSE80) and multiple cancer cell lines: 4T1 (breast cancer), MCF-7 (breast cancer), and HeLa (cervical cancer). As shown in Fig. 2C, under identical imaging conditions with 10 µM ICy-Cb, tumor cells exhibited significantly brighter fluorescence compared to normal cells, with quantitative analysis revealing 5-8 folds higher fluorescence intensity in cancer cells (Fig. 2D). Flow cytometry analysis (Fig. 2E) independently confirmed these observations, providing quantitative single-cell resolution data that corroborated the microscopy findings. These results clearly indicate that cancer cells maintain higher basal H2O2 levels than normal cells, consistent with previous studies on cancer redox dysregulation. The ability of ICy-Cb to discriminate between normal and cancer cells based on their inherent  $H_2O_2$  levels demonstrates its potential utility as a cancer-specific imaging agent for diagnostic applications

In vitro cytotoxicity evaluation.

To assess the therapeutic potential of our H2Q2-activatable system, we first investigated whether altivated acy-co-could effectively generate ROS within living cells, a prerequisite for photodynamic therapy efficacy. We employed 2',7'dichlorofluorescin diacetate (DCFH-DA) as an indicator of intracellular ROS. DCFH-DA itself is non-fluorescent; however, upon cellular internalization, it undergoes hydrolysis by intracellular esterases. The resulting compound can then be oxidized by ROS to produce 2,7-dichlorofluorescein (DCF), which exhibits strong fluorescence emission. Our experimental results, as illustrated in Fig. 3B, demonstrated negligible fluorescence in cells treated with ICy-Cb in the absence of light stimulation. In contrast, when these pre-treated cells were subjected to red-light irradiation (660 nm), a significant enhancement in fluorescence intensity was observed (Fig. 3B). These findings provide compelling evidence that ICy-Cb functions as an H<sub>2</sub>O<sub>2</sub>-responsive, activatable PS capable of generating singlet oxygen upon activation, thus demonstrating its potential utility for PDT.

To further validate the intracellular singlet oxygen generation capability of our system, we employed Singlet Oxygen Sensor Green (SOSG) as a specific probe for detecting <sup>1</sup>O<sub>2</sub> in living cells. As shown in Fig. S8, 4T1 cells were initially incubated with ICy-Cb (10  $\mu$ M) for 60 minutes, followed by treatment with SOSG (5 μM) for an additional 30 minutes. Upon irradiation with 660 nm light (10 mW/cm<sup>2</sup>) for 10 minutes, a significant enhancement in green fluorescence (λex = 488 nm,  $\lambda$ em = 500-580 nm) was observed within the **ICy-Cb**-loaded cells. In contrast, control experiments conducted with ICy-Cb without light exposure exhibited minimal SOSG fluorescence, demonstrating that singlet oxygen generation is specifically triggered by light activation of the ICy-Cb system after its H<sub>2</sub>O<sub>2</sub>mediated conversion to ICy-OH. These results provide direct evidence for the intracellular production of singlet oxygen during the photodynamic therapy process, further confirming the mechanistic basis of our dual-modality therapeutic approach.

Having confirmed the ROS-generating capability of our system, we proceeded to systematically evaluate the cytotoxicity profiles of various treatment regimens involving **ICy-Cb** using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) assay, a gold standard for cell viability assessment. Initial assessment of the cytotoxicity on normal ovarian epithelial cells (IOSE80) revealed that **ICy-Cb** exhibited minimal toxicity, maintaining approximately 87% cell viability following 24-hour incubation at concentrations up to 10  $\mu$ M, regardless of light exposure conditions (Fig. S10). This favorable safety profile in normal cells substantiates the selective therapeutic potential of our designed probe, a critical consideration for clinical translation.

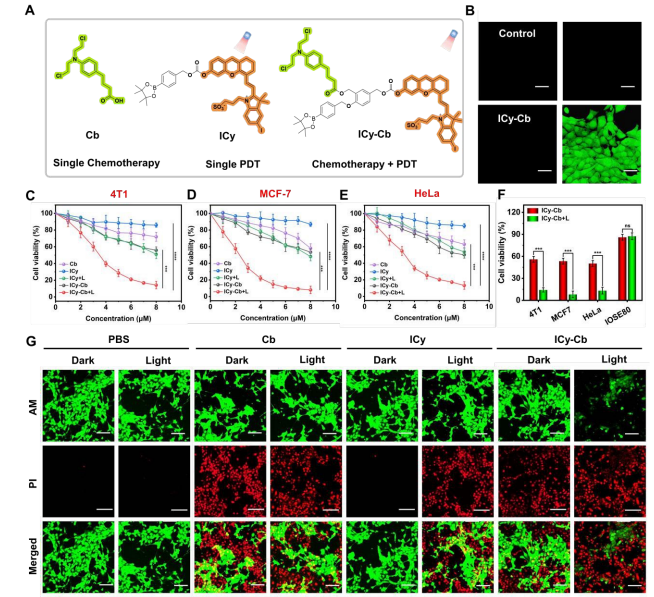
Additionally, we evaluated the cytotoxicity profiles across multiple cancer cell lines-4T1, MCF-7, and HeLa-incubated with varying concentrations of **ICy-Cb**. For comparative analysis, **ICy** and commercial **Cb** were examined under identical experimental conditions (Fig. 3A). As evidenced in Fig. 3C-F, **ICy** exhibited negligible cytotoxicity toward cancer cells under dark conditions. Notably, **Cb**, despite its status as a first-line clinical This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

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chemotherapeutic agent, demonstrated suboptimal cytotoxicity against these cancer cell lines at the tested dosage and treatment frequency. Similarly, **ICy-Cb** displayed limited cytotoxicity under dark conditions, primarily attributable to the released **Cb**, as evidenced by cellular viability trends that paralleled those observed in the **Cb** treatment group. Under light irradiation conditions, **ICy** demonstrated significant cytotoxicity, attributable to the release of **ICy-OH** that facilitates effective PDT-mediated cell killing. Most notably, we observed markedly enhanced cell death when cells were treated with **ICy-** **Cb** followed by light irradiation (Fig. 3C-F). This augmented cytotoxic effect results from the simultaneous action of **Cy OFA** mediated phototoxicity and **Cb**-induced chemotherapeutic effects, effectively achieving the intended dual-modality therapeutic approach. This synergistic effect highlights the significant advantage of our rationally designed  $H_2O_2$ activatable theranostic agent compared to either monotherapy approach, demonstrating superior therapeutic efficacy through the combination of photodynamic and chemotherapeutic modalities.



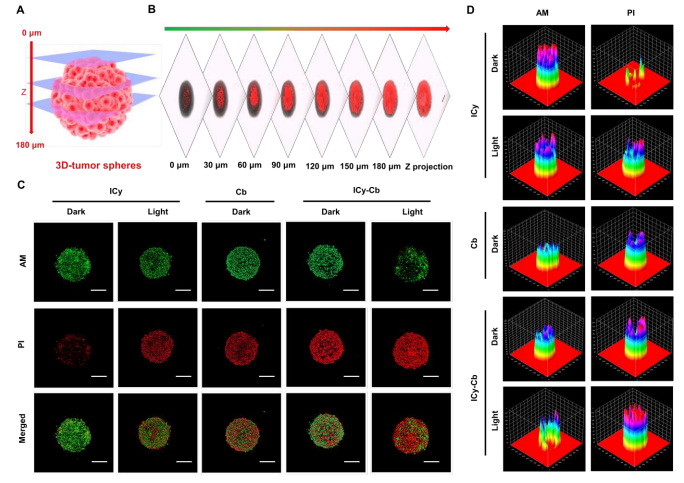
**Fig. 3.** (A) Chemical structure formula of **Cb** (single Chemotherapy), **ICy** (single PDT) and **ICy-Cb** (chemotherapy + PDT). (B) Evaluation of ROS generation in 4T1 cells. Scale bars: 20  $\mu$ m. Evaluation of the cytotoxicity of **ICy-Cb** on (C) 4T1 cells, (D) MCF-7 cells and (E) HeLa cells at various doses under dark or light, NIR light (660 nm, 10 mW cm<sup>-2</sup>, 20 min). (F) Cytotoxicity of **ICy-Cb** (6.0  $\mu$ M) to different cell lines after different treatments. N = 5. (G) AM/PI fluorescent imaging for distinguishing living and dead cells. Scale bars represent 200  $\mu$ m, (*t* tests, *n* = 5, mean  $\pm$  SD, \**p* < 0.05, \*\**p* < 0.001, \*\*\*\**p* < 0.0001).

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To visually confirm our cytotoxicity findings, we conducted additional investigations using dual fluorescent staining with calcein acetoxymethyl ester (calcein-AM) and propidium iodide (PI). Calcein-AM generates green fluorescence exclusively in viable cells, while PI produces red fluorescence when intercalated with nucleic acids in dead cells. As depicted in Fig. 3G, these visual results corroborated our quantitative MTT data: the cytotoxic efficacy of **ICy-Cb** under dark conditions was limited and comparable to that of **Cb** alone. However, following

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light exposure, cells throughout the entire microscopic field exhibited prominent red fluorescent Dosignals, and the provide the provided of the provided of the provides compelling evidence for the potent therapeutic efficacy of our H<sub>2</sub>O<sub>2</sub>-activatable dual-modality system when both the photosensitizer and chemotherapeutic components are simultaneously activated.



**Fig. 4.** (A) Diagrammatic illustration showing the z-slice scanning methodology used for cellular sphere analysis. (B) Z-axis fluorescence imaging series of tumor spheroids following **ICy-Cb** treatment. The Z-Projection represents the composite overlay of all sequential images in the column. Scale bar represents 150 μm. (C) Visualization of spheroids using Calcium-AM/PI dual staining protocol after various experimental treatments. Scale bar indicates 150 μm. (D) Three-dimensional surface plots depicting fluorescence intensity distribution throughout the spheroids along the z-axis.

#### Penetration and toxicity testing of probe on 3D-tumor spheres.

To bridge the gap between two-dimensional cell culture findings and in vivo applications, we constructed 4T1-derived multicellular spheroids (MCSs) to systematically investigate the tumor penetration capabilities and  $H_2O_2$ -catalytic activity of **ICy-Cb** under conditions that better mimic the complexity of solid tumors. A key challenge in cancer therapy is achieving sufficient penetration of therapeutic agents throughout the tumor mass. To evaluate this critical parameter, we performed z-axis scanning confocal microscopy of tumor spheroids after incubation with **ICy-Cb** (10  $\mu$ M, 4 hours). As demonstrated in Fig. 4A and B, sequential scanning images along the z-axis (methodology illustrated in Fig. 4A) revealed that **ICy-Cb** exhibited exceptional penetration capability. Remarkably, substantial fluorescence intensity was detected even at depths of 180  $\mu$ m within the spheroids (Fig. 4B), indicating activation of the probe throughout the tumor mass. This deep penetration capability is particularly significant as it addresses one of the major limitations of conventional photodynamic therapy and chemotherapeutic approaches.

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Having established the penetration characteristics, we proceeded to evaluate the therapeutic efficacy of our dual-modality system in these 3D tumor models. Consistent with our twodimensional cell culture findings, the vast majority of cells within tumor spheroids treated with ICy under dark conditions remained viable (Fig. 4C). Spheroids exposed to Cb exhibited limited cytotoxicity, indicating that at equivalent dosages, Cb demonstrates restricted therapeutic efficacy, similar to that observed with ICy-Cb under dark conditions. These observations in three-dimensional tumor models were consistent with our previous two-dimensional cell culture findings. Furthermore, ICy-treated spheroids exhibited partial cytotoxicity when exposed to light irradiation. In contrast, ICy-**Cb** treatment followed by light exposure resulted in nearly complete cell death throughout the spheroid structure (Fig. 4C). These results maintain the same efficacy trend observed in our previous MTT experiments. Through three-dimensional surface analysis of the fluorescence images, (Fig. 4D) we confirmed our hypothesis regarding the enhanced penetration and therapeutic efficacy of our dual-modality system in three-dimensional tumor models.

To quantitatively evaluate the therapeutic efficacy in our 3D tumor model, we performed CellTiter-Glo@3D (CTG-3D) assays on spheroids following various treatment regimens. This ATPbased viability assay provides direct quantification of metabolically active cells within the 3D structures. As shown in Fig. S9, treatment with ICy-Cb followed by light irradiation (660 nm, 10 mW/cm<sup>2</sup>) resulted in the most significant reduction in luminescence intensity, indicating substantial decrease in viable cells compared to all other treatment groups. Moderate reductions in viability were observed in spheroids treated with either ICy plus light (PDT only) or ICy-Cb without light (chemotherapy only), while Cb alone exhibited limited efficacy. These quantitative data strongly align with our qualitative observations from the Calcein-AM/PI imaging, confirming that the combination of photodynamic and chemotherapeutic modalities provides superior efficacy against 3D tumor models compared to either treatment alone. The enhanced performance in these structurally complex spheroids, which better mimic solid tumors than monolayer cultures, further supports the potential translational value of our dual-modal system.

#### In vivo antitumor therapy.

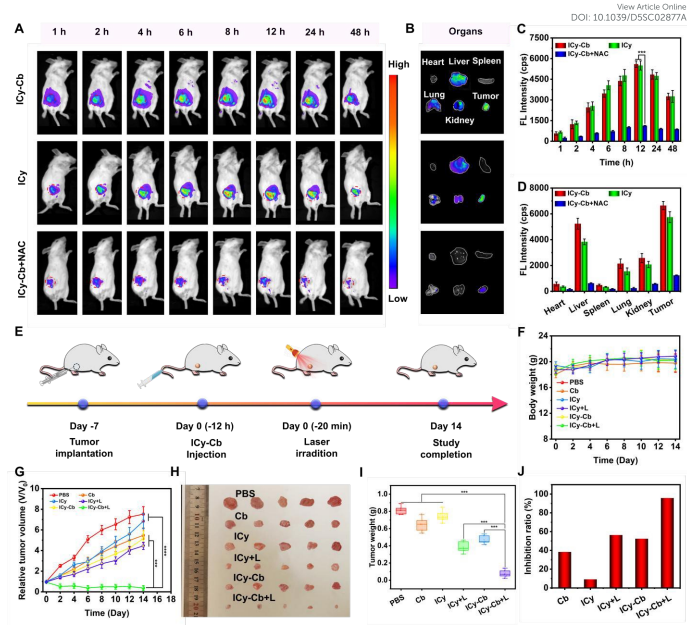
Based on the promising in vitro results, we investigated the potential of **ICy-Cb** as a tumor-targeted prodrug in vivo. We established 4T1-bearing BALB/c mouse models for these experiments. Following tail vein injection of either **ICy-Cb** or **ICy**, we monitored activation and biodistribution using an in vivo imaging system (IVIS). As a negative control, NAC was administered para-tumorally to inhibit  $H_2O_2$ . As shown in Fig. 5A and C, minimal fluorescence signals were detected during the first 2 hours post-injection of **ICy-Cb**. However, fluorescence intensity at tumor sites progressively increased over time, with substantial signals still present at 24 hours. Mice receiving **ICy** 

demonstrated comparable fluorescence activation, patterns. Conversely, tumors pre-treated with NAC showed significantly reduced fluorescence throughout the observation period (Fig. 5A and C), confirming the compounds' selective activation in high-ROS tumor environments. At the 48-hour timepoint, mice were euthanized for ex vivo imaging of tumors and major organs, revealing that most activated **ICy-Cb** or **ICy** remained localized at tumor sites, with metabolic clearance occurring through hepatic and renal pathways (Fig. 5B and D). NACpretreated mice displayed lower overall signal intensity due to reduced **ICy-Cb** activation.

Having confirmed tumor-targeting capabilities, we evaluated therapeutic efficacy in tumor-bearing mice (Fig. 5E). Seven days after subcutaneous 4T1 cell implantation, mice with comparable tumor volumes were randomly assigned to six treatment groups (n=5): (1) PBS (control), (2) Cb (chemotherapy), (3) ICy alone, (4) ICy + NIR light (PDT only), (5) ICy-Cb (chemotherapy only), and (6) ICy-Cb + NIR light (combination therapy). For light-treated groups, tumors were irradiated with 660 nm light (50 mW/cm<sup>2</sup>, 20 min) 12 hours post-injection. Throughout the 14-day treatment period, body weight and tumor dimensions were recorded every other day. Results shown in Fig. 5F-I revealed no significant weight loss across all treatment groups, indicating minimal acute toxicity of ICy-Cb (Fig. 5F). Tumor growth curves (Fig. 5G) demonstrated rapid tumor progression in the PBS control group. The clinical agent Cb showed limited tumor inhibition at the administered dose and frequency. While ICy alone had negligible impact on tumor growth, ICy-Cb without light activation produced modest tumor growth delay, attributed to the chemotherapeutic effect of Cb released following H<sub>2</sub>O<sub>2</sub> activation (Fig. 5G-I). The ICy plus light group exhibited significant tumor inhibition (TGI=57%) due to the photodynamic effect of released ICy-OH within the tumor microenvironment. Most notably, mice receiving ICy-Cb with light exposure demonstrated superior tumor growth inhibition, achieving a 92% inhibition rate compared to either monotherapy approach (Fig. 5J). This enhanced therapeutic outcome can be attributed to the synergistic effects of combined chemotherapy and photodynamic therapy. After a 14-day treatment period, the mice were humanely euthanized and the tumors were surgically excised. Subsequently, tumor weights were measured and photographic evidence was obtained to further validate the observed growth patterns (Fig. 5H and 5I). These organs were then processed into paraffin sections and subjected to hematoxylin and eosin (H&E) staining. As illustrated in Fig. S11, no discernible differences were observed in the organs of the PDT group compared to those treated with PBS. Moreover, examination revealed no abnormalities in physiological morphology or histopathological lesions, suggesting minimal toxicity of ICy-Cb to normal tissues. The comprehensive findings validate ICy-Cb as a highly efficient activatable chemo-photodynamic agent, and thereby demonstrating its significant potential for both tumor imaging and growth inhibition.

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# Conclusions

In conclusion, this study introduces a H<sub>2</sub>O<sub>2</sub>-responsive unimolecular phototheranostic agent, ICy-Cb, that strategically integrates cancer imaging and synchronized dual-modal therapy. In the structure of this engineered system, both a photosensitizer and a chemotherapeutic drug are integrated into a single molecular design that selectively reacts to the high levels of  $H_2O_2$  present in the tumor microenvironment. The molecule effectively transitions from inactive to active states upon H<sub>2</sub>O<sub>2</sub> exposure, simultaneously releasing both therapeutic components while generating near-infrared fluorescence for real-time visualization. ICy-Cb demonstrated exceptional tumor selectivity with 5 to 8 folds higher activation in cancer cells, deep tissue penetration (180  $\mu m)$  in 3D tumor models, and sustained tumor accumulation in vivo. The synergistic effect of combined chemo-photodynamic therapy resulted remarkable tumor growth inhibition (92%), substantially outperforming either monotherapy. Importantly, minimal toxicity to normal tissues was observed throughout treatment. Collectively, ICy-Cb represents a significant advancement in precision cancer theranostics by enabling spatiotemporally controlled combination therapy while minimizing systemic toxicity.

# Author contributions

J. H. and J. F. conceived and designed the experiments for this project. J. H. performed the synthetic work and carried out the characterization of the compounds. J. H. and K. L. contributed to the photophysical experiments. J. H. and M. Y. conducted data analysis and prepared the initial manuscript, with all other authors contributing to revisions. X. P. supervised the work. All authors discussed the results and provided feedback on the manuscript.

# **Conflicts of interest**

The authors declare no competing financial interests.

# **Ethical statement**

All procedures involving animals adhered to the guidelines outlined in the Guide for the Care and Use of Laboratory Animals issued by the National Institutes of Health. The research protocol received approval from the Animal Ethics Committee at Dalian University of Technology (protocol number: 2018-043).

# Data availability

The data that support the findings of this study are available in the Supporting Information of this article.

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The data supporting this article have been included as part of the Supplementary

Information.