# Chemical Science



### **EDGE ARTICLE**

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2021, 12, 13888

dll publication charges for this article have been paid for by the Royal Society of Chemistry

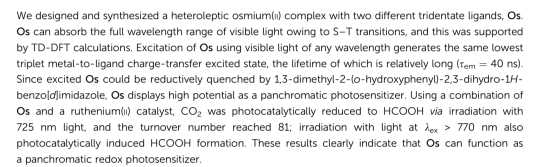
Received 23rd July 2021 Accepted 28th September 2021

DOI: 10.1039/d1sc04045f

rsc.li/chemical-science

# Development of a panchromatic photosensitizer and its application to photocatalytic CO<sub>2</sub> reduction†

Mari Irikura, Yusuke Tamaki\* and Osamu Ishitani (1)\*



#### Introduction

In recent years, various photocatalytic reactions toward artificial photosynthesis for CO<sub>2</sub> reduction,<sup>1,2</sup> H<sub>2</sub> evolution,<sup>3</sup> and water oxidation<sup>4</sup> using metal complexes have attracted attention. Because these reactions are multi-electron processes, two functions are required: a redox photosensitizer that absorbs visible light and initiates electron transfer from its excited state and a catalyst that accepts electrons or holes and activates substrates through multi-electron processes. For effective utilization of solar light, absorption of visible light over a wide range of wavelengths by redox photosensitizers is highly important. For example, light in the 280–550 nm wavelength range covers only 14% of total solar irradiance in AM1.5G conditions, while 280–800 nm light covers 40%.<sup>5</sup> The use of a panchromatic redox photosensitizer that can absorb the full wavelength range of visible light is promising for increasing the utilization efficiency of solar energy.

Photoredox catalysts, another name for redox photosensitizers, also play critical roles in many photochemical organic reactions. In these photocatalytic reactions, light absorption by co-existing components, such as a catalyst and/or substrate, frequently causes side reactions and the inner-filter effect,

absorption by the photosensitizer. For this reason, the development of redox photosensitizers that can absorb longer wavelength light is also important.

Ruthenium(II) tris-diimine complexes,  $[Ru(N^N)_3]^{2+}$   $(N^N = 1)^{2+}$ 

which lowers the efficiency of photocatalysis by inhibiting light

diimine ligand), are the most frequently used redox photosensitizers because they absorb visible light, owing to singlet metalto-ligand charge-transfer (1MLCT) transitions, and their lowest triplet MLCT (3MLCT) excited states that are rapidly and quantitatively produced by the intersystem crossing from the <sup>1</sup>MLCT states have long lifetimes ranging from several hundred ns to several µs.7,8 They also have appropriate redox potentials for various reactions. However,  $[Ru(N^N)_3]^{2+}$  as photosensitizers have the disadvantage of limited visible light absorption, typically  $[Ru(bpy)_3]^{2+}$  (bpy = 2,2'-bipyridine) can absorb only the light at  $\lambda_{abs}$  < 550 nm; therefore, many of the reported  $[Ru(N^N)_3]^{2^+}$ -type redox photosensitizers cannot utilize lowerenergy visible light. A simple strategy to extend the wavelength range of light absorption is to decrease the energy gap between the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO); however, this usually shortens the lifetime of the excited state owing to the energy gap law and decreases the reducing/oxidizing power of the complex. For this reason, the reported ruthenium(II) complexes with panchromatic and near-infrared absorption, which are employed in dye-sensitized solar cells, 9,10 cannot be utilized as redox photosensitizers for photocatalytic reactions. One example of a ruthenium(II) photosensitizer with longerwavelength absorption is a trinuclear ruthenium(II) complex bridged by 1,4,5,8,9,12-hexaazatriphenylene (HAT), [{(dmb)<sub>2</sub>- $Ru_{3}(\mu-HAT)^{6+}$  (dmb = 4,4'-dimethyl-2,2'-bipyridine), which

Department of Chemistry, School of Science, Tokyo Institute of Technology, O-okayama 2-12-1-NE-1, Meguro-ku, Tokyo, 152-8550, Japan. E-mail: tamaki@chem.titech.ac.jp; ishitani@chem.titech.ac.jp

† Electronic supplementary information (ESI) available: DFT-calculated characteristics of **Os**; emission quenching experiment; Franck-Condon analysis of emission spectrum with vibronic structure of **Os**; mirror image of S-T absorption and phosphorescence of **Os**; cyclic voltammogram of BI(OH)H; quantum yield for HCOOH production; visible absorption spectra of **Os**, **Ru(CO)** and BI(OH)H. See DOI: 10.1039/d1sc04045f

has been used as a redox photosensitizer to drive H<sub>2</sub> evolution.<sup>11</sup> In this case, the ion-pair adducts of  $[\{(dmb)_2Ru\}_3(\mu-HAT)]^{6+}$  and ascorbate anions have been reported to afford static reductive quenching and increased reducing power, which compensated for the disadvantage of a smaller HOMO-LUMO gap. However, the reducing power of this adduct is still insufficient for driving CO<sub>2</sub> reduction, which requires stronger reducing power than does H2 evolution.

Another strategy for achieving longer-wavelength absorption is to utilize the direct transition from the ground state to the triplet excited state (S-T transition). Although this is usually a spin-flip-forbidden transition, many osmium(II) complexes give S-T absorption bands owing to strong spin-orbit coupling due to the heavy atom effect of osmium. For example,  $[Os(tpy)_2]^{2+}$  (tpy = 2,2':6',2"-terpyridine) has given an S-T absorption band at 600-750 nm and been employed as a photosensitizer for organic synthesis12 and H2 evolution.13 We also previously reported that  $[Os(5dmb)_2(dmb)]^{2+}$  (5dmb = 5,5'dimethyl-2,2'-bipyridine) functions as a photosensitizer and catalyzes CO<sub>2</sub> reduction in combination with a rhenium(1) catalyst under red light irradiation ( $\lambda_{ex} > 620$  nm).<sup>14</sup> However, these osmium(II) complexes do not absorb the full wavelength range of visible light up to 800 nm (Fig. 1).

Therefore, in this study, we developed a redox photosensitizer that displays panchromatic absorption. A heteroleptic osmium(II) complex with two different tridentate ligands,  $[(mbip)Os(mtpy)]^{2+}$  (Os in Chart 1: mbip = bis(N-methylbenzimidazolyl)pyridine; mtpy = 4'-methyl-2,2':6',2''-terpyridine), was designed and synthesized. The stronger  $\sigma$ -donation of the methylbenzimidazole units should induce longerwavelength absorption. Although a similar complex, [(mbip) Os(tpy)]<sup>2+</sup>, has been reported, its photochemical and photosensitizing properties have not been investigated.15 In this study, the photophysical, photochemical, and photosensitizing properties of Os were investigated, and Os was successfully applied to the photocatalytic reduction of CO2 as the first panchromatic redox photosensitizer in combination with a ruthenium(II) catalyst, Ru(CO) (Chart 1).

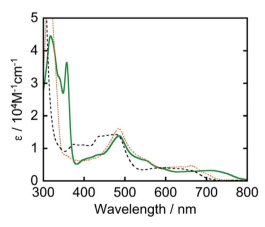


Fig. 1 UV-vis absorption spectrum of Os (green solid line) measured in DMA along with those of [Os(5dmb)<sub>2</sub>(dmb)]<sup>2+</sup> in N,N-dimethylformamide (black broken line) and [Os(mtpy)<sub>2</sub>]<sup>2+</sup> in DMA (orange dashed line).

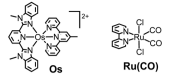
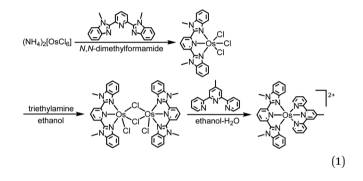


Chart 1 Structures and abbreviations of the complexes used. Counter anions were PF<sub>6</sub>

#### Results and discussion

The heteroleptic osmium(II) complex with two tridentate ligands was synthesized through the three steps shown in eqn (1). [(mbip)OsCl<sub>3</sub>] was dimerized into [Os(mbip)(μ-Cl)Cl]<sub>2</sub> to remove the byproduct of [Os(mbip)<sub>2</sub>]<sup>2+</sup> by exploiting the insolubility of [Os(mbip)(μ-Cl)Cl]<sub>2</sub> in ethanol. The target complex, Os, was isolated using ion-exchange column chromatography.



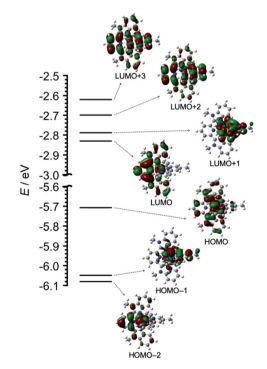


Fig. 2 DFT-calculated energy diagram and isodensity plots of frontier molecular orbitals of Os. The left portions of the structures are mbip. All orbitals were computed at an isovalue of 0.02.

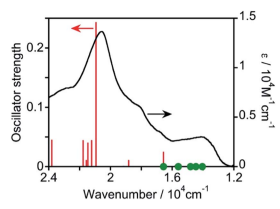


Fig. 3 Calculated oscillator strengths of S-S transitions (red lines) and UV-vis absorption spectrum of **Os** measured in DMA (black line). Green filled circles  $(\bullet)$  indicate the calculated energies of S-T transitions.

The ultraviolet-visible (UV-vis) absorption spectrum of **Os** measured in *N,N*-dimethylacetamide (DMA) is shown in Fig. 1. **Os** gave longer-wavelength absorption bands at 620–800 nm ( $\lambda_{\rm max}=709$  nm,  $\varepsilon=3.20\times10^3$  M $^{-1}$  cm $^{-1}$ ) attributable to S-T transitions to  $^3$ MLCT excited states as well as bands attributed to  $\pi-\pi^*$  transitions of the tridentate ligands at 300–360 nm and  $^1$ MLCT transitions at  $\lambda_{\rm max}=487$  nm ( $\varepsilon=1.38\times10^4$  M $^{-1}$  cm $^{-1}$ ). The S-T transition bands should be observed because of the heavy atom effect of osmium, which enhances spin–orbit coupling. The spectra of other osmium(II) complexes that have been used as photosensitizers are also shown in Fig. 1. It should be noted that only **Os** can absorb full-wavelength range of visible light with a relatively high molar absorptivity.

The geometry, molecular orbitals, and absorption spectrum of  $\mathbf{Os}$  were analyzed using density functional theory/time-dependent density functional theory (DFT/TD-DFT) calculations using Gaussian 16. The geometry optimization of  $\mathbf{Os}$  was performed in DMA using the B3LYP exchange-correlation functional and LanL2DZ basis set for osmium and 6-311++G(d,p) for the ligands. Tridentate ligands mbip and mtpy were coordinated on osmium perpendicular to each other.

Fig. 2 shows the frontier molecular orbitals of **Os**, and Table S1† summarizes their characteristics. HOMO-2, HOMO-1, and HOMO received major contributions from the osmium(II) center and minor contributions from the ligands. The energy levels of the LUMO and LUMO+1 were very close ( $\Delta E = 0.04$  eV). The LUMO and LUMO+1 received major contributions from mbip and mtpy, respectively. LUMO+2 and LUMO+3 were distributed on both mbip and mtpy.

Then, TD-DFT calculations were performed based on the geometry optimized using DFT calculations. Fig. 3 displays the calculated oscillator strengths of the excitations and the observed spectrum, and Table 1 summarizes the characteristics of the excitations. The observed  ${}^{1}MLCT$  absorption band at  $\lambda_{max}$ = 487 nm was consistent with the singlet-to-singlet (S-S) transition at 20 940 cm $^{-1}$  (= 478 nm, #17 in Table 1) with a high oscillator strength of 0.242. This excitation received a major contribution from the HOMO-1  $\rightarrow$  LUMO+1 transition (57%) and a partial contribution from the HOMO $-2 \rightarrow LUMO$  (24%) and HOMO → LUMO+2 (16%) transitions, indicating that the excitation is predominantly the MLCT from osmium(II) to the mtpy ligand. Because TD-DFT calculations using Gaussian 16 do not consider spin-orbit couplings, the oscillator strength of every S-T transition cannot be calculated. A spin-forbidden S-T transition gains strength by borrowing the intensity of the oscillator strength of the spin-allowed S-S transition  $(f_s)$ . Considering the coupling of S-T transitions with S-S transitions, effective coupling should occur only between S-T and S-S transitions involving different d orbitals and common  $\pi^*$ orbitals.17-20 As shown in Table 1, therefore, the S-T transitions of #1 and #3 couple with the S-S transition of #17, which has high oscillator strength ( $f_s = 0.242$ ), although the energy gaps are relatively large ( $\Delta E_{\rm S-T} = 6-7 \times 10^3 \ {\rm cm}^{-1}$ ). The S-T transitions of #2 and #4 strongly couple with the S-S transition of #6, which has low oscillator strength ( $f_s = 0.024$ ) because of the smaller energy gap ( $\Delta E_{S-T} = 1-2 \times 10^3 \text{ cm}^{-1}$ ). The S-T transition of #7 can couple weakly with the S-S transitions of #18 ( $f_s$  = 0.044), #21 ( $f_s = 0.039$ ), and #23 ( $f_s = 0.044$ ). Thus, the S-T transitions of #1 to #4 should gain oscillator strength by

Table 1 Excitation characteristics of Os<sup>a</sup>

#	Energy/cm <sup>-1</sup>	Oscillator strength	Multiplicity	Major transitions		
1	14 054		Triplet	$H \rightarrow L (98\%)$		
2	14 470		•	$H-2 \rightarrow L (82\%)$	$H-1 \rightarrow L+1 (15\%)$	
3	14 819			$H \rightarrow L+1 (96\%)$		
4	15 613			$H-2 \rightarrow L (15\%)$	$H-1 \rightarrow L+1 (79\%)$	
7	16 579			$H \rightarrow L+2 (30\%)$	$H \rightarrow L+3 (60\%)$	
6	16 564	0.024	Singlet	H → L+1 (98%)		
12	18 820	0.010	Ü	$H-1 \rightarrow L+1 (11\%)$	$H \rightarrow L+2 (81\%)$	
17	20 940	0.242		$H-2 \rightarrow L (24\%)$	$H-1 \rightarrow L+1 (57\%)$	$H \rightarrow L+2 (16\%)$
18	21 219	0.044		$H-1 \rightarrow L+2 (92\%)$	,	,
21	21 466	0.039		$H-2 \rightarrow L+2 (99\%)$		
23	21 778	0.044		$H-2 \rightarrow L+3 (98\%)$		

<sup>&</sup>lt;sup>a</sup> Calculated using TD-DFT at the B3LYP/LanL2DZ (Os)/6-311++G(d,p) level of theory. The HOMO and LUMO are represented by H and L, respectively.

**Edge Article** 

0.5 0.3 0.1 -1.4 -1.6 -1.8 -2
Potential / V vs. Ag/AgNO<sub>3</sub>

Fig. 4 Cyclic voltammograms of Os (0.5 mM) in an Ar-saturated DMA solution containing  $Et_4NBF_4$  (0.1 M) as a supporting electrolyte with a Ag/AgNO<sub>3</sub> (10 mM) reference electrode. Scan rate was 0.2 V s<sup>-1</sup>.

coupling with spin-allowed S–S transitions. They have lower energy than the lowest-energy S–S transition (#6) and their energy is consistent with the observed absorption band in the lower-energy region (1.3–1.6  $\times$  10 $^4$  cm $^{-1}$ ). Therefore, these S–T transitions should contribute to absorption in the longer-wavelength region.

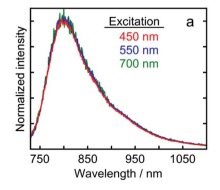
Fig. 4 shows the cyclic voltammogram of **Os** measured in an Ar-saturated DMA solution. Three reversible redox couples were observed at +0.35, -1.57, and -1.89 V  $\nu s$ . Ag/AgNO<sub>3</sub>. The oxidation wave at  $E_{1/2}^{\rm ox}=+0.35$  V was attributed to the one-electron oxidation of osmium(II), and the reduction waves at  $E_{1/2}^{\rm red}=-1.57$  and -1.89 V were attributed to the subsequent one-electron reduction of the tridentate ligands. These results indicate that the one-electron-reduced species (OERS) of **Os** has relatively strong reducing power and is stable at least on the time scale of cyclic voltammetry measurement at a scan rate of 0.2 V s<sup>-1</sup>.

Os exhibited phosphorescence at  $\lambda_{max}=795$  nm in an Arsaturated DMA solution upon excitation with visible light ( $\lambda_{ex}=456$  nm). The emission quantum yield was  $\Phi_{em}=0.3\%$ . The emission lifetime was determined to be  $\tau_{em}=40$  ns from single-exponential fitting of the transient emission (Fig. S1†). Although longer-wavelength absorption and a longer lifetime in the excited state are commonly in a trade-off relationship

because of the energy gap law which gives faster non-radiative decay at smaller energy gap, Os displayed panchromatic absorption and a relatively long lifetime, enabling initiation of the electron transfer process. These results indicate that **Os** has essential photophysical properties as a redox photosensitizer. Fig. 5 displays the emission spectra following excitation with visible light at three different wavelengths and the excitation spectrum derived from emission intensity. The emission spectra of **Os** following the excitation of the <sup>1</sup>MLCT or <sup>3</sup>MLCT absorption band at  $\lambda_{ex} = 450$ , 550, and 700 nm had very similar shapes (Fig. 5a). The shape of the excitation spectrum derived from the emission intensity at  $\lambda_{\text{max}} = 795$  nm is quite similar to that of the absorption spectrum of Os (Fig. 5b). These results clearly indicate that both the <sup>1</sup>MLCT and <sup>3</sup>MLCT excited states of **Os** relax to the same lowest <sup>3</sup>MLCT excited state very rapidly, and this excited state deactivates to the ground state through radiative and non-radiative decay processes. Therefore, visiblelight irradiation of **Os** at any wavelength generates the same lowest  ${}^{3}\text{MLCT}$  excited state with  $\tau_{em}=40$  ns.

Fig. S2† shows the emission spectra of Os in CO<sub>2</sub>-saturated DMA solutions containing 1,3-dimethyl-2-(o-hydroxyphenyl)-2,3-dihydro-1*H*-benzo[*d*]imidazole (BI(OH)H) at five different concentrations. The emission intensities decreased at higher BI(OH)H concentrations, indicating that BI(OH)H quenches the excited state of Os (Fig. S2a†). By using Stern-Volmer plots of the emission quenching (Fig. S2b†) and the lifetime of the emission ( $\tau_{\rm em}=40$  ns), the quenching rate constant was determined to be  $k_q = 1.8 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$ . The quenching fraction of excited **Os** due to BI(OH)H (0.2 M) was  $\eta_q = 60\%$ according to the equation  $\eta_q = k_q \tau_{em} [BI(OH)H]/(1 + k_q \tau_{em} [-1])$ BI(OH)H]); the reaction conditions were the same as those for photocatalytic CO2 reduction described below. To determine the product after excited Os was quenched by BI(OH)H, the absorption spectral changes during photoirradiation were measured.

The absorption spectra of a  $CO_2$ -saturated DMA solution containing only **Os** (0.1 mM) and BI(OH)H (0.2 M), without a catalyst, during irradiation and the difference spectra from the post- and pre-irradiation spectra are shown in Fig. 6a and b, respectively. Fig. 6b clearly shows that new broad absorption bands appeared at  $\lambda_{max} = 530$  and 800 nm and increased during



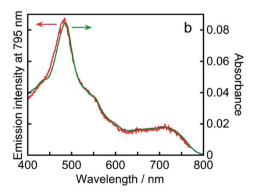
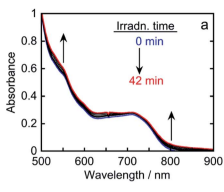


Fig. 5 (a) Emission spectra of Os in Ar-saturated DMA with excitation wavelengths of 450, 550, and 700 nm. (b) Excitation spectrum derived from emission intensity at  $\lambda_{max} = 795$  nm in DMA (red line) and absorption spectrum of Os (green line).



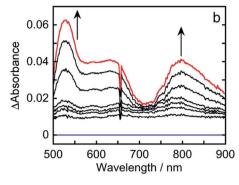


Fig. 6 (a) Absorption spectral changes of a  $CO_2$ -saturated DMA solution containing **Os** (0.1 mM) and BI(OH)H (0.2 M) during irradiation at 480 nm with a light intensity of  $2.6 \times 10^{-8}$  einstein s<sup>-1</sup> (0–42 min at 6 min intervals) and (b) difference spectra from post- and pre-irradiation spectra. The blue and red lines represent the spectra before irradiation and after irradiation for 42 min, respectively.

irradiation. The shapes of the difference spectra were fairly similar to that of the OERS of Os (Os: $^-$ ) obtained through electro-spectroscopy using an optically transparent thin-layer electrochemical (OTTLE) cell (Fig. 7). The absorption bands due to photoirradiation disappeared after aeration of the solution, which also supports the formation of Os: $^-$ , because this reduced species should be oxidized by  $O_2$ . These results clearly indicate that the  $^3$ MLCT excited state of Os was reductively quenched by BI(OH)H to give Os: $^-$  (eqn (2)).

$$\mathbf{Os} \xrightarrow{h\nu} *\mathbf{Os} \xrightarrow{\mathrm{BI(OH)H}} \mathbf{Os}^{-} \tag{2}$$

This reductive quenching process was also supported by thermodynamics. The 0-0 transition energy of **Os** was determined to be  $E_{00}=1.59$  eV from the Franck–Condon fitting of an emission spectrum with the vibronic structure measured in frozen DMA (Fig. S3†). It is noteworthy that this value was consistent with the  $E_{00}=1.6$  eV obtained from the mirror image of S–T absorption and phosphorescence (Fig. S4†). The reduction potential of **Os** in the lowest excited state was calculated to be  $E(*Os/Os^*)=E_{1/2}^{\rm red}+E_{00}=-1.57+1.59=+0.02$  V  $\nu$ s. Ag/AgNO $_3$ . The oxidation potential of BI(OH)H was measured to be  $E_{1/2}^{\rm vol}(BI(OH)H^*)$ -BI(OH)H)=-0.09 V from the cyclic voltammograms with a fast scan rate of 200 V s $^{-1}$ 

using a micro glassy carbon working electrode (Fig. S5†). Therefore, electron transfer from BI(OH)H to the excited state of **Os** in the reductive quenching process should be exothermic. Hence, we can summarize the photochemistry of **Os** in the presence of BI(OH)H as follows: **Os** can absorb the full wavelength range of visible light, giving the lowest <sup>3</sup>MLCT excited state, which is reduced by BI(OH)H to generate its relatively stable OERS.

The photocatalytic reduction of CO<sub>2</sub> was performed using **Os** as a redox photosensitizer. Ru(bpy)(CO)<sub>2</sub>Cl<sub>2</sub> (**Ru(CO)**), which has been used frequently as a catalyst for CO<sub>2</sub> reduction, <sup>21–23</sup> was employed because its reduction potential has been reported to be  $E_p = -1.51$  V vs. Ag/AgNO<sub>3</sub>, <sup>24</sup> that is, electron transfer should proceed thermodynamically from **Os** ·  $(E_{1/2}(\textbf{Os/Os} \cdot \textbf{Os})) = -1.57$  V) to **Ru(CO)**. A CO<sub>2</sub>-saturated DMA solution containing **Os** (50  $\mu$ M), **Ru(CO)** (50  $\mu$ M), and BI(OH)H (0.2 M) as a sacrificial electron donor was irradiated using a 725 nm LED light source, affording HCOOH with high selectivity (Fig. 8). After irradiation for 40 h, 8.1  $\mu$ mol of HCOOH was produced along with 0.3  $\mu$ mol of CO and a negligible amount of H<sub>2</sub> (<0.01  $\mu$ mol), corresponding to turnover numbers of TON<sub>HCOOH</sub> = 81, TON<sub>CO</sub> = 3, and TON<sub>H2</sub> < 1. The selectivity for HCOOH was 96%. The quantum yield for HCOOH formation was determined to be

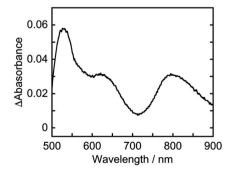


Fig. 7 Difference absorption spectrum from before and after electrolytic one-electron reduction of Os. A CO<sub>2</sub>-saturated DMA solution containing Os (0.5 mM) and Et<sub>4</sub>NBF<sub>4</sub> (0.1 M) as an electrolyte was reduced at -1.75 V using a UV-vis OTTLE cell with a Pt mesh working electrode and Ag/AgNO<sub>3</sub> reference electrode.

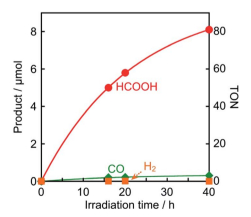


Fig. 8 Photocatalytic formation of HCOOH ( $\bullet$ ), CO ( $\bullet$ ), and H<sub>2</sub> ( $\blacksquare$ ): CO<sub>2</sub>-saturated DMA solutions (2 mL) containing Os (50  $\mu$ M), Ru(CO) (50  $\mu$ M), and BI(OH)H (0.2 M) were irradiated with 725 nm light.

 Table 2
 Products of photocatalysis after irradiation for 3 had

		Product/µmol (TON)		
Entry	Absence	НСООН	CO	$H_2$
1	_	1.9 (19)	0.05	n.d.
$2^b$	Os	0.04	n.d.	n.d.
$3^b$	Ru(CO)	n.d.	n.d.	n.d.
$4^b$	BI(OH)H	n.d.	n.d.	n.d.
5 <sup>c</sup>	Light irradiation	n.d.	n.d.	n.d.
$6^d$	$CO_2$	n.d.	n.d.	n.d.

<sup>a</sup> A CO<sub>2</sub>-saturated DMA solution (2 mL) containing Os (50 μM), Ru(CO) (50 μM), and BI(OH)H (0.2 M) was irradiated with 725 nm light.
<sup>b</sup> Photocatalytic reactions were performed without Os, Ru(CO), or BI(OH)H.
<sup>c</sup> The solution was placed in the dark for 3 h.
<sup>d</sup> An Arsaturated solution was used.

 $\Phi_{\rm HCOOH} = 0.061\%$  (Fig. S6†). Since the quenching fraction was  $\eta_{\rm q}=60\%$  when [BI(OH)H] = 0.2 M, the efficiency of HCOOH formation after the reductive quenching was  $\Phi_{\text{HCOOH}}/\eta_{\text{q}} =$ 0.1%. Table 2 summarizes the results of control experiments without Os, Ru(CO), BI(OH)H, light irradiation, or CO2. Entry 1 shows the result for the system with all components; that is, Os, Ru(CO), and BI(OH)H with irradiation for 3 h under CO<sub>2</sub> gave  $TON_{HCOOH} = 19$ . Irradiation of solutions without **Os** (entry 2), Ru(CO) (entry 3), or BI(OH)H (entry 4) did not lead to even a catalytic amount of the products. Control experiments in the dark (entry 5) or using an Ar-saturated solution (entry 6) yielded no photocatalysis products. Because only Os absorbs irradiated light at 725 nm under the photocatalytic reaction conditions (Fig. S7†), these results indicate that **Os** functioned as a redox photosensitizer and drove the photocatalytic formation of нсоон.

To determine the source of the carbon atom in the produced HCOOH, isotope labeling experiments were performed using  $^{13}\text{CO}_2$ . Fig. 9a and b show the  $^{13}\text{C}\{^1\text{H}\}$  NMR spectra of a DMF- $d_7$  solution containing **Os** (0.5 mM), **Ru(CO)** (50  $\mu$ M), and BI(OH)H (0.2 M) under a  $^{13}\text{CO}_2$  (447 mmHg) atmosphere before and after

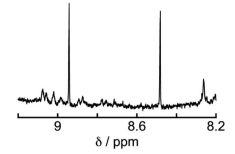


Fig. 10 <sup>1</sup>H NMR spectrum of the solution in Fig. 9 after irradiation.

irradiation at 725 nm using an LED for 60 h, respectively. After irradiation, a new signal appeared at 166.7 ppm, which was attributed to an equilibrium mixture of H13COOH and  ${\rm H}^{13}{\rm COO}^{-}$  (Fig. 9b). A doublet ( ${}^{1}J_{\rm CH} = 183~{\rm Hz}$ ) signal at 8.71 ppm in the <sup>1</sup>H NMR spectrum of the same solution (Fig. 10) was attributed to the methine proton of the equilibrium mixture of H<sup>13</sup>COOH and H<sup>13</sup>COO<sup>-</sup>. The absence of a clear singlet signal at 8.71 ppm attributable to the methine proton of an equilibrium mixture of H<sup>12</sup>COOH and H<sup>12</sup>COO<sup>-</sup> indicates that HCOOH was produced through the reduction of CO2. An identical J value of 183 Hz from the <sup>13</sup>C-<sup>1</sup>H coupling observed in the <sup>13</sup>C NMR spectrum without <sup>1</sup>H decoupling (Fig. 9c) also supports the formation of H<sup>13</sup>COOH. Therefore, we can conclude that CO<sub>2</sub> can be photocatalytically reduced to HCOOH using a system consisting of Os as the photosensitizer, Ru(CO) as the catalyst, and BI(OH)H as the reductant.

It is noteworthy that this system provides the first example of photocatalytic  $CO_2$  reduction driven by only light at  $\lambda_{ex}$  > 700 nm. We investigated the possibility of using even longer wavelengths. Irradiation with light only at  $\lambda_{ex}$  > 770 nm using a xenon lamp with a short-cut filter (LI0780, Asahi Spectra Co.) to a  $CO_2$ -saturated DMA solution (4 mL) of **Os** (0.6 mM), **Ru(CO)** (50  $\mu$ M), and BI(OH)H (0.2 M) for 5.5 h also gave 4.5  $\mu$ mol of HCOOH, although those using  $[Os(5dmb)_2(dmb)]^{2+}$  or

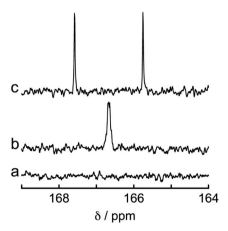


Fig. 9  $^{13}$ C( $^{1}$ H) NMR spectra of a DMF- $d_7$  solution of **Os** (0.5 mM), Ru(CO) (50  $\mu$ M), and Bl(OH)H (0.2 M) under a  $^{13}$ CO<sub>2</sub> (447 mmHg) atmosphere (a) before and (b) after 60 h of irradiation. (c)  $^{13}$ C NMR spectrum without  $^{1}$ H decoupling of the same solution after irradiation.

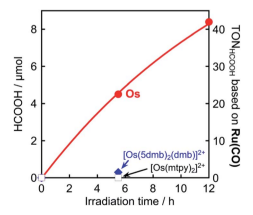


Fig. 11 Photocatalytic formation of HCOOH using light at  $\lambda_{ex} > 770$  nm and Os (  $\odot$ ),  $\left[ \text{Os}(5\text{dmb})_2(\text{dmb}) \right]^{2+} ( )$ , or  $\left[ \text{Os}(\text{mtpy})_2 \right]^{2+} ( )$  as a redox photosensitizer: CO<sub>2</sub>-saturated DMA solutions (4 mL) containing osmium(II) photosensitizer (0.6 mM), Ru(CO) (50  $\mu$ M), and BI(OH)H (0.2 M) were irradiated.

1.5

 $[Os(5dmb)_2(dmb)]^{2+}$ 

 $[Os(mtpy)_2]^{2}$ 

 Photosensitizer
 Irradiation time/h
 HCOOH/μmol
 TON<sub>HCOOH</sub>OS
 TON<sub>HCOOH</sub>Ru(CO)

 Os
 5.5
 4.5
 1.9
 23

 12
 8.4
 3.5
 42

0.3

Table 3 Photocatalytic formation of HCOOH using light at  $\lambda_{ex} > 770 \text{ nm}^a$ 

5.5

 $[Os(mtpy)_2]^{2^+}$  instead of **Os** scarcely produced HCOOH (Fig. 11, Table 3). This should be due to the poor absorption of  $[Os(5dmb)_2(dmb)]^{2^+}$  and  $[Os(mtpy)_2]^{2^+}$  at  $\lambda > 750$  nm as shown in Fig. 1. Further irradiation for 12 h in the same condition using **Os** gave 8.4 µmol of HCOOH which corresponds to  $TON_{HCOOH} = 3.5$  based on **Os** used and  $TON_{HCOOH} = 42$  based on **Ru(CO)** used. Therefore, **Os** can function as a panchromatic redox photosensitizer.

## Experimental

#### General procedures

<sup>1</sup>H NMR spectra were obtained in chloroform-d or acetonitrile- $d_3$ using a JEOL ECA400II (400 MHz) system to identify the synthesized ligands and complexes. The residual protons of chloroform-d or acetonitrile- $d_3$  were used as internal standards. Electrospray ionization-mass spectrometry (ESI-MS) was performed using a Shimadzu LCMS-2010A system with acetonitrile as the mobile phase. UV-vis absorption spectra were measured using a JASCO V-670 spectrophotometer. Prior to the emission measurements, the solutions were purged with Ar to remove dissolved O2. Emission spectra and excitation spectra derived from emission intensity were measured using a HORIBA Fluorolog-3-21 spectrofluorometer equipped with an NIR-PMT R5509-43 near-infrared detector. The emission lifetime was determined using a HORIBA FluoroCube time-correlated singlephoton counting system. The excitation light source was a NanoLED-460 pulse lamp (456 nm; pulse duration: 1.3 ns). The emission quantum yield was determined using a Quantaurus-QY Plus C13534-01 quantum yield analyzer (HAMAMATSU) using a xenon lamp with a band-pass filter (475  $\pm$  25 nm) as the light source. Emission-quenching experiments were performed in CO<sub>2</sub>-saturated solutions containing Os and BI(OH)H at five different concentrations. The quenching rate constant,  $k_q$ , was obtained from the slope of the Stern-Volmer plots of emission intensity against BI(OH)H concentration. The redox potentials of Os were measured in an Ar-saturated DMA solution containing Os (0.5 mM) and Et<sub>4</sub>NBF<sub>4</sub> (0.1 M) as a supporting electrolyte via cyclic voltammetry using an ALS CHI-620Ex electrochemical analyzer with a glassy carbon working electrode (diameter: 3 mm), Ag/AgNO<sub>3</sub> (10 mM) reference electrode, and Pt counter electrode. The scan rate was 0.2 V s<sup>-1</sup>. To determine the redox potential of BI(OH)H, a fast scan rate of 200 V s<sup>-1</sup> was applied while using a micro glassy carbon electrode (diameter: 33 μm).

#### **Emission spectral fitting**

Franck–Condon analysis of the emission spectrum obtained in DMA frozen using liquid nitrogen was conducted to determine the 0-0 transition energy,  $E_{00}$ , according to the equation below:<sup>25</sup>

0.1

$$I(\tilde{\nu}) = \sum_{\substack{n_1 = 0 \\ n_2 = 0}}^{5} \left( \frac{E_{00} - n_1 \tilde{\nu}_1 - n_2 \tilde{\nu}_2}{E_{00}} \right)^4 \left( \frac{S_1^{n_1}}{n_1!} \right) \left( \frac{S_1^{n_2}}{n_2!} \right)$$

$$\exp \left[ -4 \log 2 \left( \frac{\tilde{\nu} - E_{00} + n_1 \tilde{\nu}_1 + n_2 \tilde{\nu}_2}{\tilde{\nu}_{1/2}} \right)^2 \right]$$

where  $I(\nu)$  is the normalized intensity in the emission spectrum (wavenumber),  $S_i$  are the Huang–Rhys parameters of vibronic couplings reflecting the distortion in the acceptor modes,  $\nu_i$  are the energies of the acceptor modes, and  $\nu_{1/2}$  is the FWHM of the 0-0 vibronic component in the emission spectrum.

#### **Photocatalysis**

Photocatalytic reactions were performed in DMA solutions containing Os (50  $\mu$ M), Ru(CO) (50  $\mu$ M), and BI(OH)H (0.2 M). After it was purged with CO<sub>2</sub> for 20 min, the solution was irradiated. To determine the TON, a 2 mL solution in an 11 mL test tube was irradiated using a merry-go-round-type photoirradiation apparatus with an LED light source (725 nm), Iris-MG (CELL System Co.). The quantum yield was determined using a QYM-01 Shimadzu photoreaction quantum yield evaluation system; a 3.5 mL solution in a quartz cubic cell (11 mL; light pass length: 1 cm) was irradiated with 480 nm light from a 300 W xenon lamp equipped with a band-pass filter (480 nm; FWHM: 10 nm). The temperature of the solution during irradiation was maintained at 25  $\pm$  0.1  $^{\circ}$ C using an IWAKI CTS-134A constant temperature system. The gaseous products of the photocatalysis, namely, CO and H2, were analyzed using gas chromatography-thermal conductivity detector (GC-TCD) (GL Sciences GC323). HCOOH was analyzed using a capillary electrophoresis system (Agilent Technologies 7100L).

#### UV-vis absorption spectral changes during irradiation

A CO $_2$ -saturated DMA solution (4 mL) containing **Os** (0.1 mM) and BI(OH)H (0.2 M) in a quartz cubic cell (light pass length: 1 cm) was irradiated at 480 nm using a xenon lamp equipped with a 480 nm band-pass filter (Asahi Spectra Co.) and a 5 cm-long water filter. During irradiation, UV-vis absorption spectral changes were measured using a Photal MCPD-9800

<sup>&</sup>lt;sup>a</sup> A CO<sub>2</sub>-saturated DMA solution (4 mL) containing osmium(II) photosensitizer (0.6 mM), **Ru(CO)** (50 μM), and BI(OH)H (0.2 M) was irradiated using light at  $\lambda_{ex}$  > 770 nm.

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

Open Access Article. Published on 28 September 2021. Downloaded on 7/29/2025 12:09:29 AM

**Edge Article Chemical Science** 

spectrophotometer. The temperature of the solution was maintained at 25  $\pm$  0.1  $^{\circ}$ C using an IWAKI CTS-134A constant temperature system.

#### Electrochemical spectroscopy

A UV-vis OTTLE cell (light pass length: 1.0 mm) equipped with a Pt mesh working electrode, Ag/AgNO<sub>3</sub> (10 mM) reference electrode, and Pt counter electrode was employed for UV-vis electrochemical spectroscopy. UV-vis absorption spectral changes were measured using a Photal MCPD-9800 spectrometer. A CO<sub>2</sub>-saturated DMA solution containing Os (0.5 mM) and Et<sub>4</sub>NBF<sub>4</sub> (0.1 M) as the supporting electrolyte was electrolyzed using an ALS CHI-620Ex electrochemical analyzer.

#### **Ouantum chemical calculation**

The calculations were performed using the Gaussian 16 program<sup>16</sup> using the B3LYP exchange-correlation functional. Geometry optimization was performed in DMA using a general basis set with the Los Alamos National Laboratory effective core potential LanL2DZ basis set for osmium and 6-311++G(d,p) for the other elements. The geometry was fully optimized without symmetry constraints. Frequency calculations were performed with the same level of theory to ensure that the optimized geometry was a local minimum. The fractional contributions of various groups to each molecular orbital were calculated using GaussSum<sup>26</sup> based on Mulliken population analysis.

TD-DFT excited state calculations of the singlet-singlet and singlet-triplet transitions were performed in DMA using the geometry optimized by DFT calculations; the same LanL2DZ and 6-311++G(d,p) basis sets were used.

#### Isotope labeling experiment using <sup>13</sup>CO<sub>2</sub>

An isotope labeling experiment using <sup>13</sup>CO<sub>2</sub> was performed in a DMF- $d_7$  solution containing Os (0.5 mM), Ru(CO) (50  $\mu$ M), and BI(OH)H (0.2 M). The solution was deaerated using the freezepump-thaw method prior to the introduction of <sup>13</sup>CO<sub>2</sub> (447 mmHg). 13C(1H), 13C, and 1H NMR spectra were measured using a JEOL ECA400II spectrometer (100 MHz for <sup>13</sup>C NMR and 400 MHz for <sup>1</sup>H NMR) before and after irradiation for 60 h using a LED (725 nm) of Iris-MG (CELL System co.) as a light source. The residual carbons and protons of DMF- $d_7$  were used as internal standards for these measurements.

#### **Materials**

DMA was dried over 4 Å molecular sieves, distilled under reduced pressure, and kept under Ar before use. Et4NBF4 was dried under vacuum at 100 °C overnight prior to use. Ru(bpy)(CO)<sub>2</sub>Cl<sub>2</sub>,<sup>27</sup> BI(OH)H,<sup>28,29</sup> bis(N-methylbenzimidazolyl) pyridine (mbip),<sup>30</sup> and 4'-methyl-2,2':6',2"-terpyridine (mtpy)<sup>31</sup> were synthesized according to methods reported in the literature.

#### **Synthesis**

[Os(mbip)Cl<sub>3</sub>]. A N,N-dimethylformamide solution (13 mL) containing  $(NH_4)_2[Os^{IV}Cl_6]$  (258 mg, 0.59 mmol) and bis(N-

methylbenzimidazolyl)pyridine (205 mg, 0.61 mmol) was refluxed for 1 h. After the solution had cooled to room temperature, water was added. The resulting black precipitate was collected. Yield: 335 mg (0.53 mmol, 90%).

 $[Os(mbip)(\mu-Cl)Cl]_2$ . A N<sub>2</sub>-saturated ethanol solution (35 mL) containing [Os(mbip)Cl<sub>3</sub>] (333 mg, 0.53 mmol) and triethylamine (0.9 mL) was refluxed for 2 h under a  $N_2$  atmosphere. The black precipitate was filtered hot and washed with ethanol and diethyl ether. Yield: 157 mg (0.13 mmol, 50%).

[(mbip)Os(mtpy)](PF<sub>6</sub>)<sub>2</sub> (Os). A mixed solution of ethanol (8.5 mL) and H<sub>2</sub>O (8.5 mL) containing Os(mbip)(μ-Cl)Cl (160 mg, 0.13 mmol) and 4'-methyl-2,2':6',2"-terpyridine (66 mg, 0.27 mmol) was heated at 180 °C for 1 h using a microwave reactor, Monowave 200 (Anton Paar). After the solvent was removed, the product was purified using ion-exchange column chromatography (packing material: SP Sephadex C-25; eluent: acetonitrilewater (1:1 v/v) containing NH<sub>4</sub>PF<sub>6</sub> (0-14 mM)). Yield: 69 mg (0.060 mmol, 24%). <sup>1</sup>H NMR (400 MHz, acetonitrile- $d_3$ )  $\delta$ /ppm: 8.75 (s, 2H), 8.68 (d, 2H, J = 8.2 Hz), 8.30 (d, 2H, J = 7.8 Hz), 7.71(dd, 1H, J = 8.2, 8.2 Hz), 7.61-7.53 (m, 4H), 7.39 (dd, 2H, J = 7.8,7.8 Hz), 7.21 (d, 2H, J = 5.5 Hz), 7.00–6.94 (m, 4H), 6.00 (d, 2H, J= 8.2 Hz), 4.52 (s, 6H), 3.36 (s, 3H). ESI-MS (acetonitrile) m/z: 388 ( $[M - 2PF_6^{-}]^{2+}$ ), 923 ( $[M - PF_6^{-}]^{+}$ ). Anal. Calcd for  $C_{37}^{-}$ H<sub>30</sub>F<sub>12</sub>N<sub>8</sub>OsP<sub>2</sub>·H<sub>2</sub>O: C, 40.96; H, 2.97; N, 10.33. Found: C, 40.83; H, 3.03; N, 10.39.

#### Conclusion

Os absorbed the entire wavelength range of visible light up to 800 nm, generating the same triplet lowest excited state. The lifetime of the resulting excited state was  $\tau_{\rm em} = 40$  ns, which is relatively long and enabled the initiation of electron-transfer processes. Owing to longer-wavelength absorption by S-T transitions, Os exhibited both panchromatic absorption and a relatively long lifetime in the excited state. Excited Os was reductively quenched by BI(OH)H with a rate constant of  $k_q =$  $1.8 \times 10^8 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$ , giving an OERS with a reducing power of  $E_{1/2}$  $_{2}(Os^{\bullet -}) = -1.57 \text{ V} \text{ vs. Ag/AgNO}_{3}$ . In combination with **Ru(CO)** as a catalyst, Os functioned as a redox photosensitizer and drove the photocatalytic reduction of CO2 to HCOOH under light irradiation at  $\lambda_{\rm ex} = 725$  nm. This is the first example of photocatalytic CO<sub>2</sub> reduction *via* irradiation with  $\lambda_{ex} > 700$  nm light.

#### Author contributions

YT and OI designed the study. MI and YT performed all experiments. YT and OI wrote the manuscript.

#### Conflicts of interest

There are no conflicts to declare.

# Acknowledgements

The work was supported by JSPS KAKENHI Grant Number JP18H05355 and JP17H06440 in Scientific Research on Innovative Areas "Innovations for Light-Energy Conversion (I4LEC)", and IP18K14238.

#### References

- 1 H. Takeda, C. Cometto, O. Ishitani and M. Robert, *ACS Catal.*, 2017, 7, 70–88.
- 2 Y. Tamaki and O. Ishitani, ACS Catal., 2017, 7, 3394-3409.
- 3 H. Ozawa and K. Sakai, *Chem. Commun.*, 2011, 47, 2227-2242.
- 4 S. Fukuzumi, J. Jung, Y. Yamada, T. Kojima and W. Nam, *Chem.-Asian J.*, 2016, **11**, 1138–1150.
- 5 G. P. Smestad, F. C. Krebs, C. M. Lampert, C. G. Granqvist, K. L. Chopra, X. Mathew and H. Takakura, Sol. Energy Mater. Sol. Cells, 2008, 92, 371–373.
- 6 C. K. Prier, D. A. Rankic and D. W. C. MacMillan, *Chem. Rev.*, 2013, 113, 5322–5363.
- 7 D. W. Thompson, A. Ito and T. J. Meyer, *Pure Appl. Chem.*, 2013, **85**, 1257–1305.
- 8 A. Juris, V. Balzani, F. Barigelletti, S. Campagna, P. Belser and A. von Zelewsky, *Coord. Chem. Rev.*, 1988, **84**, 85–277.
- 9 A. Hagfeldt, G. Boschloo, L. Sun, L. Kloo and H. Pettersson, Chem. Rev., 2010, 110, 6595–6663.
- 10 F. Bella, C. Gerbaldi, C. Barolo and M. Grätzel, *Chem. Soc. Rev.*, 2015, 44, 3431–3473.
- 11 Y. Tsuji, K. Yamamoto, K. Yamauchi and K. Sakai, *Angew. Chem.*, *Int. Ed.*, 2018, **57**, 208–212.
- 12 B. D. Ravetz, N. E. S. Tay, C. L. Joe, M. Sezen-Edmonds, M. A. Schmidt, Y. Tan, J. M. Janey, M. D. Eastgate and T. Rovis, ACS Cent. Sci., 2020, 6, 2053–2059.
- 13 Y. Miyake, K. Nakajima, K. Sasaki, R. Saito, H. Nakanishi and Y. Nishibayashi, *Organometallics*, 2009, **28**, 5240–5243.
- Y. Tamaki, K. Koike, T. Morimoto, Y. Yamazaki and
   O. Ishitani, *Inorg. Chem.*, 2013, 52, 11902–11909.
- 15 J.-Y. Shao and Y.-W. Zhong, *Inorg. Chem.*, 2013, **52**, 6464-6472
- 16 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega,
  - G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota,

O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo,

R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda,

- R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman, and D. J. Fox, *Gaussian 16, Revision A.03*, Gaussian, Inc., Wallingford CT, 2016.
- 17 Z. Abedin-Siddique, Y. Yamamoto, T. Ohno and K. Nozaki, *Inorg. Chem.*, 2003, 42, 6366–6378.
- 18 Z. Abedin-Siddique, T. Ohno, K. Nozaki and T. Tsubomura, *Inorg. Chem.*, 2004, 43, 663–673.
- 19 K. Nozaki, J. Chin. Chem. Soc., 2006, 53, 101-112.
- 20 S. Obara, M. Itabashi, F. Okuda, S. Tamaki, Y. Tanabe, Y. Ishii, K. Nozaki and M. Haga, *Inorg. Chem.*, 2006, 45, 8907–8921.
- 21 J.-M. Lehn and R. Ziessel, J. Organomet. Chem., 1990, 382, 157–173.
- 22 H. Ishida, K. Fujiki, T. Ohba, K. Ohkubo, K. Tanaka, T. Terada and T. Tanaka, *J. Chem. Soc., Dalton Trans.*, 1990, 2155–2160.
- 23 Y. Kuramochi, O. Ishitani and H. Ishida, *Coord. Chem. Rev.*, 2018, 373, 333–356.
- 24 Y. Kuramochi, J. Itabashi, K. Fukaya, A. Enomoto, M. Yoshida and H. Ishida, *Chem. Sci.*, 2015, **6**, 3063–3074.
- 25 G. H. Allen, R. P. White, D. P. Rillema and T. J. Meyer, J. Am. Chem. Soc., 1984, 106, 2613–2620.
- 26 N. M. O'boyle, A. L. Tenderholt and K. M. Langner, J. Comput. Chem., 2008, 29, 839–845.
- 27 P. A. Anderson, G. B. Deacon, K. H. Haarmann, F. R. Keene, T. J. Meyer, D. A. Reitsma, B. W. Skelton, G. F. Strouse, N. C. Thomas, J. A. Treadway and A. H. White, *Inorg. Chem.*, 1995, 34, 6145–6157.
- 28 E. Hasegawa, T. Seida, N. Chiba, T. Takahashi and H. Ikeda, *J. Org. Chem.*, 2005, **70**, 9632–9635.
- 29 X.-Q. Zhu, M.-T. Zhang, A. Yu, C.-H. Wang and J.-P. Cheng, *J. Am. Chem. Soc.*, 2008, **130**, 2501–2516.
- 30 J. J. Concepcion, J. W. Jurss, M. R. Norris, Z. Chen, J. L. Templeton and T. J. Meyer, *Inorg. Chem.*, 2010, 49, 1277–1279.
- 31 Y. Tamaki and O. Ishitani, *Faraday Discuss.*, 2017, 198, 319–335.