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Understanding the mechanism of direct visible-light-activated [2 + 2] cycloadditions mediated by Rh and Ir photocatalysts: combined computational and spectroscopic studies†

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The mechanism of [2 + 2] cycloadditions activated by visible light and catalyzed by bis-cyclometalated Rh(III) and Ir(III) photocatalysts was investigated, combining density functional theory calculations and spectroscopic techniques. Experimental observations show that the Rh-based photocatalyst produces excellent yield and enantioselectivity whereas the Ir-photocatalyst yields racemates. Two different mechanistic features were found to compete with each other, namely the direct photoactivation of the catalyst–substrate complex and outer-sphere triplet energy transfer. Our integrated analysis suggests that the direct photocatalysis is the inner working of the Rh-catalyzed reaction, whereas the Ir catalyst serves as a triplet sensitizer that activates cycloaddition via an outer-sphere triplet excited state energy transfer mechanism.

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Introduction

Photocatalysts bearing transition metals have emerged as practical and valuable synthetic tools for organic chemists. These catalysts can form reactive radicals^{1–3} and triplet intermediates^{4,5} under relatively mild conditions and facilitate challenging transformations that are difficult to achieve by traditional synthetic methodologies. A successful photocatalytic reaction produces photogenerated radicals or triplet excited state intermediates in a well-controlled manner, while minimizing the production of undesired intermediates that result in byproducts and contaminate the catalyst. In this regard, there is a dire need for understanding the inner-workings of photocatalytic reactions to promote the rational design of novel photocatalytic transformations. A photocatalytic transformation is comprised of two reaction phases: (i) a photo-physical phase that activates substrates via photoexcitation, intersystem crossing (ISC), and electron or energy transfer, and

(ii) a reaction phase, wherein the activated substrates undergo desired bond formation and dissociation events. Unraveling the detailed mechanisms of photo-activated reactions requires advanced spectroscopic techniques such as time-resolved photoluminescence and transient absorption measurements,⁶ that have begun to be explored only recently.⁷ Computational studies^{8–11} have characterized reactive excited state intermediates, but mechanistic details of how the charge/energy transfer processes occur during the photochemical reaction remain poorly understood.

Asymmetric induction is of particular interest and constitutes a significant challenge in photocatalysis. Recent advances include the development of visible-light sensitized asymmetric [2 + 2] cycloadditions mediated by triplet excited energy transfer (TEET), which were pioneered by Bach,^{12–14} Yoon^{15–20} and Meggers.^{21–23} Photochemical [2 + 2] cycloaddition reactions engender a cyclobutyl functional group, that can not only produce valuable products but can also garner important intermediates en route to larger ring systems.^{24,25} Compared to photocatalysis utilizing single electron transfer (SET) pathways, photocatalytic transformations based on TEET mechanisms have broader substrate scope and withstand relatively mild reaction conditions.^{26–32} As such, there is growing interest in the development of asymmetric photocatalysts exploiting the TEET process, where the photocatalyst not only populates triplet excited state intermediates but also induces a chiral environment for stereoselective bond formation. Bach and co-workers adopted chiral Lewis acid and lactam photosensitizers to

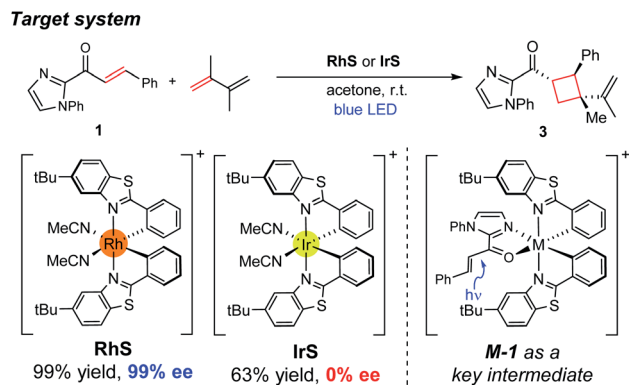
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Scheme 1 Experimental conditions of the enantioselective [2 + 2] cycloaddition reaction investigated in this study.

construct a hydrogen bonded substrate–catalyst complex to control the facial selectivity of intramolecular cycloaddition.^{33,34} Yoon and co-workers employed a dual catalysis approach, wherein the photocatalyst generates reactive intermediates and a chiral Lewis acid cocatalyst controls the stereoselectivity and enhances reactivity.²⁰ Meggers designed chiral-at-metal platforms that enable visible-light-activated [2 + 2] photocycloadditions with pronounced enantio- and diastereoselectivities.^{21,35}

Herein, we investigated the detailed mechanism of an enantioselective [2 + 2] photocycloaddition reported previously (Scheme 1).²¹ The transformation utilizes a chiral-at-metal Rh catalyst (**RhS**) that enables the [2 + 2] photocycloaddition of enones and terminal alkenes with high enantioselectivity and diastereoselectivity. Interestingly, when using an analogous Ir catalyst (**IrS**), a racemic product mixture was obtained with a moderate yield. The different outcomes of the two transition-metal photocatalysts drew our attention, as the two isolobal metal centres were expected to exhibit similar reactivities. In order to comprehend these unexpected outcomes, we combined computational and spectroscopic analyses that characterize both the photophysical and the reaction phases of the reactions. Specifically, we delineated the mechanistic details of the photophysical phase of the reactions that activate the substrate–catalyst complex to access the triplet-state electronic manifold and rationalized the different outcomes from the Rh- and Ir-photocatalysts.

Results and discussion

Catalytic cycle of direct photocatalysis

Fig. 1 summarizes the proposed photocatalytic cycle of the [2 + 2] cycloaddition. Previously, the direct photoexcitation of the catalyst–substrate complex was suggested as a key mechanistic event to explain the high enantioselectivity of the Rh catalyst.²¹ The reaction begins with the substitution of two acetonitrile ligands in the metal centers of the photocatalyst by the enone substrate **1** to afford the chelated metal complexes **Rh-1**, that absorbs blue light to access the singlet excited states **¹Rh-1*. The generated singlet excited species undergoes fast internal**

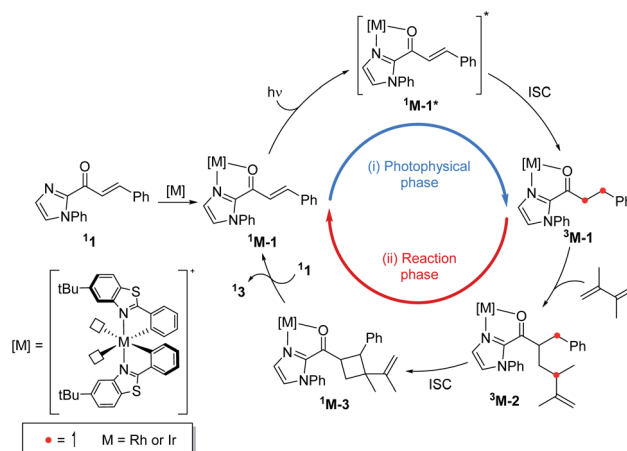


Fig. 1 The catalytic cycle of the direct absorption mechanism for [2 + 2] photocycloaddition. (M = Rh or Ir). The unpaired electrons are expressed with red dots.

conversion (IC) to the first singlet excited state (Kasha's rule³⁶), followed by intersystem crossing (ISC) to form the triplet state intermediates **³Rh-1**. The generated triplet excited state species undergoes a stepwise C–C bond formation with terminal alkenes to afford the diradical intermediate species **³Rh-2**. A radical recombination step completes the formation of the [2 + 2] cycloaddition product **¹Rh-3**, which is a highly exergonic and likely barrierless process.

Fig. 2 summarizes the computed reaction energy profiles of the [2 + 2] cycloadditions, initiated from the photoexcited triplet intermediates. We used density functional theory calculations

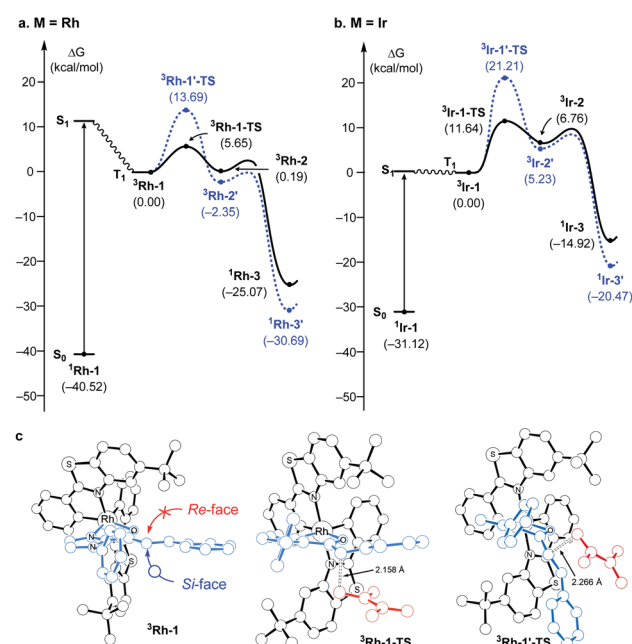


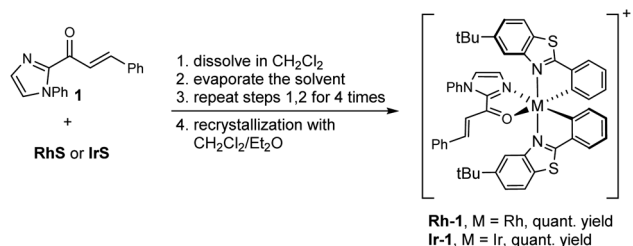
Fig. 2 (a) Computed energy profile following the direct absorption mechanism with the Rh- and (b) Ir- photocatalyst; (black: Si-face attack, blue: Re-face attack, plain: favored, dashed: disfavored) (c) optimized structures of **³Rh-1**, **³Rh-1-TS**, and **³Rh-1'-TS**.

to analyze the reaction energy profiles (see the ESI† for the details). For the Rh photocatalyst, the computed energy profile explained the observed enantioselectivity, as shown in Fig. 2a. $^3\text{Rh-1}$ can engage the terminal alkene and the α,β -carbons of enone of the substrate **1**. There are two possible faces of the metal-bound enone offered for the photocycloaddition reaction, as illustrated in Fig. 2c. The *Si*-face attack traverses the transition state $^3\text{Rh-1-TS}$ with an activation barrier of $5.7 \text{ kcal mol}^{-1}$, which is 8 kcal mol^{-1} lower in energy than the corresponding barrier of the *Re*-face attack at $13.7 \text{ kcal mol}^{-1}$ ($^3\text{Rh-1'-TS}$). This substantial difference in the activation energy is in line with the experimental observation of the Rh photocatalytic system, where an enantiomeric excess of $>99\%$ was observed. A comparison of the transition structures suggested that the *Re*-face imposes a significant steric demand on the approaching diene substrate at the corresponding transition structure ($^3\text{Rh-1'-TS}$) due to a bulky ^tBu group of the LX ligand.^{8,22} The likely short-lived 1,4-diradical intermediates $^3\text{Rh-2}$ and $^3\text{Rh-2'}$ are found at 0.2 and $-2.4 \text{ kcal mol}^{-1}$, relative to the initial triplet intermediate ($^3\text{Rh-1}$), respectively. A reverse ISC from the triplet ($^3\text{Rh-2}$ and $^3\text{Rh-2'}$) to singlet biradical state followed by radical recombination completes the cyclization ($^1\text{Rh-3}$ and $^1\text{Rh-3'}$). We were unable to locate the singlet biradical intermediate, as the C-C coupling occurs spontaneously.

In contrast, the computed reaction energy profile did not explain the observed racemic outcome from the Ir photocatalyst (Fig. 2b). The barriers for the initial C-C bond formations are notably higher with $^3\text{Ir-1-TS}$, that is located at $11.6 \text{ kcal mol}^{-1}$ from the triplet state intermediate, whereas that of the alternative transition state $^3\text{Ir-1'-TS}$ is $21.2 \text{ kcal mol}^{-1}$. Albeit the computed activation barriers are $5\text{--}7 \text{ kcal mol}^{-1}$ higher than those of the Rh-catalyst, the reactions are amenable at room temperature,³⁷ suggesting that the Ir-photocatalyst also exhibits pronounced enantioselectivity. The discrepancy between the computational results and the experimental outcomes strongly suggested that the proposed mechanistic pathway is no longer valid for the Ir-photocatalyst. In the following, we demonstrate that $^3\text{Ir-1}$ is an off-cycle intermediate and the products are formed by a different mechanism, by means of the photophysical and electrochemical measurements and the computational analyses of the photochemical phase.

Photophysical and electrochemical studies

Our computational analysis of the mechanism of the $[2 + 2]$ cycloadditions suggested that the reactions follow different



Scheme 2 Synthesis of **Rh-1** and **Ir-1**. The complexes were prepared as their hexafluorophosphate salts.

pathways depending on the transition metal complexes used. To explore these possibilities, we prepared the intermediates **Rh-1** and **Ir-1** (Scheme 2) and studied their photophysical and electrochemical properties.

Fig. 3 shows the results of photophysical measurements of the substrate **1**, metal complexes **RhS** and **IrS**, and substrate bound intermediates (**Rh-1** and **Ir-1**), in CH_2Cl_2 at room temperature and in a $\text{CH}_2\text{Cl}_2 : \text{CHCl}_3$ 1 : 1 (v/v) rigid matrix at 77 K. The absorption spectra of **RhS** (cyan line, Fig. 3a) and **Rh-1** (blue line, Fig. 3a) are quite similar and they exhibit bands at 380 nm and 385 nm respectively, which were assigned to singlet metal-to-ligand charge transfer ($^1\text{MLCT}$) states based on our computational study (*vide infra*). **Rh-1** presents an additional low-energy tail up to 500 nm. On the other hand, the absorption spectra of the two Ir(III) metal complexes were quite different from each other (Fig. 3c): **IrS** exhibits the lowest-energy absorption band peaked at 398 nm, while **Ir-1** has a weak absorption band centered at 640 nm with a tail up to 800 nm. The low-energy absorption band of **Ir-1** can be tentatively assigned to a triplet metal-to-ligand charge transfer ($^3\text{MLCT}$) state. As iridium possesses a much higher spin-orbit coupling compared to rhodium, a spin-forbidden electronic transition can be observed more easily.³⁸

As far as emission is concerned, the enone substrate **1** exhibits a weak phosphorescence at 77 K (red line, Fig. 3b) with a lifetime of 11 ms. The metal complexes **RhS** and **IrS** display phosphorescence bands in a rigid matrix at 77 K with the maximum at 514 nm and 527 nm, respectively (Fig. 3b and d): the former presents a well-resolved vibrational structure and a longer lifetime (τ) of 88 μs compared to the latter ($\tau = 5.6 \mu\text{s}$), suggesting a stronger LC characteristic of the electronic

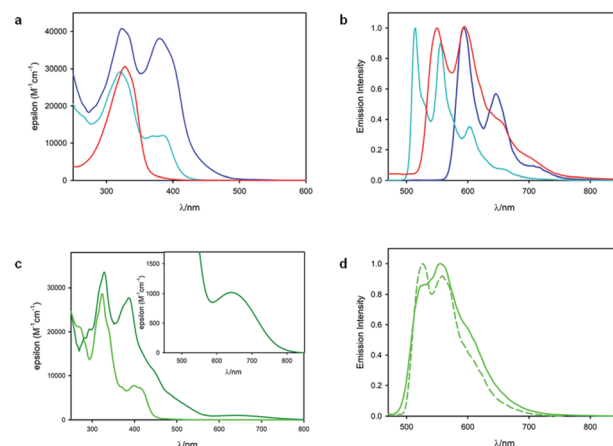


Fig. 3 (a) Absorption spectra of **RhS** (cyan line), **1** (red line) and **Rh-1** (blue line) in CH_2Cl_2 at room temperature. (b) Normalized phosphorescence spectra at 77 K of **Rh-1** (blue line, $\lambda_{\text{ex}} = 450 \text{ nm}$), **RhS** (cyan line, $\lambda_{\text{ex}} = 350 \text{ nm}$) and **1** (red line, $\lambda_{\text{ex}} = 350 \text{ nm}$) in $\text{CH}_2\text{Cl}_2 : \text{CHCl}_3$ 1 : 1 (v/v). (c) Absorption spectra of **IrS** (light green line) and **Ir-1** (dark green line) in CH_2Cl_2 at room temperature. The inset shows the low-energy band of **Ir-1**. (d) Normalized phosphorescence spectra of **IrS** at RT (solid line) in de-oxygenated CH_2Cl_2 solution and at 77 K (dashed line) in $\text{CH}_2\text{Cl}_2 : \text{CHCl}_3$ 1 : 1 (v/v); $\lambda_{\text{ex}} = 400 \text{ nm}$. Of note, we did not observe the phosphorescence signal from **Ir-1**.

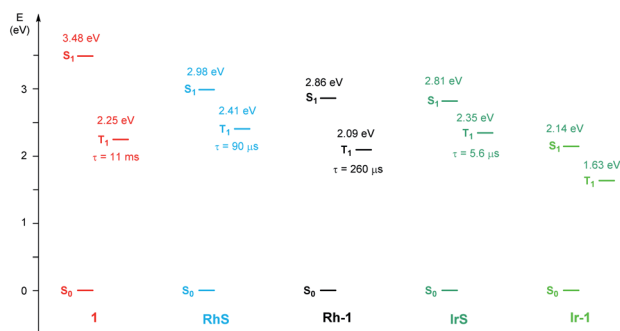


Fig. 4 Schematic Jablonski diagram for **1**, **RhS**, **Rh-1**, **IrS**, and **Ir-1** at 77 K.

transition for the **RhS** compared to the **IrS**. The substrate-bound intermediates **Rh-1** and **Ir-1** exhibit distinctly different emissive properties: **Ir-1** is not emissive both in deoxygenated CH_2Cl_2 solution at room temperature and in the rigid matrix at 77 K. We tentatively assign this dark state as $^3\text{MLCT}$, which rapidly decays to the ground state by non-radiative processes. This result agrees with the conclusions drawn from the computational studies that this state is not catalytically active (*vide infra*). **Rh-1** displays a vibrationally structured phosphorescence band with a maximum at 593 nm at 77 K and a lifetime of 260 μs. Based on the vibrational structure and lifetime, this band can be tentatively assigned to a ^3LC transition centered on ligand **1**. Fig. 4 shows the values of the lowest singlet (S_1) and triplet excited states (T_1) evaluated using absorption and emission spectra of the analyzed compounds.

We also determined the electrochemical properties of the metal complexes (Fig. 5). **RhS** undergoes a chemically irreversible oxidation with $E_{\text{pa}} = +1.76$ V at 0.5 V s^{-1} and a chemically reversible reduction at -1.55 V (partial chemical reversibility was attained at scan rate $>2 \text{ V s}^{-1}$). **Rh-1** displays a chemically irreversible oxidation at $+1.41$ V and a chemically reversible reduction at -0.71 V. The MLCT characteristic of the Ir complexes is compatible with the measured electrochemical properties. The energy difference between the first oxidation and the first reduction is 1.98 V, much lower than in **IrS** (2.44 V).

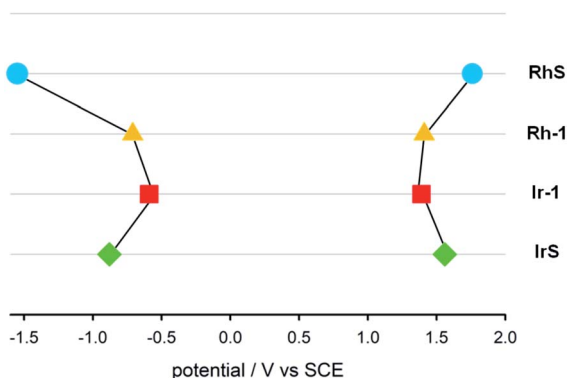


Fig. 5 Diagram of the redox potentials ($E_{1/2}$ or E_{pa} at 0.5 V s^{-1} in CH_2Cl_2 relative to SCE) of **RhS** (light-blue circle), **Rh-1** (yellow triangle), **Ir-1** (red square) and **IrS** (green diamond).

Table 1 Estimated redox potentials of the excited state transition metal complexes^a

Compound	$E([\text{M}]^+ / [\text{M}])$	$E([\text{M}] / [\text{M}]^-)$
RhS	-0.65	0.86
Rh-1	-0.68	1.38
IrS	-0.79	1.47
Ir-1 ^b	-0.26	1.04

^a The unit of potential is V. ^b We used the computed ground-triplet gap (1.54 eV) for the estimation (Fig. 6c).

IrS undergoes a chemically reversible oxidation at $+1.56$ V and a chemically reversible reduction at -0.88 V. **Ir-1** displays a chemically reversible oxidation at $+1.39$ V and a chemically reversible reduction at -0.59 V.

The redox potentials of the triplet excited state species precluded the photoredox activation of the substrates. Table 1 summarizes the estimated redox potentials of the four transition metal complexes. The redox potentials were computed using the ground state and the experimental ground state to triplet state energy gaps (Fig. 4 and 5).³⁹ The estimated reduction potentials (E_{red}) ranged from -0.79 V to -0.26 V and the oxidation potentials (E_{ox}) spanned from 0.86 V to 1.47 V. The redox potentials of the transition metal complexes were weak to reduce/oxidize the substrates effectively: the computed reduction (E_{red}) and oxidation potentials (E_{ox}) of the substrate **1** are -1.69 V and 1.56 V vs. SCE, respectively and those of dimethylbutadiene are -3.68 V and 1.42 V vs. SCE, respectively.

Computational studies of the excited state species

To understand the differences in the excited state electronic manifolds, **Rh-1** and **Ir-1** were further analyzed by means of excited-state electronic structure methods. The photophysical activation steps are summarized in Fig. 6a. The chelated metal complex species (**M-1**) consisting of M(III)-d^6 centres can absorb light to access singlet excited state species, that is subject to non-radiative internal conversion (IC) to the first singlet excited state species (S_1). Our computational analysis suggested the S_1 species bears a singlet MLCT characteristic (**M-1- $^1\text{MLCT}$**), where the metal centre is formally oxidized to a M(IV)-d^5 state and the ligand is reduced (*vide infra*). Intersystem crossing (ISC) mediated by the spin-orbit coupling allows the conversion of the singlet excited species to the triplet excited state species, wherein the alkene moiety of the enone substrate is activated to become a diradical, manifesting the characteristics of the ligand-centered (LC) triplet excited state (**M-1- ^3LC**). In the LC state, two independent electrons are localized at the π and π^* orbitals of the enone while the metal centre is reduced by one electron. Once the substrate is activated to access the triplet LC state, the reaction phase becomes operational as long as the photophysical deactivation is slower than the activation of the cycloaddition reaction. Triplet LC states can proceed further to become a triplet MLCT state, that is denoted as **M-1- $^3\text{MLCT}$** . Experimental evidence suggested the triplet MLCT states are

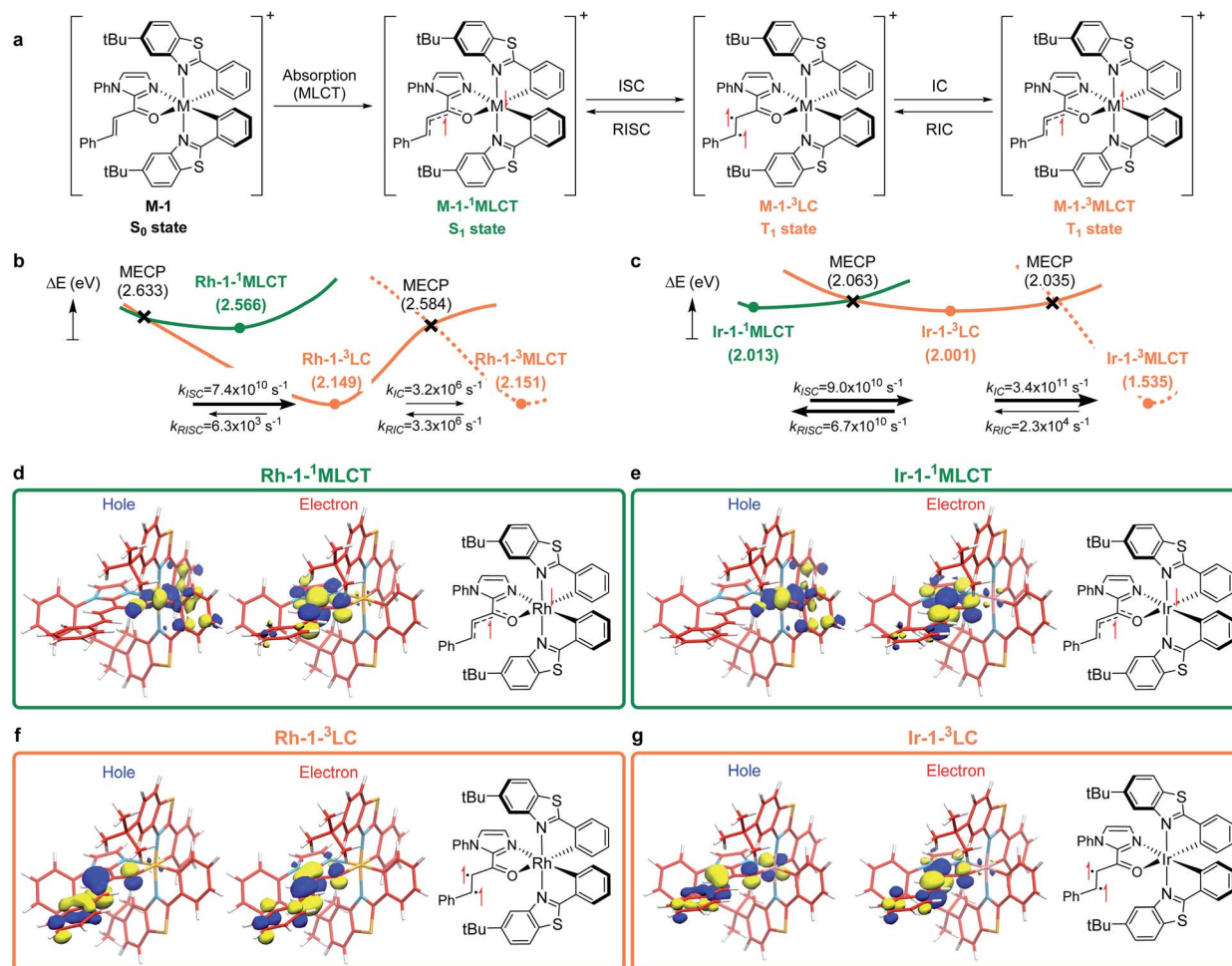


Fig. 6 (a) The photophysical route to access the triplet LC and MLCT states. Computed excited state energy surfaces of (b) Rh-1 and (c) Ir-1. The renditions of natural transition orbitals of the identified excited singlet (M-1-³MLCT) and triplet state (M-1-³LC) species of Rh-1 (d and f) and Ir-1 (e and g). Filled circles represent the energy of the optimized geometries on the corresponding energy surfaces; crosses designate minimum energy crossing points (MECPs) of two energy surfaces. Electronic energies were computed at the (TD)CAM-B3LYP-D3/6-31G(d,p)/LANL2DZ level of the theory.

prone to deactivation prior to engendering desired cycloaddition products.

We utilized time-dependent density functional theory (TD-DFT) calculations of the excited state species to characterize the energetics and electronic nature of the photoexcited intermediate species that are implicated in the photochemical phase of the reaction. Firstly, we optimized the first singlet excited state geometries of the chelated metal complexes Rh-1-¹MLCT and Ir-1-¹MLCT. The computed adiabatic energies relative to those of the optimized ground state geometries are 2.57 and 2.01 eV, respectively, in good agreement with the energetics of singlet MLCT states measured from the emission spectra (Fig. 3 and 4). Natural transition orbital (NTO)⁴⁰ analysis confirmed that the singlet excited state species bears a strong MLCT characteristic (Fig. 6d and e), where a metal d-orbital participates in the hole and the lowest π^* orbital of the enone contributes to the electron state. The observation is in line with the orbital contributions to the singlet excited state (Table S3[†]): a singlet MLCT state species comprises HOMO to LUMO

excitations, where the HOMO contains d-orbitals of the metal center and the LUMO consists of the π^* -orbital of enone.

We also optimized geometries following the triplet excited state manifold bearing the characteristics of π - π^* ligand-centered (LC) excitations (Rh-1-³LC and Ir-1-³LC) and the MLCT excitations (Rh-1-³MLCT and Ir-1-³MLCT). For the Rh complex, both the triplet LC and MLCT states (Rh-1-³LC and Rh-1-³MLCT) are lower in electronic energy by 0.4 eV than that of the singlet MLCT species Rh-1-¹MLCT. For the triplet LC state, significant contribution from the substrate π - π^* orbitals was observed, whereas the d- π^* transition was dominant for the triplet MLCT state (Table S3[†]). The adiabatic electronic energy of the Ir-complex Ir-1-³LC was calculated at 2.00 eV from that of the optimized ground state geometry, which was nearly isoenergetic to that of the singlet MLCT state Ir-1-¹MLCT. The triplet MLCT of the Ir complex (Ir-1-³MLCT) was 0.47 eV lower in energy than that of the singlet MLCT state (Ir-1-¹MLCT) and the triplet LC state (Ir-1-³LC). Again, the renditions of NTO of the triplet species confirmed that the excited species possess



the LC and the MLCT characteristics, as highlighted in Fig. 6f, g, and S1†. Similar to the Rh-complex, the $d-\pi^*$ transition in the $^3\text{MLCT}$ state and the substrate $\pi-\pi^*$ transition in the ^3LC state were identified (Table S3†). The energy difference between **Ir-1- $^3\text{MLCT}$** and **Ir-1- ^3LC** explained the higher activation barrier of the first bond formation step in the $[2+2]$ cycloaddition of **Ir-1**, than that of the **Rh-1** (Fig. 3), as the direct photocatalysis using the Ir complex necessitates the presence of **Ir-1- ^3LC** that is higher in energy than **Ir-1- $^3\text{MLCT}$** .

The calculated rates of ISC and ICs suggested that the Rh-photocatalyst populates triplet LC states with efficiency, which explains the enantioselective $[2+2]$ cycloaddition from the Rh-photocatalyst. The rates of ISC between the singlet MLCT and the triplet LC states were calculated based on Marcus theory,⁴¹ as summarized in Table 2. For the Rh-complex, the computed rate of ISC from the singlet MLCT **Rh-1- $^1\text{MLCT}$** to the triplet LC state **Rh-1- ^3LC** was $k_{\text{ISC}}(\text{Rh}) = 7.4 \times 10^{10} \text{ s}^{-1}$, whereas that of the reverse process was $k_{\text{RISC}}(\text{Rh}) = 6.3 \times 10^3 \text{ s}^{-1}$. The ratio of forward and reverse ISC rates demonstrates that the intersystem crossing is an irreversible process. The computed rates of internal conversions between **Rh-1- ^3LC** and **Rh-1- $^3\text{MLCT}$** were $k_{\text{IC}}(\text{Rh}) = 3.2 \times 10^6 \text{ s}^{-1}$ and $k_{\text{RIC}}(\text{Rh}) = 3.3 \times 10^6 \text{ s}^{-1}$, respectively. This finding indicated that the photophysical conversion of **Rh-1- ^3LC** to **Rh-1- $^3\text{MLCT}$** is 4000-fold slower than the ISC from **Rh-1- $^1\text{MLCT}$** whose lifetime is expected to be in the order of microseconds. The conversions between the singlet and triplet MLCT states were anticipated to be much slower. The computed rate of ISCs from **Rh-1- $^1\text{MLCT}$** to **Rh-1- $^3\text{MLCT}$** was $3.2 \times 10^5 \text{ s}^{-1}$ and the rate of the reverse process was negligibly small relative to the other rates of ISCs and ICs. This finding manifested the El-Sayed rule of ISCs: both the singlet and triplet MLCT states share similar orbital symmetries (Fig. 6d, e, and S1†), in which the ISCs are inefficient.⁴² Of note, at the minimum energy crossing point (MECP) of **Rh-1- $^1\text{MLCT}$** and **Rh-1- ^3LC** , the bond length of $\text{C}_\alpha-\text{C}_\beta$ is 1.347 Å, which is close to that of the optimized ground state (Table S1†), indicating that the intersystem crossing occurs in the Marcus inverted region.⁴³

Based on the computations of the rates of ISCs and ICs of **Ir-1**, we concluded that the ^3LC state of the Ir-substrate complex is deactivated before engaging in photocycloaddition. The computed rates of ISC from **Ir-1- $^1\text{MLCT}$** to **Ir-1- ^3LC** was $k_{\text{ISC}}(\text{Ir}) = 9.0 \times 10^{10} \text{ s}^{-1}$, whereas that of the reverse ISC from **Ir-1- ^3LC** to **Ir-1- $^1\text{MLCT}$** was $k_{\text{RISC}}(\text{Ir}) = 6.7 \times 10^{10} \text{ s}^{-1}$. The singlet MLCT and triplet LC states of the Ir-complexes are isoenergetic, affording comparable forward and reverse rates. The internal conversion from **Ir-1- ^3LC** to **Ir-1- $^3\text{MLCT}$** was anticipated to be a fast and irreversible process. The rate of internal conversion ($k_{\text{IC}}(\text{Ir})$) was computed as $3.5 \times 10^{11} \text{ s}^{-1}$ which was comparable to that of intersystem crossings. The rate of the reverse process from **Ir-1- $^3\text{MLCT}$** to **Ir-1- ^3LC** ($k_{\text{RIC}}(\text{Ir})$) was calculated to be $2.3 \times 10^4 \text{ s}^{-1}$, that is 10^7 -fold slower than that of the forward process. The finding suggested that the triplet LC state of **Ir-1** (**Ir-1- ^3LC**) is short lived and cannot engage in the $[2+2]$ cycloaddition with the diene.

Comparisons of the electronic structures of Rh and Ir systems

We demonstrated above that the triplet LC state of the Rh complex is kinetically stable, whereas that of the Ir complex is not. We further investigated electronic transitions within frontier molecular orbitals (FMOs) of both of the chelated metal complexes, which rationalized the differences between the $^1\text{MLCT}$ and ^3LC states of the Rh and Ir systems.

Fig. 7a highlights the electronic transitions that occur during the photochemical phase. In the electronic ground state, a metal complex **M-1** features the characteristics of d^6 -octahedral transition metal complexes, wherein one of the d-orbitals of the metal center (d_{xy}) constitutes the HOMO level and the π and π^* levels from the enone substrate **1** constitutes one of the high-lying doubly occupied orbitals and the LUMO level, respectively. The MLCT from the metal d_{xy} level to the π^* -level of the ligand creates a **M-1- $^1\text{MLCT}$** state, which formally oxidizes the metal center while reducing the ligand. The ISC promotes one electron from the π level of the substrate to the half-filled d_{xy} orbital, resulting in the formation of the ^3LC state (**M-1- ^3LC**). The reverse ISC (RISC) conveys the backward

Table 2 Detailed parameters for the computations of k_{ISC} and k_{IC}^a

Molecule	Transition	K	V	ΔG	A
Rh-1	$^1\text{MLCT} \rightarrow ^3\text{LC}$	7.4×10^{10}	17.14	-9.61	12.74
	$^3\text{LC} \rightarrow ^1\text{MLCT}$	6.3×10^3	17.14	9.61	12.74
	$^3\text{LC} \rightarrow ^3\text{MLCT}$	3.2×10^6	7.56	0.03	20.06
	$^3\text{MLCT} \rightarrow ^3\text{LC}$	3.3×10^6	7.56	-0.03	20.06
	$^1\text{MLCT} \rightarrow ^3\text{MLCT}$	3.2×10^5	0.83	-9.58	28.94
Ir-1	$^3\text{MLCT} \rightarrow ^1\text{MLCT}$	2.0×10^{-2}	0.83	9.58	28.94
	$^1\text{MLCT} \rightarrow ^3\text{LC}$	9.0×10^{10}	16.90	-0.17	2.49
	$^3\text{LC} \rightarrow ^1\text{MLCT}$	6.7×10^{10}	16.90	0.17	2.49
	$^3\text{LC} \rightarrow ^3\text{MLCT}$	3.4×10^{11}	31.98	-9.76	11.33
	$^3\text{MLCT} \rightarrow ^3\text{LC}$	2.3×10^4	31.98	9.76	11.33
	$^1\text{MLCT} \rightarrow ^3\text{MLCT}$	6.5×10^1	0.11	-9.93	42.41
	$^3\text{MLCT} \rightarrow ^1\text{MLCT}$	1.5×10^{-8}	0.11	9.93	42.41

^a The unit of the rates is s^{-1} . The units of electronic coupling and energy are cm^{-1} and kcal mol^{-1} , respectively.

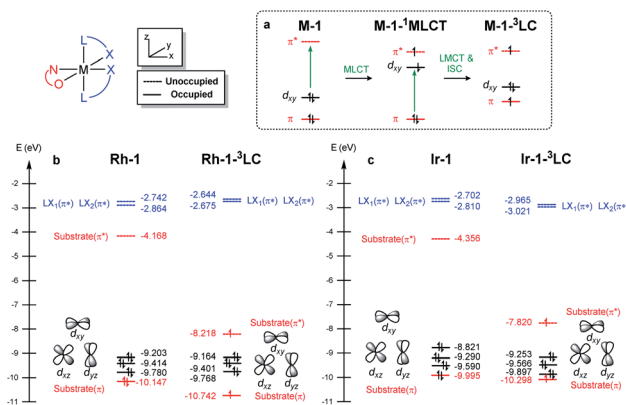


Fig. 7 (a) Electronic excitations and (b) frontier molecular orbitals of **M-1** computed at the B3LYP-D3/cc-pVTZ(-f)/LACV3P** level of theory.

process, bringing back the singlet MLCT state. We may conceptualize the ISC and RISC processes comprising two independent processes: (i) the transition of an electron between the substrate π orbital to the singly occupied d_{xy} orbital (ligand-to-metal charge transfer (LMCT) and metal-to-ligand charge transfer (MLCT)) and (ii) the spin-flip to convert between singlet and triplet states. Our computational results suggested that the spin-orbit coupling (V_{soc}) for both Rh and Ir systems are comparable (Table 2). In this regard, the energy gaps of π - d_{xy} and π - π^* play important roles in the ISC and RISC processes, as the energy gaps determine the electronic energies of $^1\text{MLCT}$ and ^3LC states.

Fig. 7b and c summarize the energy levels of frontier molecular orbitals that participate in the photoactivated electronic transitions. We investigated d - π^* gaps of the optimized singlet ground state geometries to approximate the gap of the singlet MLCT states. The computed d - π^* gap of **Rh-1** was 2.46 eV and that of **Ir-1** was 2.00 eV, in qualitative agreement with the energetics of **M-1**- $^1\text{MLCT}$ states measured from the fluorescence spectra: the excitation energy of **Rh-1** was 2.86 eV being 0.72 eV higher than that of **Ir-1** (Fig. 4). The observed difference in d_{xy} - π^* gaps can be explained by the pronounced relativistic contraction of Ir that elevates the occupied d -levels of Ir and lowers the unoccupied π^* -level of the ligand.⁴⁴ The orbital energy difference between π - π^* gaps of **Rh-1** and **Ir-1** is comparable: 3.50 eV and 3.66 eV, respectively. This is in line with the finding that the adiabatic energies of **M-1**- ^3LC states of both systems are comparable (Fig. 6). In short, our analysis of the FMOs highlighted that the labile characteristic of **Ir-1** compared to **Rh-1** originates from the higher d_{xy} of the Ir center than that of Rh.

The mechanism of triplet energy transfer of the Ir photocatalyst

We further investigated substrate-chelated Ir(III) complex **Ir-1** with regard to its stability and reactivity to expand our insights into the mechanism of the Ir-catalyzed $[2 + 2]$ photocycloaddition. The circular dichroism (CD) spectra of **Ir-1** before and after irradiation with blue LEDs for 20 h under an inert gas atmosphere gave us information about the chiral stability under reaction conditions. In acetone as solvent, which we used as the main solvent for the $[2 + 2]$ photocycloadditions, we were able to observe a slow racemization of the complex indicated by different heights of maxima and minima (Fig. S9†). The conjecture of decomposition upon irradiation as the reason for the distinct spectra was omitted due to the similarity of the curves as well as $^1\text{H-NMR}$ measurements showing no major decomposition upon illumination in acetone (Fig. S9†). To further evaluate the possible photoracemization of **IrS** during a $[2 + 2]$ photocycloaddition, we investigated the chiral stability of **IrS** in the presence of a large excess of 2-acyl imidazole substrate **1**, by measuring CD spectra before and after illumination (Fig. S11†). The spectrum prior to irradiation strongly resembles the spectrum of **A-Ir-1** indicating that the substrate-coordinated complex is formed instantaneously. Importantly, the CD spectrum after exposure to light looks almost identical,

implying that an excess of substrate protects the stereogenic center of the complex against racemization. Therefore, a racemization of the iridium catalyst during photolysis can be excluded as a reason for the formation of the racemic $[2 + 2]$ photocycloaddition product.

Ruling out the possibility of the direct photocatalysis pathway of **Ir-1** prompted us to search for an alternative mechanistic pathway that can explain the racemic product formation from the **IrS** system. Mechanisms that include the bimolecular activation between substrate **1** and the diene without the chiral environment of **IrS** may lead to the racemic products. This includes two modes of activation of **1**, either a redox pathway to furnish an open shell intermediate or an outer-sphere energy transfer pathway to access the triplet state of the substrate. We precluded the photoredox pathway, as the measured redox potentials of the metal catalysts and the catalyst-substrate bound complexes are not strong enough to oxidize/reduce the substrate **1** (*vide supra*). After much effort, we concluded that the triplet energy transfer pathway is the most likely mechanism for the Ir-photocatalysis that was substantiated by our experimental findings. First, we found that **Ir-1** catalyzes the photocycloaddition of methyl cinnamate. As shown in Table 2, **Ir-1** catalyzes the $[2 + 2]$ photocycloaddition of (*E*)-methyl cinnamate (entries 1 and 3). No conversion was observed when the catalyst was omitted (entries 2 and 4). Since methyl cinnamate cannot coordinate to iridium, this finding strongly suggests that **Ir-1** serves as a triplet photosensitizer and activates the cinnamate substrates to access the triplet excited state manifold. Of note, Reiser and coworkers recently reported that bis-cyclometalated iridium complexes can serve as effective photocatalysts for the photosensitized $[2 + 2]$ cycloaddition of cinnamate esters.⁴⁵

The triplet energy transfer mechanism is outlined in Fig. 8. First, the ground state photocatalyst ^1PC absorbs visible light to become the singlet excited state $^1\text{PC}^*$, followed by intersystem crossing that yields the photocatalyst in its triplet excited state ^3PC . Then the ^3PC transfers its energy to the enone substrate **1** to generate the triplet state species $^3\text{1}$, which undergoes alkene addition to form a 1,4-biradical intermediate $^3\text{2}$. Finally, ISC into the singlet state occurs yielding cyclobutane species $^1\text{3}$.

Fig. 9 shows the computed energy profile following the triplet sensitization mechanism. Once the photocatalyst harvests light energy, an outer-sphere energy transfer can promote the enone substrate **1** to become a reactive triplet state species $^3\text{1}$. We showed that **Ir-1** may act as a photosensitizer in

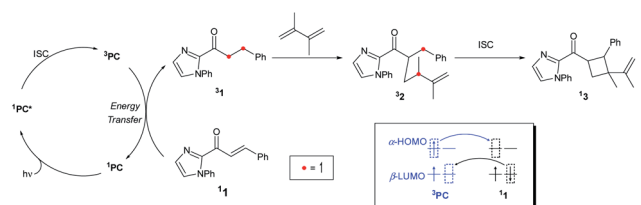


Fig. 8 Triplet energy transfer mechanism for the $[2 + 2]$ cycloaddition reaction.



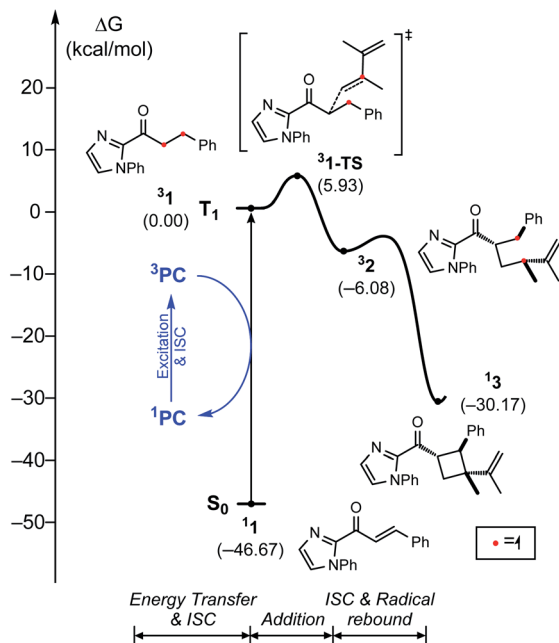


Fig. 9 Energy profile with triplet energy transfer mechanism.

Table 3 [2 + 2] Photocycloaddition of methyl cinnamate catalyzed by Ir-1^a

Entry	Solvent	t (h)	Conversion ^b (%)	5 ^b (%)	6 + 7 ^{b,c} (%)
1	Acetone	20	70.4	60.1	10.3
2 ^d	Acetone	20	0	—	—
3	DMF	44	50.4	16.4	34.0
4 ^d	DMF	44	0	—	—

^a Conditions: a flame dried Schlenk flask was charged with methyl cinnamate (0.10 mmol) and Ir-1 (2 mol%) under an inert gas atmosphere. The indicated solvent (0.5 mL) was added and the resulting mixture was degassed thoroughly *via* freeze-pump-thaw for three cycles. Subsequently, the reaction mixture was stirred for the indicated time under an inert gas atmosphere in front of blue LEDs (24 W, 10 cm distance). ^b Conversion and yield determined by ¹H-NMR of the crude mixture after solvent evaporation. ^c Combined ¹H-NMR yields of the two diastereomers δ - and β -truxinate. ^d Reaction was performed without a catalyst.

solution (Table 3), whose triplet excitation energy is 1.63 eV. IrS may be another candidate as the photosensitizer as it shows a sufficient triplet energy of 2.35 eV (Fig. 4). Once generated, **31** will react with a diene to yield 1,4-diradical species **32** *via* transition state **31-TS** with the activation barrier of 5.9 kcal mol⁻¹. According to our photophysical measurements, the **31** has a long lifetime of 11 ms, that can withstand the low activation barrier for the first C–C bond formation step (**31-TS**). The last phase of the reaction, followed by the radical rebound step, that produces cyclobutane species **13** is endothermic by –30.2 kcal mol⁻¹ relative to **31**. From the calculated results of the triplet energy transfer pathway, we believe that the triplet sensitization mechanism could be key to explaining the observed racemates by Ir photocatalysis.

Conclusions

Intrinsic mechanistic differences investigated by combined computational and experimental studies shed light on the roles of Rh or Ir catalysts in a visible-light induced [2 + 2] photocycloaddition reaction. While a chiral-at-rhodium catalyst showed a high level of enantioselectivity, the use of a chiral iridium catalyst led to racemate formation. Our integrated approach led us to conclude that Rh photocatalysis activates the substrate *via* direct photoexcitation of the catalyst–substrate complex **Rh-1** traversing the ³LC state upon excitation. In contrast, the analogous **Ir-1** showed a ³MLCT characteristic, which is a non-emissive state, and cannot engage asymmetric induction due to its short lifetime. Iridium mediated photocatalysis, however, utilizes an alternative pathway, in which an outer-sphere triplet energy transfer is the underlying mechanism of the photocatalysis. The lower d– π^* energy gap of **Ir-1** compared to the **Rh-1** can explain the origin of this striking difference, thereby highlighting the importance of choosing the metal center in photocatalytic systems. We anticipate that the described mechanistic aspects highlighting the differences between isolobal rhodium and iridium catalysts will provide additional insights into designing new photocatalytic systems.

Data availability

Further details of experimental procedure, photophysical measurements, electrochemical studies, computational methods, relevant coordinates, and energies are available in the ESI.

Author contributions

PC, JP, and MHB conceived the project. HJ and MH, and JP carried out the computational analysis in consultation with MHB; MM and MV performed the photophysical measurements in consultation with PC; PSS, XH, and MH carried out the synthesis and characterization in consultation with EM. MHB, JP, HJ, and MH wrote the manuscript, and PC and EM contributed to writing the manuscript.

Conflicts of interest

There are no conflicts to declare.

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