In vivo O$_2$ imaging in hepatic tissues by phosphorescence lifetime imaging microscopy using Ir(III) complexes as intracellular probes

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Phosphorescence lifetime imaging microscopy (PLIM) combined with an oxygen (O$_2$)-sensitive luminescent probe allows for high-resolution O$_2$ imaging of living tissues. Herein, we present phosphorescent Ir(III) complexes, (btp)$_2$Ir(acac-DM) (Ir-1) and (btp-OH)$_3$Ir (Ir-2), as useful O$_2$ probes for PLIM measurement. These small-molecule probes were efficiently taken up into cultured cells and accumulated in specific organelles. Their excellent cell-permeable properties allowed for efficient staining of three-dimensional cell spheroids, and thereby phosphorescence lifetime measurements enabled the evaluation of the O$_2$ level and distribution in spheroids, including the detection of alterations in O$_2$ levels by metabolic stimulation with an effector. We took PLIM images of hepatic tissues of living mice by intravenously administrating these probes. The PLIM images clearly visualized the O$_2$ gradient in hepatic lobules with cellular-level resolution, and the O$_2$ levels were derived based on calibration using cultured cells; the phosphorescence lifetime of Ir-1 gave reasonable O$_2$ levels, whereas Ir-2 exhibited much lower O$_2$ levels. Intravenous administration of NH$_4$Cl to mice caused the hepatic tissues to experience hypoxia, presumably due to O$_2$ consumption to produce ATP required for ammonia detoxification, suggesting that the metabolism of the probe molecule might affect liver O$_2$ levels.

Molecular oxygen (O$_2$) is essential for cellular function and is continuously supplied to whole tissues in the body to maintain homeostasis. Oxygen deprivation (hypoxia) in tissues is associated with the pathophysiology of various diseases such as cerebral infarction, fatty liver disease, chronic kidney disease, diabetic retinopathy, and cancer$^{1-3}$. Understanding the cellular and molecular mechanisms underlying hypoxia-associated diseases requires O$_2$ imaging technology capable of detecting the tissue oxygen status in real time and with high resolution. In vivo O$_2$ detection methods that have been developed so far include oxygen electrodes$^{4,5}$, blood oxygenation level dependent magnetic resonance imaging (Bold MRI)$^{6,7}$, positron emission tomography (PET) with a hypoxia tracer$^8$, electron paramagnetic resonance (EPR) oximetry$^{9,10}$, hypoxia markers$^{11-13}$, and optical imaging$^{14-16}$. These methods can work in vivo, but they have advantages and limitations in terms of applicable targets, spatial resolution, tissue permeability, convenience, reversibility, etc. Of these known methods, optical imaging utilizing O$_2$ quenching of probe phosphorescence has great advantages in that high-resolution O$_2$ images can be obtained in a reversible manner, although it is limited to the detection of a penetration depth through tissues of ~1 cm even under near-infrared excitation$^{17}$.

To visualize tissue O$_2$ levels with high sensitivity and high stability, various phosphorescent metal complexes have been developed over the past few decades$^{14-16,18-15}$. The most common compounds used as luminescent O$_2$ probes include Pt(II) and Pd(II) porphyrins, Ru(II) complexes, Pt(II) complexes, and Ir(III) complexes, which give intense phosphorescence in the visible to near-infrared wavelength regions with reasonably long lifetimes (>1.0 μs). Oxygen probes targeting various biological tissues of interest have been developed by modifying these...
O₂-sensing luminophores: cell-penetrating conjugates of Pt(II)-porphyrins20,21, dendritic Pt(II)- and Pd(II)-porphyrins22–24, Ru(II) complex derivatives targeting the cell nucleus25, ratiometric O₂ probes based on Ir(III) complexes26,27, etc.

With the use of phosphorescent probes, wide-field luminescence lifetime measurements using a gated CCD camera have enabled whole-body O₂ imaging including tumor tissues, ocular fundus, and specific organs28–31. In contrast, phosphorescence lifetime imaging microscopy (PLIM)32,33 allows high-resolution O₂ imaging at the cellular level, providing precise information on microenvironment along with real-time changes in O₂ levels. The PLIM method has been applied to O₂ imaging of giant cells34, cell spheroids20,24,35,36, neurospheres37,38 and the epithelium of rat and human colon tissues38 using cell-permeable small-molecule and nano-particle probes. By using dendrimerized Pt-porphyrin probes, the PLIM method has been successfully applied to microvascular and interstitial O₂ imaging of the brain39,40, bone marrow41,42, and retinal tissue43 in vivo.

In contrast, few studies have reported tissue oxygen distributions in vivo using an intracellular O₂ probe44. We have recently developed small-molecule O₂ probes based on Ir(III) complexes18,19. Among them, BTPDM1 ((btp)2Ir(acac-DM); Ir-1) and (btp-OH)3Ir ((btp-OH)3Ir; Ir-2) (Fig. 1A) allows for the measurement of O₂ levels in hypoxic tumors45,46 and high-resolution O₂ imaging in renal cortex in vivo47,48. To improve the reliability of the O₂ level as measured with a small-molecule probe, it is necessary to quantify O₂ levels in different tissues in vivo and investigate various issues: localization and stability of probes in tissues, calibration of emission lifetime in tissues (cell-specific calibration), the influence of probe administration on the target tissues, etc.

In this study, we established an in vivo O₂ imaging method using small-molecule probes by investigating the O₂ distribution in the hepatic lobules of living mice by PLIM measurements with two structurally related Ir(III) complexes, Ir-1 and (btp-OH)3Ir (Ir-2) (Fig. 1A). The hepatic lobule is a fundamental structural unit of the liver with a hexagonal shape (Fig. 1B)49,50. In the lobule, blood runs unidirectionally from the vertices of the hexagon called portal triads, through microcirculatory networks known as sinusoids, to a central vein (CV) in the middle of the lobule. The portal triad contains two distinct inlets for blood flow: a well-oxygenated hepatic artery and a poorly-oxygenated portal vein, the latter of which constitutes ~70% of total liver blood flow. Therefore, even under physiological conditions, liver parenchymal cells and hepatocytes surrounding the portal triad are
exposed to relatively low oxygen concentrations, and pericentral hepatocytes experience much lower oxygen concentrations, forming a zonal heterogeneous distribution of oxygen along the sinusoids in the hepatic lobule. These intralobular oxygen distributions, however, are considered to be prerequisite for normal liver metabolic functions in a zone-specific manner. In addition, this metabolic zonation is often disrupted in disease conditions with aberrant oxygen distribution in the hepatic lobules. To deepen our knowledge about the dynamic alterations in liver metabolism in both physiological and pathophysiological conditions, real-time imaging of intracellular oxygen tension in situ is a promising method with the development of oxygen-sensitive probes.

We first compared the photophysical properties and oxygen sensitivity of Ir-1 and Ir-2 in solution and cultured cells. Next, we investigated the feasibility of these probes for the measurements of O_2 distribution in cell spheroids. Based on these results, we performed PLIM measurements of hepatic tissues of mice using Ir-1 and Ir-2 to reveal the probe performances.

### Results

#### Photophysical properties.

Emission brightness and O_2 sensitivity are basic characteristics required for an optical O_2 probe. The emission brightness of the probes under the same concentration can be evaluated by the product (εΦ_p) of the molar absorption coefficient (ε) at the excitation wavelength and the phosphorescence quantum yield (Φ_p) under the absence of oxygen, and the O_2 sensitivity by k_qτ_p, where k_q and τ_p are the bimolecular quenching rate constant by O_2 and the phosphorescence lifetime under the absence of oxygen. The first absorption band of Ir-1 appears at around 476 nm in MeCN (Fig. 1A), which originates from singlet metal-to-ligand-charge-transfer (1MLCT) transition. This band is slightly blue-shifted in Ir-2, giving a smaller ε value (2800 dm^3mol^-1 cm^-1) compared with that (5400 dm^3mol^-1 cm^-1) of Ir-1 at the excitation wavelength (488 nm) in our PLIM measurements. Ir-1 and Ir-2 exhibit red emission extending to the near infrared region and give vibrational structures with maximum wavelengths at 612 and 601 nm, respectively. Both complexes are highly emissive in MeCN (Φ_p=0.26 for Ir-1 and 0.28 for Ir-2). Judging from the ε values at 488 nm and the almost equal Φ_p, Ir-1 is superior to Ir-2 in brightness for excitation at 488 nm.

The O_2 sensitivity of a probe in solution can be evaluated experimentally from the ratio (τ_p/q) of the phosphorescence lifetimes under aerated and deaerated conditions. The obtained τ_p/q values for Ir-1 and Ir-2 in MeCN were 72 and 97, and the k_q values were calculated to be 7.59×10^4 mmHg^-1 s^-1 for Ir-1 and 8.22×10^4 mmHg^-1 s^-1 for Ir-2 (Table 1). Using the solubility of oxygen in MeCN at an oxygen partial pressure (pO_2) of 0.21 atm, the quenching rate constant can be converted from pressure units to concentration units. The converted k_q values (6.38×10^8 M^-1 s^-1 for Ir-1 and 6.91×10^8 M^-1 s^-1 for Ir-2) were close to the diffusion-controlled rate of bimolecular reactions in MeCN (52). The τ_p values were then converted to the τ_p/q values compared with metalloporphyrins and Ru(II) complexes (Table 1). The O_2 sensitivity is slightly higher in Ir-2 than Ir-1 owing to the longer intrinsic lifetime and larger k_q value of Ir-2. The intrinsic lifetimes (5~10 μs) of Ir-1 and Ir-2 are shorter than those (10~100 μs) of Pt-porphyrins, but longer than those of typical Ru(II) complexes used as O_2 probes (53), and thus the O_2 sensitivity of Ir-1 and Ir-2 is considered to be comparable with Pt-porphyrins and higher than Ru(II) complexes.

#### Characterization of intracellular O_2 probes.

To evaluate the cellular uptake efficiencies of Ir-1 and Ir-2, we compared the emission intensities of human colorectal adenocarcinoma (HT-29) cells incubated with these complexes (5 μM, incubation time: 2 h) using a plate reader. Here, we used their prototype complex BTP ((btp)_2Ir(acac); btp = benzothienylpyridine, acac = acetylacetone) (54) as a reference compound. The emission these complexes (5 μM, incubation time: 2 h) using a plate reader. Here, we used their prototype complex BTP ((btp)_2Ir(acac); btp = benzothienylpyridine, acac = acetylacetone) (54) as a reference compound. The emission

| Probe | λ_{max}^{abs} (nm) | τ_p (μs) | τ_p/τ_0 | Φ_p | k_q (10^9 mmHg^-1 s^-1) |
|-------|-------------------|----------|---------|------|------------------------|
| Ir-1  | 476               | 5.86     | 81.4    | 0.26 | 7.59                   |
| Ir-2  | 601               | 7.31     | 75.4    | 0.28 | 8.22                   |

Table 1. Photophysical parameters of Ir-1 and Ir-2 in MeCN at room temperature. τ_p/τ_0 and Φ_p denote the phosphorescence lifetime and quantum yield taken in degassed solutions, and τ_p and Φ_p denote the phosphorescence lifetime and quantum yield taken in aerated solutions.
the probes (500 nM or 1 µM) for 2 h prior to microscopic measurements. Therefore, under these experimental conditions, the effect of the probes on cell activity is considered to be sufficiently small.

Photostability is another important characteristic required for an intracellular O2 probe. We investigated the photostability of Ir-1 and Ir-2 in HT-29 cells by taking emission images of HT-29 cells under continuous 488 nm laser pulse irradiation with our PLIM system. The cells were incubated with Ir-1 or Ir-2 (1 µM) for 2 h under 21% or 2.5% O2 conditions, and PLIM images of the HT-29 cells were taken at every 50 scans until 550 scans (Fig. S4). During this irradiation time, the phosphorescence lifetimes of Ir-1 and Ir-2 were almost constant, although the phosphorescence intensity changed slightly by 488 nm light irradiation. Also, no significant change was observed in the cell morphology. In this study, the signals accumulated with 50 scans were usually averaged to obtain a single PLIM image. Therefore, Ir-1 and Ir-2 were found to have sufficient photostability for obtaining clear PLIM images.

Quantification of O2 levels. To quantify the oxygen levels in tissues based on lifetime measurements, we need a calibration curve that represents the relationship between lifetime and pO2. Since Ir-1 and Ir-2 are likely to accumulate in specific organelle membranes, the pO2 dependence of the phosphorescence lifetime in cells is expected to be different from that in solution. Therefore, we calibrated the lifetime by measuring PLIM images of HT-29 cells incubated with Ir-1 or Ir-2 under 21, 15, 10, 5, and 0% O2 conditions at 37 °C (Fig. 2A,B); we added 10 µM antimycin A (AntA) to the medium prior to the PLIM measurements to suppress the oxygen consumption by cellular respiration, and in the experiments under N2, saturated conditions we added Na2SO3 (500 mM) into the medium to remove oxygen remaining in the culture. The average lifetime of an entire image was plotted against the pO2 according to the Stern–Volmer equation, τp0/τp = 1 + kq pO2. A linear relationship was obtained for both complexes (Fig. 2C), and the kq values for Ir-1 and Ir-2 were derived to be 4.22 × 103 mmHg−1 s−1 and 5.36 × 104 mmHg−1 s−1 along with the τp0, 5.20 µs and 6.18 µs, respectively. These values were used to quantify oxygen levels in cell spheroids from phosphorescence lifetimes (τp).

Evaluation of O2 distribution in cell spheroids. In the past decade, three-dimensional (3D) spheroid systems have received much attention in fields such as drug discovery, cancer research, and toxicology. They provide a more physiologically-relevant environment and organ-like microarchitecture compared with conventional 2D cell cultures and better mimic the crucial tumor tissue properties and microenvironment. The excellent cell-permeable properties of Ir-1 and Ir-2 suggested their potential for in-depth staining of 3D cell spheroids and thus the potential of these probes for visualization of the O2 distribution within spheroids and living tissues.

So, we first acquired the PLIM images of HT-29 cell spheroids that were incubated with Ir-1 or Ir-2. Here, each probe (500 nM) was added to the medium after the spheroids were formed, and the spheroids were further incubated with the probe for 24 h prior to PLIM measurements. The bright-field images taken on different planes in the z direction (Fig. 3) shows that spheroids with a diameter of 150–200 µm were almost uniformly stained, including the core, by Ir-1 and Ir-2. We found from the Z-stack phosphorescence lifetime images of an HT-29 spheroid stained with Ir-1 (Fig. 3B) and Ir-2 (Fig. 3C) that the cells closer to the center of the spheroid gave longer lifetimes, i.e. lower O2 levels compared with those in the peripheral region. For quantitative evaluation of the O2 gradient in a spheroid, we investigated the line profiles of the average phosphorescence lifetime on the plane approximately 20 µm from the bottom (Fig. 3D). We also measured the lifetimes along the central area in the z-direction (Fig. 3E), and derived the pO2 distribution (Fig. 3D,E) based on the τp0 and kq obtained for HT-29 cultured cells. Both probes showed that the pO2 is reduced near the center and bottom surface of the spheroids, because the oxygen supply from the bottom direction is cut off by the bottom glass, and also the peripheral cells consume oxygen carried by diffusion from the culture medium. The degree of hypoxia in the core depended on the spheroid size (Fig. S3) and the respiratory activity as will be described below.

We next investigated whether Ir-1 and Ir-2 can image changes in the oxygen status of spheroids caused by metabolic stimulus. We used FCCP (carbonyl cyanide-(trifluoromethoxy)phenylhydrazone) and AntA to alter the oxygen status of HT-29 spheroids by stimulating metabolic processes; FCCP is an uncoupler of oxidative phosphorylation in mitochondria that disrupts ATP synthesis by transporting protons across the membrane and thus increases the O2 consumption rate. AntA is known to inhibit the mitochondrial electron transport chain from cytochrome b to cytochrome c1 and suppress O2 consumption. Metabolic stimulation of HT-29 cell spheroids by FCCP and AntA significantly changed the PLIM images (Fig. 4). The lifetime images of the spheroid plane ~30 µm from the bottom (Fig. 4B,C) demonstrated that upon addition of FCCP to the medium, the lifetimes of both probes quickly increased within ~10 min, especially in the core. The oxygen partial pressures calculated from the lifetimes based on the calibration in Fig. 2 indicate that the pO2 decreased sharply to less than 20 mmHg by increased O2 consumption, and then it slowly recovered. Conversely, the lifetimes of both probes rapidly dropped within 10 min upon metabolic stimulation using AntA (Fig. 4D,E), corresponding to the increase in pO2 by suppression of O2 consumption. The pO2 changes of spheroids reflected the metabolic effects of FCCP and AntA, demonstrating that Ir-1 and Ir-2 can be used to track the oxygen status of cell spheroids. These results confirmed the high penetrating ability of Ir-1 and Ir-2 into spheroids and their feasibility as tissue O2 probes.

In vivo O2 imaging of hepatic tissues. Since we confirmed that Ir-1 and Ir-2 can be efficiently taken up into spheroids and evaluate the oxygen status, we next attempted to visualize the oxygen gradient of hepatic lobules by in vivo PLIM measurements with Ir-1 or Ir-2 as a probe. Each probe (250 nmol in 25 µL dimethyl sulfoxide, diluted using PBS to 175 µL) was administered intravenously to anesthetized mice. Approximately 10 min after probe administration, the abdomen was opened to expose the liver, and the phosphorescence lifetime images of hepatic tissues (~10 µm from the surface of the liver) were measured around the central vein (CV) using different magnifications. Both probes gave clear PLIM images of hepatic tissues with cellular-level
Figure 2. Intensity (upper) and PLIM (lower) images of HT-29 cells stained with (A) Ir-1 and (B) Ir-2 under different $pO_2$ in an incubator. The average phosphorescence lifetimes are shown below each image. Cells were treated with Ant A (5–21% $O_2$) and Na$_2$SO$_3$ (0% $O_2$). Scale bar: 50 µm. (C) Stern–Volmer plots of $\tau_p^0/\tau_p$ as a function of $pO_2$ for Ir-1 (red) and Ir-2 (blue) partitioned into HT-29 cells under different $pO_2$ in an incubator. Error bars: S.D.
resolution (Fig. 5). Here, the areas that appear black are the regions where the emission intensity from the probe is extremely low, and these mainly correspond to the nucleus and sinusoid that have low probe uptake. It can be seen from the PLIM images measured with a 40 × objective lens that Ir-1 and Ir-2 are internalized into hepatocytes that line up along sinusoids. The arrangement of hepatocytes and lifetime distribution indicate that these images correspond to a hexagonal hepatic lobule (Fig. 1B) and that CV exists in the area with the longest lifetime (displayed in orange). The region surrounding the CV is considered to be the portal vein (PV) because it has a much shorter lifetime, i.e. much higher oxygen level. The direction of blood flow seen in the video image of hepatic tissues (Video S1) also indicates that the phosphorescence lifetime in the vicinity of CV is longer than that in the surrounding PV. PLIM images obtained with Ir-1 and Ir-2 show a similar lifetime gradient that increases from the PV to CV.

To quantify the O2 levels in specific locations in the lobules from the phosphorescence lifetimes, we performed calibration of the phosphorescence lifetimes using an AML 12 (alpha mouse liver 12) cell line that was established from mouse hepatocytes (Fig. S6). Using the $\tau_p^0$ and $k_q$ of Ir-1 and Ir-2 as determined in a monolayer of AML 12 cells, we evaluated the average $pO_2$ of the regions of interest (ROIs) that can be attributed to CV, PV, and the intermediate region (IR); the phosphorescence lifetime of Ir-1 gave reasonable O2 levels (24 ± 6.1 mmHg for CV, 32 ± 4.9 mmHg for IR, and 39 ± 4.2 mmHg for PV), whereas Ir-2 exhibited much lower O2 levels (3 ± 1.8 mmHg for CV, 5 ± 1.8 mmHg for IR, and 7 ± 2.7 mmHg for PV) (Table 2). These results indicated that PLIM images obtained with Ir-1 and Ir-2 clearly visualized the oxygen gradient from PV to CV. However, the O2 levels derived from the phosphorescence lifetimes differed between Ir-1 and Ir-2.

To confirm that Ir-1 and Ir-2 retain their spectral properties in hepatic tissues after systemic administration, we measured the emission spectra of the livers of living mice. Each probe molecule (250 nmol) was administered through the tail vein, and the emission images and spectra of livers were taken using a fluorescence microscope with an excitation filter of 450–500 nm and emission filter of > 532 nm (Fig. S7). The observed emission spectra with maxima at 619 nm for Ir-1 and 607 nm for Ir-2 were slightly red-shifted compared with those in MeCN,
but they were in good agreement with those in AML 12 cells, demonstrating that these probes maintain their spectral properties in hepatic tissue.

The reason why Ir-2 gave an abnormally low $O_2$ level seems to be related to the function of the liver to metabolize xenobiotics. So, to clarify how clearance of harmful metabolic byproducts and detoxification of xenobiotics affect hepatic $O_2$ levels, we investigated the detoxification of ammonia in the liver, which has the function of converting toxic ammonia to urea by the urea cycle or to glutamine by glutamine synthesis in hepatocytes. With increased metabolic activity of the liver, oxygen consumption should be enhanced to produce ATP. We intravenously administered $NH_4Cl$ (0.13 g/kg) to anesthetized mice at ~ 30 min after 250 nmol Ir-1 injection into the tail vein. PLIM images of hepatic lobules measured 10 min after the administration of $NH_4Cl$ exhibited a marked increase in the phosphorescence lifetime of Ir-1 (Fig. 6A,B). The average phosphorescence lifetimes

Figure 4. Imaging of the oxygen status of an HT-29 cell spheroid upon metabolic stimulation with FCCP or AntA. (A) Schematic representation of a spheroid in media containing FCCP or AntA. PLIM images were taken at 30 μm from the bottom after oil sealing. (B,C) Variation in PLIM images (left), and their average phosphorescence lifetime and $pO_2$ (right) of an HT-29 cell spheroid stained with (B) Ir-1 and (C) Ir-2 by metabolic stimulation with FCCP (1 μM) at 5 min and 6 min, respectively, and (D) Ir-1 and (E) Ir-2 by metabolic stimulation with AntA (1 μM) at 5 min. Scale bar: 100 μm.
(τₚ) and pO₂ of different ROIs in the pericentral region (ROI1) and periportal regions (ROI2 and ROI3) showed that the lifetimes decreased over time, and the O₂ levels recovered to the original levels in approximately 1 h.

To clarify the difference in in vivo probe performance between Ir-1 and Ir-2, we performed O₂ imaging experiments on liver tissues using another Ir(III) complex, (btp)₂Ir(acac-2OH) (Ir-3), which has a similar structure to Ir-1 (Fig. S8A). The τ₀ and kₑ of Ir-3 in AML 12 cells were determined to be 4.80 µs and 4.82 × 10³ mmHg⁻¹ s⁻¹. The PLIM image of an Ir-3-administered mouse exhibited a clear lifetime gradient and gave an average pO₂ of 16 ± 3.2 mmHg in CV, 21 ± 3.4 mmHg in IR, and 24 ± 3.8 mmHg in PV (Fig. S8C). The hepatic O₂ levels obtained with Ir-3 were close to those obtained with Ir-1, although Ir-3 gave somewhat smaller pO₂ values.

Figure 5. PLIM images of the hepatic surface of a mouse administered (A) Ir-1 and (B) Ir-2. The color bar indicates phosphorescence lifetime in µs. Left images: 10 × objective lens, center images: 20 × objective lens, right images: 40 × objective lens. Scale bar: 200 µm (left images), 100 µm (center images), and 50 µm (right images). CV: central vein, PV: portal vein. (C, D) Phosphorescence lifetime of Ir-1 and Ir-2 and pO₂ in hepatic lobules. Average of phosphorescence lifetime and pO₂ are shown. N = 23 ROIs for CV, 23 for IR (intermediate region between CV and PV), and 24 for PV in 5 mice administered Ir-1. N = 21 ROIs for CV, 23 for MR, and 27 for PV in 5 mice administered Ir-2. *p value < 0.01 by 2-tailed unpaired t test. Error bar: S.D.
Scientific Reports

O2, these are considered to be reasonable values given the oxygen levels than those reported for intravascular and 24 ± 6.1 mmHg, respectively. Although the hepatic oxygen levels obtained in our study are somewhat lower those in MeCN. One possible reason lies in the concentration quenching in organelle membranes. The τp complex bearing a cyclometalating 3-di(2-pyridyl)benzene-based moiety35, and click-assembled oxygen-sensing pO2 of 59.8 mmHg in portal vessels and 38.9 mmHg in central venules. Our (bovine serum albumin)-bound Pd-TCPP (Pd(II)-meso-tetra(4-carboxyphenyl)porphine) in hepatic tissues of steroids. Although the τp O2 cannot be ruled out, but its contribution appears to be relatively small because the τp partitioned into DMPC liposomes tended to shorten with increasing probe concentration in solution (Table S1),

Table 2. Phosphorescence lifetimes (τp) of Ir-1 and Ir-2 at around CV, IR, and PV in hepatic lobules and the pO2 values calculated from the lifetime. N = 23 ROIs for CV, 23 for IR, and 24 for PV in 5 mice administered Ir-1. N = 21 ROIs for CV, 23 for IR, and 27 for PV in 5 mice administered Ir-2.

| Area  | τp (μs) | pO2 (mmHg) |
|-------|---------|------------|
|       | Ir-1    | Ir-2       | Ir-1    | Ir-2             |
| CV    | 3.09 ± 0.31 | 5.78 ± 0.28 | 24 ± 6.1 | 3 ± 1.8          |
| IR    | 2.73 ± 0.18 | 5.45 ± 0.27 | 32 ± 4.9 | 5 ± 1.8          |
| PV    | 2.48 ± 0.14 | 5.10 ± 0.33 | 39 ± 4.2 | 7 ± 2.7          |

In contrast to the results in the liver tissue, Ir-1 and Ir-2 gave similar O2 tensions in the kidney (Fig. S9). Intravenously administered probes rapidly migrated from the vasculature into the proximal tubule cells and were hardly excreted in the urinary space during the observation period. The average O2 tension of tubular cells in each image was derived using calibration lines obtained for human kidney 2 (HK-2) cells to be 45 ± 9.8 mmHg for Ir-1 and 36 ± 4.3 mmHg for Ir-2 (Fig. S10).

Discussion

We first verified that Ir-1 and Ir-2 exhibit desired photophysical properties as O2 probes in cells and cell spheroids. Although the τp values (5–10 μs) of Ir-1 and Ir-2 are much smaller than those of Pt(II)- and Pd(II)-porphyrins, their oxygen sensitivity is sufficiently high due to their large kp values. In addition, the moderately long lifetimes of the Ir(III) complexes have the advantage of increasing the counting efficiency in PLIM measurements and thus shortening the image acquisition time.

One key issue in applying small-molecule intracellular oxygen probes for in vivo O2 measurements is the method of lifetime calibration. We quantified the tissue pO2 based on the kp and τp determined using a cell line that is the same as or close to the cell type in the target organ in which the probe is accumulated. The O2 quenching rate constants in HT-29 and AML 12 cells were reduced by an order of magnitude compared with those in solution (see Table 1). Since lipophilic Ir-1 and Ir-2 are likely to accumulate in organelle membranes after they pass through the plasma membranes of living cells, the much smaller kp values in cells can be attributed, at least in part, to the decrease in the diffusion rates of O2 and Ir(III) complexes in organelle membranes. Further reductions in kp values can be caused by the binding of excited probe molecules to proteins in the organelle membrane in which they are incorporated. The τp values of Ir-1 and Ir-2 in cells were decreased by 10–15% compared with those in MeCN. One possible reason lies in the concentration quenching in organelle membranes. The τp of Ir-1 partitioned into DMPC liposomes tended to shorten with increasing probe concentration in solution (Table S1), suggesting that concentration quenching may have occurred. Cross-sensitivity to endogenous species other than O2 cannot be ruled out, but its contribution appears to be relatively small because the τp values of Ir-1 and Ir-2 in cells are close to those in MeCN. It was also confirmed that phosphorescence lifetime is almost independent of pH and the presence of glutathione under physiological conditions (Tables S2 and S3).

One of the major advantages of Ir-1 and Ir-2 is their extremely high cellular uptake efficiencies. The O2 imaging experiments using HT-29 cell spheroids (Fig. 3) demonstrated that Ir-1 and Ir-2 are efficiently internalized into the spheroids with a diameter of ~200 μm. The penetrative abilities of some molecular O2 probes into cell spheroids reported so far are compared in Table S4. These probes include glucose conjugates of Pt(II)-mesotetrasik-(pentfluorophenyl)porphyrin20, Pt(II)-5, 10, 15, 20-tetrakis-(4-carboxyphenyl)porphyrin36, a Pt(II) complex bearing a cyclometalating 3-di(2-pyridyl)benzene-based moiety35, and click-assembled oxygen-sensing nanoconjugates with Pd(II)tetracarboxytetraphenylporphyrin as a phosphorescent core24. Comparing each probe for its concentration in culture medium and the incubation time required for observation of clear luminescence images, it can be seen that Ir-1 and Ir-2 have excellent cell permeability. Furthermore, PLIM measurements with these probes made it possible to track oxygen levels and distributions in spheroids. These properties are very useful for investigating the oxygen status in tissues.

The oxygen gradient in the hepatic lobule has been demonstrated by applying the hypoxia marker 2-nitroimidazole to liver tissue sections52. High-resolution visualization of hepatic oxygen distribution in vivo has been performed by Paxian et al.48 by intravital emission microscopy using [Ru(phen)3]2+ (Tris(1,10-phenanthroline) ruthenium(II)) as an O2 probe. They observed a continuous increase in emission intensity of [Ru(phen)3]2+ from for its concentration in culture medium and the incubation time required for observation of clear luminescence images, so that intensity-based measurements cannot assess the precise pO2 and O2 gradient. As for hepatic tissue oxygen levels, Kietzmann and Jungermann54,55 have reported the pO2 to be 60–65 mmHg in the periportal blood and 30–35 mmHg in the perivenous blood, and they suggested the pO2 level to be approximately 15 mmHg lower in periportal and perivenous cells. Tsukada and Suematsu56 have investigated the average pO2 of hepatocytes near PV and CV were 39 ± 4.2 mmHg and 24 ± 6.1 mmHg, respectively. Although the hepatic oxygen levels obtained in our study are somewhat lower than those reported for intravascular pO2, these are considered to be reasonable values given the oxygen levels to which hepatocytes are exposed. To the best of our knowledge, the PLIM images shown in Fig. 5 are the first to visualize the oxygen concentration gradient in hepatic lobules in vivo using an intracellular O2 probe.
Figure 6. (A,B) Variation of PLIM images of hepatic lobules after administration of NH$_4$Cl following Ir-I injection into the tail vein of a mouse. Scale bar: 200 µm. (C,D) Variation of average phosphorescence lifetime and $p$O$_2$ of ROIs in the pericentral region (ROI1) and periportal regions (ROI2 and ROI3) shown in B. Error bar: S.D.
Imaging the spatiotemporal $pO_2$ changes within the liver microarchitecture provides useful information about the oxygen response and dynamics of the liver at the cellular level during physiological stimulation. Our PLIM measurements with Ir-1 showed that intravenous administration of NH$_4$Cl resulted in a rapid decrease in hepatic tissue $O_2$ levels, and its recovery rate depended on the position in the lobule (Fig. 6). Hepatocytes are known to display considerable functional differences depending on their position along the porto-central axis of the liver lobule, and ammonia detoxification in the liver occurs by two pathways: consumption by urea synthesis in perivenous area and removal by glutamine synthesis in periportal region. Since these processes require $O_2$ consumption, ammonia stimulation may produce hypoxic regions in the hepatic lobules depending on the $O_2$ consumption rate of hepatocytes.

In HT-29 cell spheroids, Ir-1 and Ir-2 showed almost the same $O_2$ levels and distribution, whereas in hepatic tissues of living mice, Ir-2 gave much lower oxygen levels: 7 ± 2.7 mmHg for the periportal region and 3 ± 1.8 mmHg for the perivenous region (Table 2). In kidneys, on the other hand, both probes accumulated in tubular cells and showed similar oxygen levels: 45 ± 9.8 mmHg for Ir-1 and 36 ± 4.3 mmHg for Ir-2 (Fig. S9). These results suggest that the administration of Ir-2 to mice causes hypoxia in hepatic tissues due to some specific metabolic processes in the liver. Since spectral measurements (Fig. S7) revealed that the luminescent properties of Ir-1 and Ir-2 were maintained in liver tissue, it is likely that Ir-2 is excercised in the bile through solubilization processes that do not affect the luminescent properties such as glucuronidation at the OH group. The luminescence spectrum of the extract from bile 6 h after probe administration showed that Ir-1 was excercised through the bile duct along with bile (Fig. S11). Considering that Ir-3, which has the same acac moiety as Ir-1, gave similar liver tissue $O_2$ levels as Ir-1, the lipophilic Tris-ligand structure of Ir-2 may be related to its metabolism in the liver.

Conclusions
Small-molecule $O_2$ probes, Ir-1 and Ir-2, were efficiently taken up into cells when added to the culture media of monolayer cells and cell spheroids, and this enabled reversible $pO_2$ measurements in the cells. PLIM measurements of mouse livers following intravenous administration of Ir-1 or Ir-2 allowed high-resolution $O_2$ imaging of hepatic tissues, exhibiting an $O_2$ gradient from the pericentral to periportal regions in hepatic lobules. The phosphorescence lifetime of Ir-1 gave reasonable hepatic $O_2$ levels after calibrating the lifetime using cultured AML 12 cells. Furthermore, Ir-1 allowed visualization of the $O_2$ changes in hepatic tissues stimulated by ammonia. However, Ir-2 gave a much lower $pO_2$ compared with Ir-1, which may be due to the toxicity of Ir-2, which promoted detoxification in the liver. These results reveal that Ir(III) complexes allow imaging of spatiotemporal changes in oxygen levels within the tissue microarchitecture in vivo, but some complexes may influence oxygen consumption in the liver when used for oxygen imaging of hepatic tissues.

Methods
Materials. The Ir(III) complexes used in this study were synthesized and characterized according to the methods described in the Supplementary Information.

Acetominitrile (MeCN; Kanto Chemical, spectroscopic grade) and NH$_4$Cl (Kanto Chemical, special grade) were used as received. Anti Amycin A from Streptomyces sp. (Ant A) and carboxyl cyanide 4-(trifluoromethoxy) phenylhydrozone (FCCP, 98%) were purchased from Sigma-Aldrich. Cell Counting Kit-8 (CCK-8) was purchased from Dojindo Laboratories.

$^1$H-NMR spectra were recorded with a JEOL JNM-ECS400 (400 MHz) spectrometer in DMSO-$d_6$. $^1$H-NMR chemical shifts were referenced to tetramethylsilane. The apparent resonance multiplicity was described as s (singlet), d (doublet), dd (double doublet), t (triplet), and m (multiplet). ESI–MS spectra were recorded on an Applied Biosystems API 2000 mass spectrometer.

Photophysical properties in solution. Absorption spectra were recorded on a UV/Vis spectrophotometer (Ubest-V550, JASCO). Emission spectra were measured with a system consisting of a monochromatized Xe arc lamp, a sample holder, and a photonic multichannel analyzer (C10027-01, Hamamatsu Photonics), and were fully corrected for spectral sensitivity between 200–950 nm. Phosphorescence lifetimes in solution were measured with a lifetime measurement system (Quantaurus-Tau C11367G, Hamamatsu Photonics) based on the time-correlated single photon counting (TCSPC) method. Phosphorescence quantum yield was measured with an absolute photoluminescence quantum yield measurement system (C9920-02, Hamamatsu Photonics) that consisted of a Xe arc lamp, a monochromator, an integrating sphere, a multichannel detector, and a personal computer.

Cell and cell spheroid cultures. Human colorectal adenocarcinoma (HT-29) cells and HeLa cells were purchased from the American Type Culture Collection (ATCC). Alpha mouse liver 12 cells (AML12 cells) were kindly provided by Prof. T. Inagaki of the Laboratory of Epigenetics and Metabolism, IMCR, Gunma University. Human kidney 2 cells (HK-2 cells) were kindly provided by Prof. M. Nangaku of the Graduate School of Medicine, the University of Tokyo. HT-29 cells were incubated in McCoy’s 5A medium (Gibco) containing 10% fetal bovine serum (FBS), penicillin (50 units/mL), and streptomycin (50 µg/mL). AML12 cells and HK-2 cells were incubated at 37 °C in a mixture of Dulbecco’s Modified Eagle’s Medium (DMEM) and Ham’s F-12 Nutrient Mixture containing 10% FBS, penicillin (50 units/mL), and streptomycin (50 µg/mL). All cells were grown in a humidified 5% CO$_2$ enriched atmosphere at 37 °C. In PLIM imaging, all cells were cultured on glass-based dishes, and the medium was changed to McCoy’s 5A or DMEM/F-12 without phenol red and FBS prior to the measurements.

HT-29 cell spheroids were prepared using a low attachment 96-well NanoCulture Plate (NCP; NCP-LH96, ORGANOGENICS) with a micro-honeycomb pattern on the film that comprises the bottom of the plate. McCoy’s
5A medium (100 μL) containing 10% FBS, penicillin (50 units/mL), and streptomycin (50 μg/mL) was added to each well in the NCP, and the NCP was centrifuged at 10 × 100 rpm for 5 min to remove the bubbles in the media in the wells. Then HT-29 cells (1.0 ×10^6 cells/mL) suspended in 100 μL McCoy’s 5A medium were added to each well in the NCP. The cells were incubated in the NCP at 37 °C, 5% CO_2 for 4 days to form the HT-29 cell spheroids.

**PLIM imaging of cultured cells.** PLIM images were measured using an inverted fluorescence microscope (IX73, Olympus) equipped with a picosecond diode laser (BDL-SMC, Becker & Hickl; wavelength: 488 nm, pulse width: 40–90 ps, repetition rate: 50 MHz) and a confocal scanning system (DCS-120, Becker & Hickl). A 4× objective (UPlanSApo; NA 0.16, Olympus), 10× objective (UPlanSApo; NA 0.40, Olympus), 20× objective (UPlanFLN; NA 0.70, Olympus), 40× oil objective (UPlanFLN; NA 1.30, Olympus), or 100× oil objective (UPlanSApo; NA 1.40, Olympus) were used for PLIM imaging. The emission, as guided through a 1 mm pin-hole, was collected through a longpass filter and detected with a hybrid detector module (HPM-100–40, Becker & Hickl). Time-resolved emission measurements were performed with a time-correlated single photon counting (TCSPC) system (Simple-Tau-150-DX, Becker & Hickl). The emission signals were acquired at a resolution of 128×128 pixels, and the decay curve in each pixel was fitted to single- or double-exponential decay function. In the case of double-exponential decay, the amplitude average lifetime was used for Stern–Volmer analyses. PLIM images were analyzed with SPImage data analysis software (Becker and Hickl). In PLIM imaging of cell spheroids, HT-29 cell spheroids were incubated in the McCoy’s 5A medium containing a final concentration of 500 nMIr-1 or Ir-2 for 24 h at 37 °C, 5% CO_2 prior to the measurements.

**In vivo PLIM measurements.** All protocols for animal experiments were approved by the Ethical Committee on Animal Experiments of Gunma University (18-018), and all animal experiments were conducted in accordance with the institutional guidelines. In vivo PLIM measurements were carried out using six- to eight-week-old Balb/c male mice (CLEA Japan). After general anesthesia, probe solution (200 μL) containing 250 nmol Ir(III) complex in 10% DMSO/saline was injected into the tail vein, and the liver or kidney was exposed for ~10 min before PLIM experiments.

**Calibration of the phosphorescence lifetime.** The oxygen partial pressure in cells was obtained from the phosphorescence lifetime based on a calibration using cultured cells. HT-29, AML12, or HK-2 cells were seeded on glass-based dishes and stained with Ir-1 or Ir-2 (500 nM, 2 h). PLIM images were acquired for the stained cells at 21%, 15%, 10%, 5%, and 0% O_2 in an incubator at 37 °C. Here, in the 5–21% O_2 experiments, cells were incubated with Ant A (10 μM, >0.5 h) to block cellular respiration, and in the 0% O_2 experiments, fresh medium containing Na_2SO_4 (500 mM) was used to remove existing O_2 in the medium. The average phosphorescence lifetimes of the PLIM images taken under different pO_2 were plotted according to the Stern–Volmer equation, \( \tau_p^0 = 1 + \frac{kq}{\tau_p}pO_2 \), to determine the bimolecular quenching rate constant, \( kq \). The pO_2 of cells, spheroids, and tissues was obtained from the phosphorescence lifetime (\( \tau_p \)) using the \( kq \) and \( \tau_p^0 \).

**Subcellular localization of probes.** HeLa cells were cultured in DMEM (Gibco) with 10% FBS, penicillin (50 units/mL), and streptomycin (50 μg/mL), and were grown under a 5% CO_2 atmosphere at 37 °C. The cells were stained in glass bottom imaging dishes (Greiner) and allowed to adhere for 24 h, incubated with Ir-1 (500 nM) or Ir-2 (1 μM) for 2 h, washed 3 times with DMEM, and then the medium was replaced with DMEM (FluoroBrite, Gibco) without FBS. Luminescence microscopic images were obtained with an inverted microscope (IX71, Olympus) equipped with an electron multiplying CCD camera (Evolve 512, PHOTOMETRICS) driven by MetaMorph software.

**Evaluation of cytotoxicity using CCK-8 assay.** HT-29 cells (2.5 ×10^4 cells/well) were seeded into a 96-well flat bottom plate (Greiner) for 48 h. The cells were incubated with various concentrations of each probe for 24 h at 37 °C under a 5% CO_2 atmosphere. The medium was removed, and the cells were washed gently with McCoy’s 5A medium without phenol red. Cell Counting Kit-8 reagent (CCK-8, Dojindo) was added to each well, and incubation was continued for 1 h. The absorbance at 450 nm of each well referenced at 650 nm was recorded using a microplate reader (Infinite 200 PRO, Tecan). Cell viability (% of control) was calculated as
\[
\frac{A_{\text{sample}} - A_{\text{blank}}}{A_{\text{control}} - A_{\text{blank}}} \times 100
\]
where \( A_{\text{sample}} \) is the absorbance of cells exposed to the probe, \( A_{\text{control}} \) is the absorbance of cells without probe, and \( A_{\text{blank}} \) is the absorbance of wells without cells.

**Mitochondrial membrane potential.** HT-29 cells (3.0 ×10^4 cells/well) were seeded into a 96-well flat bottom plate (Greiner) for 48 h. The cells were stained with Ir-1 or Ir-2 (1 μM, 2 h). The medium was removed, and the cells were washed gently with McCoy’s 5A medium without phenol red. 5,5′,6,6′-tetrachloro-1,1′-3,3′-tetraethyl-benzimidazolylcarbocyanine iodide (JC-1, Thermo Fisher Scientific) (5 μM) was added to each well, and incubation was continued for 30 min. The medium was removed, and the cells were washed gently with McCoy’s 5A medium without phenol red. The emission intensities of JC-1 were measured using a microplate reader (Infinite 200 PRO, Tecan) equipped with a gas control module (GCM, Tecan). The excitation wavelength was 488 nm, and the monitor wavelength was 595 nm for Red emission (aggregate) and 540 nm for green emission (monomer).
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Author contributions
T.Y., K.M. and S.T. designed the experiment. K.M., A.K. and S.S. carried out the experiments. S.T., N.G. and K.M. wrote the manuscript. All authors reviewed the manuscript.

Competing interests
The authors declare no competing interests.

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