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Singlet oxygen ($^{1}O_{2}$) plays a crucial role in various biological processes, including cellular signaling and oxidative stress, necessitating reliable methods for its detection and quantification. However, detecting 10_2 in biological systems is challenging due to its high reactivity and short lifespan. Here, we developed dually-labelled polymeric micelles (SOSG@Cy5-PIC/m) capable of detecting $^{1}O_{2}$ and reporting probe positions. These micelles are formed using oppositely charged diblock copolymers, Cy5-conjugated PEGpoly(α,β -aspartic acid) and Cy5-conjugated PEG[1]poly([5-aminopentyl]- α,β -aspartamide), which selfassemble in aqueous solution. Singlet Oxygen Sensor Green (SOSG) molecules, which fluoresce upon interaction with ¹O₂, are integrated into the micelle core by condensation to the amines in the poly([5aminopentyl]- α , β -aspartamide) block. The micelle core is crosslinked by forming amide bonds between the carboxylates in the poly(α,β [1]aspartic acid) segment and the amines in the poly([5-aminopentyl]- α,β -aspartamide) block, ensuring stability in biological environments. The micelles are around 30 nm and the dyes in their core are protected by a dense PEG shell. The micelles quantitatively detect 10_2 in media and enable visualization of ${}^{1}O_{2}$ in 3D cellular spheroids under induced cellular stress. Potential applications include advancing research tools for understanding ${}^{1}O_{2}$, diagnostics for oxidative stressrelated diseases and exploring targeted therapies, such as photodynamic therapy. Future developments may enhance sensitivity and targeting capabilities for broader biomedical applications.

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Dual-Labelled Polymeric Micelles for Singlet Oxygen Reporting in 7 Biological Systems

- 8 Yasuhiro Nakagawa^{1, 2, 3†}, Hayato Laurence Mizuno^{2,3}, Yuta Ushimaru¹, Jumpei Norimatsu¹,
- 9 Kazunori Igarashi^{1, 2}, Keita Masuda¹, Madoka Takai¹, Yasutaka Anraku^{2, 3}, Horacio Cabral^{1, 2*}

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Visualizing singlet oxygen $({}^{1}O_{2})$ in biological systems could greatly enhance our understanding of its biological roles and offer near diagnostics and therapeutics. However, ${}^{1}O_{2}$ is unstable and highly reactive, making its detection in living systems a significal challenge. To address this, we have developed dually-labelled polymeric micelles designed to trace both the location and levels 48 ${}^{1}O_{2}$.

Reactive oxygen species (ROS), e.g., super oxide anions (O₂))6 hydrogen peroxide (H_2O_2), and 1O_2 , are important mediators in Δa variety of biological and pathological processes [1, 2]. The ROS are involved in essential biological functions, such as signalling metabolism, and regulation of the immune system [3]. When ROS and overproduced or antioxidants are depleted, these reactive species cause oxidative stress, oxidizing lipids, proteins, carbohydrates, RNA2 and DNA, which leads to cellular damage. Such damage is linked 53 various diseases. including cardiovascular disease. neurodegenerative disorders, and cancer [4, 5, 6, 7]. Moreover, high ROS levels can be used for therapeutically damaging cancer cells [8]. In fact, a major physiological mechanism to kill tumour cells is the production of ROS by neutrophils and macrophages [9]. Moreover, certain anticancer drugs can induce ROS production. For instance, oxaliplatin, a widely used chemotherapeutic agent [10], can directly affect mitochondria, inducing mitochondrial permeability transition and ROS generation [11, 12]. Thus, an improved understanding of the role of ROS in living systems can provide opportunities for developing innovative diagnostic and therapeutic strategies.

36 Detection of ${}^{1}O_{2}$ has been attracting much attention, as it is 37 involved in various biological phenomena by direct reaction with

1. Department of Bioengineering, Graduated School of Engineering, The University of Tokyo, 7-3-1 Hongo, Bunkyo-ku, Tokyo, 113-8656, Japan 7

2. Innovation Center of NanoMedicine, Kawasaki Institute of Industrial Promotion, 3-25-14 Tonomachi, Kawasaki-ku, Kawasaki, 210-0821, Japan

3. Department of Materials Science and Engineering, Graduated School of Materials and Chemical Technology, Tokyo Institute of Technology, 2-12-1 Ookayama, Meguro-ku, Tokyo, 152-8550, Japan † Y.N. has moved to Mitsui Chemicals, Inc. (Tokyo, Japan).

Supplementary Information available: Materials and methods. SeeL DOI: 10.1039/x0xx00000x biomolecules and regulation of intracellular signals [13, 14]. However, because ¹O₂ has a short lifetime, high reactivity, and low concentration, its detection has been challenging. Several approaches have been developed for the detection of ¹O₂, including direct measurement of ¹O₂ phosphorescence (transition light from singlet to triplet state) at 1280 nm [15], electron spin resonance spectroscopy (ESR) [16], photo/chromo-sensitizers [17], and fluorescent probes [18]. Despite these major efforts, in situ observation is still limited. Challenges include low phosphorescence quantum efficiency (\sim 10⁻⁶ for aqueous systems and even smaller for systems containing ¹O₂ quencher) [<u>19</u>], inefficient conversion of ¹O₂ to stable radical species for ESR [16], toxicity and cellular damage from photo/chromo-sensitizers [20], and low stability and uncertain distribution of fluorescent probes in tissues and cells. Thus, innovative approaches are still necessary for effective ¹O₂ sensing in biological systems.



Figure 1. Schematic illustration of SOSG/Cy5-labeled PIC micelles (SOSG@Cy5-PIC/m). The conjugated SOSG enables the detection of ${}^{1}O_{2}$ by changing its fluorescent characteristic, while the Cy5 constantly reports the position of the probes in biological systems.

Polymeric micelles, *i.e.*, core-shell polymeric structures, have demonstrated great potential for targeted therapy and diagnosis by delivering a wide variety of bioactive compounds loaded in their core [21]. Moreover, the shell of the polymeric micelles can protect the

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63 bioactive agents in the core from biological degradation. Thus? 64 polymeric micelles can navigate and spatiotemporally control the function of the payloads in biological settings [22]. Polymeric 65 micelles can be assembled in aqueous environments by controling 66 the interaction of the forming block copolymers, such as $\frac{163}{123}$ 67 hydrophobicity and electrostatic interaction between the $c\overline{qr_{4}}$ 68 forming segments. Particularly, polyion complex (PIC) micelles have 69 70 been prepared using poly(ethylene glycol) (PEG) based oppositeba charged segments of the block copolymer pair of PEG-poly(anip)な 71 72 and PEG-poly(cation). These PIC micelles (PIC/m) offer substantial 73 advantages as reporter agents, as they allow effective size control, 74 softness, and versatile drug loading capability [23].

Here, we developed a nano-scaled reporter based on PIC/m that is capable of indicating its distribution in living systems and sensing the presence of ${}^{1}O_{2}$ in their surroundings. These reporters were made by mixing oppositely charged diblock copolymers, i.e., Cy5conjugated PEG-poly(α, β -aspartic acid) (PEG-PAsp) and Cy5-580 conjugated PEG-poly([5-aminopentyl]- α , β -aspartamide) (PEG-P(Asp-<u></u>81 AP)), in an aqueous solution [24]. The PIC/m were also stabilised by ₹82 cross-linking the core through 1-ethyl-3-(3-√suo 184 dimethylaminopropyl)carbodiimide (EDC) coupling to avoid micelle dissociation in biological environments. As cross-linked PIC 85 structures are still permeable to small biomolecules [24], Singlet \$86 Oxygen Sensor Green (SOSG) molecules, which are precise reporters of ${}^{1}O_{2}[25, 26]$, were conjugated to the core of the micelles to provide specific sensitivity for fluorescent detection of ${}^{1}O_{2}$ (Figure 1) [18] ü88 <u>5</u>89 After purification to remove the unreacted SOSG and EDC, the size Ĕ90 and polydispersity index of the dual-labelled micelles (SOSG@d/29 ັ ສິ91 PIC/m) were 31 nm and 0.08, respectively (Figure 2a). <u>8</u>92 incorporation of Cy5 and SOSG was confirmed by fluoresceந்த் .≌93 spectrometry (Figure 2b), with the emission spectrum of the <u>9</u>294 SOSG@Cy5-PIC/m showing peaks at 540 nm that correspond to SOSG ^{te}95 and 680 nm that relate to Cy5. These results indicate successible preparation of SOSG@Cy5-PIC/m.

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The ${}^{1}O_{2}$ detection capacity of SOSG@Cy5-PIC/m was evaluated 17 8 by using EP, which is a convenient reagent for this purpose, as it $\frac{1}{238}$ 9 quantitatively generate ${}^{1}O_{2}$ in solution by simply heating above $3\overline{139}$ λa without using light or photosensitisers. The ${}^1\text{O}_2$ dose-dependent)() fluorescence emission of SOSG@Cy5-PIC/m showed a gradual 11 102 increase of the peak at 539 nm (Figure 2c), similar to the behavior observed with free SOSG. The fluorescence intensity of SOSG@d/43103 104 PIC/m was stronger than that of free SOSG under 1×10^{-3} M 16^{4}_{2} presumably due to reduced inter-SOSG energy transfer that cauted 105 106 quenching of free SOSG. The increase in fluorescence intensity 107 exhibited high linearity in relation to ${}^{1}O_{2}$ concentration, indicating that the micelles can quantitatively detect ${}^{1}O_{2}$ (Figure 249108 Importantly, the Cy5 fluorescence remained unaffected by ${}^{1}O_{2}$ level 9 109 110 (Figure 2e), reinforcing its reliability as a stable marker for PIC/m 111 concentration. This stability enables the use of Cy5 fluorescence normalize SOSG signal, providing a robust ratiometric approach 151 112 152 113 precise quantification of ¹O₂ in biological samples. 153

The time-dependent cellular uptake of SOSG@Cy5-PIC/m 1/54 114 115 examined in monolayer and spheroid cultured cells of huma55 116 pancreatic adenocarcinoma (BxPC3) in vitro. Polymeric micelles willo 117 their size precisely controlled below 50 nm display great permeability

against tumour tissues [21]. Thus, time-dependent internalizations of SOSG@Cy5-PIC/m in both cell culture systemsowere/observed by confocal laser scanning microscopy (CLSM) (Figures 3a-c). The internalization of the micelle in each cell was evaluated as relative fluorescent units (RFU) per cell or spheroid. The cells cultured in monolayer showed a time-dependent increase of the RFU/cells, which became saturated at 6 h incubation (Figure 3b). Spheroid cultured cells also showed a time-dependent increase of the RFU/spheroid signal. However, the fluorescence intensity in the spheroids kept increasing even after 24 h incubation (Figure 3c).

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Figure 2. Characterization of the SOSG@Cy5-labeled PIC/m (SOSG@Cy5-PIC/m). (a) Size distribution of SOSG@Cy5-PIC/m determined by DLS. (b) Fluorescent spectra of SOSG@Cy5-PIC/m, SOSG, and Cy5 (Excitation (Ex) = 460 nm). (c) ¹O₂ dose-dependent fluorescence intensity of SOSG@Cy5-PIC/m (Ex = 460 nm). (d) ¹O₂ dose-dependent fluorescence intensity of SOSG@Cy5-PIC/m at an emission wavelength of 530 nm. (e) ¹O₂ dose-dependent fluorescence intensity of Cy5 in PIC/m (Ex = 633 nm).

The time-dependent penetration of the SOSG@Cy5-PIC/m into the spheroid core of SOSG@Cy5-PIC/m was analysed by taking the fluorescence profile across the spheroids and summarised as histograms (Figure 3c). These histograms indicate that the SOSG@Cy5-PIC/m penetrated the spheroids time-dependently. At 1 h incubation, the micelles were barely detected in the periphery of the spheroids. Extending the incubation time to 3-6 h increased the intensity of micelles in cells located at the border of the spheroids. This also resulted in a gradual decay of the fluorescence profile toward the spheroid core. After 24 h incubation, the micelles were observed throughout the entire spheroid, including the centre of the spheroid. These results suggest that imaging the spheroid culture at 24 h is an optimal time point for detecting ¹O₂, as the micelles are distributed throughout the entire cluster of cells

Spheroids more accurately mimic the 3D structure, cell-cell interactions, and microenvironment of tissues in vivo. Therefore, we addressed the proof-of-concept for in situ detection of ¹O₂ by using SOSG@Cy5-PIC/m in the spheroids of BxPC3 cells. In this experiment, oxaliplatin, which is a clinically approved drug that can induce ${}^{1}O_{2}$ by mitochondria damage [12], was used as a trigger for activating the SOSG signal in the micelles. Thus, free SOSG and SOSG@Cy5-PIC/m

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were first incubated with the spheroid-cultured BxPC3 cells for $2\frac{1}{2}$ 157 Then, oxaliplatin was administrated to the cells and incubated for $\frac{1}{197}$ 158 159 , 3-, 6- and 24-h to induce ${}^{1}O_{2}$ generation. The distribution of the 160 micelles and the generation of ${}^{1}O_{2}$ inside the cells were imaged **198** 161 CLSM. The SOSG signal from free SOSG and SOSG@Cy5-PIC/m was 162 not observed in the cells exposed to oxaliplatin for 1 to 6 h (data not 163 shown). However, the ¹O₂ molecules produced after 24 h incubation 164 with oxaliplatin were detectable by the SOSG in the SOSG@Cy5-165 PIC/m (Figure 4a). In contrast, the signal from free SOSG was 166 undetectable. Previous studies have indicated that the access of 167 SOSG into living mammalian cells presents major hurdles [25] and 168 that SOSG is sensitive to extracellular ¹O₂ molecules, which activates **1**69 the probes before being internalized by the cells [27]. A comparison .¥70 of the fluorescence intensity between free SOSG and SOSG@Cy5-1 1 1 1 1 7 1 1 7 1 1 1 7 1 PIC/m revealed a 15-fold increase in the SOSG signal intensity (Figure 4b). The fluorescence intensities of the SOSG in SOSG@Cy5-PIC/m £73 per spheroid were also evaluated by normalizing the SOSG signal **1**74 with the Cy5 signal (Figure 4c). Compared to the spheroids treated <u></u>**1±**75</u> with saline, the oxaliplatin-treated spheroids showed almost 2-fold Ī76 higher SOSG/Cy5 ratio (p < 0.05). These results support the ability of <u></u> <u></u> <u></u> <u></u> <u></u> 77 SOSG@Cy5-PIC/m to detect ¹O₂ in cellular systems. Given that ROS generation is commonly employed to induce cell death in cancer **1179** therapies, SOSG@Cy5-PIC/m offers a valuable tool for monitoring the therapeutic efficacy of such treatments. Additionally, the system's capability to encapsulate ROS-generating drugs, such an oxiplatin used in this study, within its core enables a theranoptig approach that can simultaneously treat and track therapeutic efforts with a single carrier. With further improvements and tunings SOSG@Cy5-PIC/m holds the potential of introducing a new paradigging in cancer theranostics, enabling the concurrent generation 2007 detection of ROS. 208



189 Figure 3. (a) Time-lapse images of monolayer/spheroidal cultured BxPC3 cells 190 with SOSG@Cy5-PIC/m observed by confocal laser-scanning microscopy (Blue: Nuclei [Ex/Emission (Em) = 405/446 nm], Red: SOSG@Cy5-PIC/m 191 [Ex/Em = 633/661-750 nm]). (b) Time-dependent uptake of SOSG@Cy5-PI6/m 192 193 for single cell/spheroid against monolayer/spheroidal cultured BxPC3 cells in = 3), and all plots were statistically analysed by the Tukey-Kramer method (194 227

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< 0.005). (c) Quantification of internalised SOSG@Cy5-PIC/m against BxPC-3 spheroid. The relative intensity (RFU/pixel) is calculated from the pixel by pixel integration of the RFU on the white dotted line in Figure 3(a) (spheroid).



Figure 4. (a) Fluorescent imaging analysis of SOSG@Cy5-PIC/m against spheroidal cultured BxPC3 cells with oxaliplatin for 24 h co-incubation (Red: Cy5 [Ex/Em = 633/661-750 nm], Green: SOSG-EP [Ex/Em = 488/499 -552 nm], yellow: colocalised micelles and SOSG). Scale bar: 50 µm. (b) Fluorescent intensity of oxaliplatin-stimulated SOSG in the micelles versus free SOSG with spheroidal culture (c) Fluorescent intensity ratio of SOSG/Cy5 in the spheroidal culture. The fluorescence intensity of SOSG/Cy5 per spheroid was calculated. Error bar means standard deviation (n = 4), and all plots were statistically analysed by the Student's t-test (*p < 0.05)

Conclusions

We successfully developed SOSG/Cy5-labeled PIC/m as probes for the detection of ¹O₂ in biological environments. The SOSG@Cy5-PIC/m probes were internalised by BxPC3 pancreatic cancer cells in a time-dependent manner and were able to penetrate into the core of spheroidal cultures of BxPC3 cells. Thus, using these probes, we successfully detected the in situ ${}^{1}O_{2}$ generated by oxaliplatin in the spheroids. This method shows promise for the local analysis and quantification of ${}^{1}O_{2}$ in biological systems. The SOSG/Cy5-labeled PIC/m could be applied in a wide variety of treatments based on the oxidative responses, leading to an improved understanding of biological mechanisms, diagnostic tools, and therapeutic strategies.

Conflicts of interest

There are no conflicts to declare.

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234 Data availability

235 A data availability statement (DAS) is submitted alongside.

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Data availability statement

The authors confirm that the data supporting the findings of this study are available within the article. Raw data that support the findings of this study are available from the corresponding author, upon reasonable request.