Nitrogen radical-triggered trifunctionalizing *ipso*-spirocyclization of unactivated alkenes with vinyl azides towards new spiroaminal frameworks

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**ABSTRACT:** Radical-mediated spirocyclization is a powerful and efficient tool to forge complex spirocyclic frameworks. Radical-mediated non-dearomative double-cyclization of linear precursors for the concomitant construction of both rings is more challenging but highly attractive. Herein, we report the first example of non-dearomative trifunctionalizing *ipso*-spirocyclization of unactivated alkenes through photoredox-catalyzed, nitrogen radical-triggered cyclization-trapping-translocation-cyclization cascade, providing a single-step modular access to architecturally new and fascinating spiroaminal frameworks through simultaneous formation of one C−C bond and two geminal C−N bonds. The developed protocol utilizes not only internal or terminal olefinic oxime esters but also olefinic amides as nitrogen radical precursors, and features broad substrate scope, varied functional group compatibility, easy scalability, and potential for product derivatization and late-stage functionalization of biologically active molecule. Importantly, the mechanistic studies including DFT calculations indicate that such photocatalytic trifunctionalizing *ipso*-spirocyclization undergoes a radical relay cascade of intramolecular 5-*exo-trig* cyclization, intermolecular radical trapping, 1,5-hydrogen
atom transfer, and sequential $5$-endo-trig cyclization, which would open up a new reaction mode of alkenes.

Spirocyclic skeletons widely occur in a plethora of naturally occurring products with broad structural diversities and diverse biological activities$^{1-4}$, and are increasingly being incorporated into drug candidates$^{5,6}$ (Fig. 1a). Furthermore, the inherent three-dimensional (3D) conformational constraint character of the spirocyclic skeletons enables them as one of the excellent ligand$^{7-10}$ and organocatalyst$^{11,12}$ frameworks in the field of asymmetric catalysis. In this context, an elegant radical-mediated spirocyclization for the construction of the spirocyclic compounds, including dearomative$^{13}$ and non-dearomative$^{14}$ strategies, has been developed rapidly in the past decade. To the best of our knowledge, the existing approaches for the radical-mediated spirocyclization are solely based upon the elaboration of aromatic or aliphatic monocyclic precursors on which the second ring is newly appended, and compared with the former dearomative strategies, the latter non-dearomative strategies could provide structurally distinct spirocyclic compounds with varied functionality (Fig. 1b). However, radical-mediated non-dearomative double-cyclization of linear precursors for the concomitant construction of both rings has remained unexploited to date. Thus, the development of novel spirocyclic skeletons and evermore-effective non-dearomative spirocyclization strategies using different classes of linear starting materials are highly desirable but challenging.

On the other hand, alkenes are energetic and versatile synthons that engage in diverse organic transformations because of their abundance and rich reactivity. In recent years, the difunctionalization of alkenes is one of the most important and valuable strategies in organic synthesis, since it allows the rapid, efficient, and synergistic incorporation of two functional groups onto one C–C double bond in an atom- and step-economic fashion for enhancing the molecular complexity and diversity. Among them, the 1,2-difunctionalization of alkenes, including intermolecular and intramolecular reactions, was extensively developed as a powerful tool for the creation of two new vicinal chemical bonds$^{15-20}$. Compared with the vigorous increasing achievements of 1,2-difunctionalization, the 1,1-difunctionalization of alkenes to
access geminal difunctionalized alkanes is still underdeveloped\textsuperscript{21}; for example, the scope of alkene geminal diamination is limited to activated alkenes as well as ethylenediamine derivatives as nitrogen sources\textsuperscript{22-24}. Notwithstanding the remarkable advances achieved toward the difunctionalization of alkenes, only sparsely have the more challenging trifunctionalization of alkenes been exploited, which significantly restricted their application for the assembly of diversely multifunctionalized molecules (Fig. 1c). The reason might be mainly attributed to the lack of an effective approach to realize the trifunctionalization. For the purpose of providing structurally diverse and complex functional molecule libraries derived from privileged scaffolds (particularly spirocyclic skeletons) to meet the ever-increasing demand for the development of new drugs, ligands and organocatalysts, the exploration of novel, straightforward, and efficient approaches for the trifunctionalization of alkenes is of special interest and great significance.

Successfully merging the difunctionalization of alkenes with the selective functionalization of C(sp\textsuperscript{3})–H bonds might be one of the reliable and effective approaches to reach the formidable trifunctionalization of alkenes. Accordingly, the strategies employed for such trifunctionalization can be largely divided into two categories: 1) very few “innate trifunctionalization”, aliphatic C(sp\textsuperscript{3})–H functionalization of the newly generated difunctionalized intermediates, such as β-keto sulfones derived from activated alkenes including vinyl azides\textsuperscript{25} and styrenes\textsuperscript{26}, is solely based on the inherent reactivity of aliphatic C(sp\textsuperscript{3})–H bonds; 2) unexploited “guided trifunctionalization”, selective functionalization of inert aliphatic C(sp\textsuperscript{3})–H bonds in the resulting difunctionalized intermediates is guided by the external reagents or directing groups armed with functionalities, such as those that can be transformed into open-shell intermediates. In recent years, the iminyl radical-directed 1,5-hydrogen atom transfer (HAT) process provides an exceptional platform for the site-selective activation and functionalization of inert C(sp\textsuperscript{3})–H bonds\textsuperscript{27-30}. Meanwhile, vinyl azides have emerged as an attractive iminyl radical source generated through an intermolecular radical addition followed by the extrusion of N\textsubscript{2}. Up to now, several elegant examples, wherein the iminyl radicals derived from vinyl azides could trigger aliphatic C(sp\textsuperscript{3})–H functionalization involving 1,5-HAT process, have been reported by
Nevado$^{31}$, Guo$^{32}$, Bolm$^{33}$, and Liu group$^{34}$. Inspired by elegant work on the rich reactivities of nitrogen radical precursors$^{35-40}$, we envisioned whether the tandem radical relay strategy of the initial radical difunctionalization of alkenes and follow-up C(sp$^3$)–H functionalization involving iminyl radical-directed 1,5-HAT process could be applicable to the guided trifunctionalization of unactivated alkenes. Herein, we report for the first example of non-dearomative trifunctionalizing ipso-spirocyclization of unactivated alkenes with vinyl azides through photoredox-catalyzed, nitrogen radical-triggered cyclization-trapping-translocation-cyclization (CTTC) cascade (Fig. 1d). More importantly, one C–C bond and two geminal C–N bonds could be synergistically constructed across double bonds under this photocatalytic system to enable a straightforward single-step modular synthesis of architecturally new and intriguing spiroaminals including spirobipyrrolines and spiropyrrrole lactams.

**a** Representative examples of spirocyclic compounds: natural products, bioactive molecules, and ligands

(-)-Spirocyclic compounds: natural products, bioactive molecules, and ligands

**b** The existing approaches for the radical-mediated spirocyclization

**c** Alkene multifunctionalization

**d** This work: nitrogen radical-triggered trifunctionalizing ipso-spirocyclization of unactivated alkenes with vinyl azides

* Architecturally new spiroaminal frameworks * Non-dearomative spirocyclization of linear precursors * Alkene trifunctionalization
* Unactivated alkene: 1,2,3-tri-radical synthon * One C-C bond and two geminal C-N bonds in 1-step * 52 examples, up to 80% yield
* Broad substrate scope * Good functional group compatibility * Easy scalability
* Product derivatization * Late-stage functionalization * DFT calculation
**Fig. 1** a Representative examples of spirocyclic compounds: natural products, bioactive molecules, and ligands. b The existing approaches for the radical-mediated spirocyclization. c Different approaches for the multifunctionalization of alkenes. d This work: nitrogen radical-triggered trifunctionalizing *ipso*-spirocyclization of unactivated alkenes with vinyl azides.

**Results**

**Reaction optimization.** To test the described hypothesis, iminyl radical precursor *O*-4-trifluoromethylbenzoyl oxime ester 1a and vinyl azide 2a were initially chosen as the model substrates to explore the reaction conditions (Table 1). Unless otherwise noted, all the experiments were performed in the presence of *fac*-Ir(ppy)₃ as photocatalyst, K₂CO₃ as base, and DMSO as solvent under argon atmosphere upon irradiation of 30 W blue light-emitting diodes (LEDs) at room temperature. To our delight, after systematic optimization of the reaction parameters, including nitrogen radical precursors, photocatalyst, solvent, and base, the desired product spirobi[pyrroline] 3aa was obtained in 71% isolated yield. Based on the more oxidizing ability of electron-deficient *O*-substituted oxime esters, the utilization of other leaving groups (LG) instead of OBzCF₃ gave no better results (entries 2 and 3). Among the catalysts examined, *fac*-Ir(ppy)₃ was found to be more effective than other photocatalysts (entries 4 and 5). Furthermore, screening of solvents demonstrated that other solvents delivered no more significant improvement than that of DMSO (entries 6–8). It was worth mentioning that the yields of the desired product declined slightly when the reaction was conducted without the addition of base or argon protection (entries 9 and 10). As anticipated, no desired product could be detected without either light or photocatalyst, indicating the indispensability of both light and photocatalyst in such photocatalytic CTTC cascade (entry 11).

| Table 1 Optimization of reaction conditions⁴ |
Table 2. Variation from the standard conditions and yields (%)

| Entry | Variation from the standard conditions | Yield (%)<sup>b</sup> |
|-------|----------------------------------------|----------------------|
| 1     | none                                   | 76 (71)              |
| 2     | LG = C₆F₅CO₂⁻                         | 10                   |
| 3     | LG = 2,4-NO₂-C₆H₄O⁻                     | n.d.                 |
| 4     | PC-2 instead of fac-Ir(ppy)<sub>3</sub> | 30                   |
| 5     | PC-3 instead of fac-Ir(ppy)<sub>3</sub> | n.d.                 |
| 6     | DMF instead of DMSO                    | 34                   |
| 7     | MeCN instead of DMSO                   | 46                   |
| 8     | DME or MeOH instead of DMSO            | n.d.                 |
| 9     | no base                                | 54                   |
| 10    | no argon protection                    | 53                   |
| 11    | no light or photocatalyst              | n.d.                 |

<sup>a</sup>Reaction conditions: 1a (0.1 mmol), 2a (0.25 mmol), fac-Ir(ppy)<sub>3</sub> (2 mol %), K₂CO₃ (0.15 mmol), DMSO (1 mL), 30 W blue LEDs, argon atmosphere, r.t., 6 h, in a sealed tube. n.d. = no detection. LG = p-CF₃-C₆H₄CO₂⁻ (OBz<sub>CF₃</sub>).<sup>b</sup>Yields were determined by <sup>1</sup>H NMR using dibromomethane as an internal standard. Isolated yields in parentheses.

**Substrate scope on the vinyl azides.** With the optimized reaction conditions in hand, we next sought to investigate the substrate scope on the trifunctionalizing *ipso*-spirocyclization enabled by the iminyl radical-triggered CTTC cascade (Table 2). First, the generality of this *ipso*-spirocyclization with regard to vinyl azides was examined with olefinic oxime ester 1a as the iminyl radical precursor. A variety of vinyl azides with different electronic groups at the *para*- (3aa–3al), *meta*- (3am–3ap), or *ortho*-position (3aq) of the phenyl rings reacted smoothly with 1a to afford the corresponding spirobi[pyrrolines] in acceptable yields. Gratifyingly, this photocatalytic
trifunctionalizing *ipso*-spirocyclization tolerated a broad range of diverse functionalities such as alkyl (3aa–3ae, and 3am), phenyl (3ag), methoxyl (3ah, 3an, and 3aq), and trifluoromethyl (3ai), as well as halogen (3aj–3al, 3ao, and 3ap) which could be employed as a handle for further derivatization. Furthermore, heteroaryl-substituted vinyl azides including thiophene (3ar) and pyridine (3as) displayed good reactivity in our protocol. Notably, alkyl-substituted vinyl azide was also proved to be suitable substrate for this spirocyclization and provided 44% yield of the expected spirobi[pyrroline] 3at.

**Table 2 Substrate scope on the vinyl azides.**

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*a*
Reaction conditions: 1 (0.2 mmol), 2 (0.5 mmol), fac-Ir(ppy)$_3$ (2 mol %), K$_2$CO$_3$ (0.3 mmol), DMSO (2 mL), 30 W blue LEDs, argon atmosphere, r.t., 6 h, in a sealed tube; isolated yields based on 1 after the chromatographic purification.
**Substrate scope on the olefinic oxime esters.** Having demonstrated the varied functional group compatibility in the vinyl azide coupling partners, we subsequently set out to explore the scope of olefinic oxime esters 1 with 2a as the coupling partner. As illustrated in Table 3, a panel of aromatic oxime esters bearing electronic variation on the phen rings were also well-tolerated in this transformation, affording the corresponding spirobi[pyrrolines] 3ba–3ga in moderate to good yields. Furthermore, 2-naphthyl- (3ha) and heteroaryl-substituted oxime esters (3ia) were also subjected to the present protocol. Additionally, aromatic oxime esters with alkyl chain units at the α position were used as the substrates in this protocol and smoothly transformed into the corresponding spirobi[pyrrolines] 3ja and 3ka in 45% and 58% isolated yields, respectively. Notably, the structure of 3ka was unambiguously confirmed by X-ray crystallographic analysis (CCDC 2061499). Importantly, in addition to 1,2,2-trisubstituted olefinic oxime esters, 1,2-disubstituted (3la–3na) and 1-substituted olefinic oxime esters (3oa) could also be employed as competent substrates in this transformation to provide the desired products.

**Table 3 Substrate scope on the olefinic oxime esters.**

a
Substrate scope on the amidyl radical-triggered trifunctionalizing ipso-spirocyclization. Encouraged by the success of the unprecedented trifunctionalizing ipso-spirocyclization enabled by the iminyl radical-triggered CTTC cascade, we speculated that the high electrophilic amidyl radical\textsuperscript{41,42} could also trigger such CTTC cascade to achieve the desired trifunctionalizing ipso-spirocyclization of
unactivated alkenes for rapid construction of another new spiroaminal framework. Excitingly, upon the facile screening of parameters, amidyl radical precursor olefinic amides were indeed capable partners and readily underwent such photocatalytic trifunctionalizing \textit{ipso}-spirocyclization (Table 4). Similarly, vinyl azides with various electron-donating or electron-withdrawing groups reacted smoothly with 4a to furnish the structurally novel spiropyrroline lactams in satisfactory yields. Olefinic amides with a hindered \( N \)-Cy group (5ba), \( N \)-Ph group (5ca), as well as a removable \( N \)-Bn group (5da) could successfully participate in this trifunctionalizing \textit{ipso}-spirocyclization. Furthermore, olefinic amide with alkyl chain unit at the \( \alpha \) position could be applied as well to deliver the corresponding product 5ea. In addition, the proposed approach could address 6-\textit{exo-trig} cyclization of unactivated alkene to generate 6-membered ring lactam 5fa. Interestingly, \( N \)-protected amine was compatible with the photoredox catalysis and provided the desired spiropyrroline pyrrolidine 5ga in moderate yield.

\textbf{Table 4 Substrate scope on the amidyl radical-triggered trifunctionalizing \textit{ipso}-spirocyclization.}^a
Reaction conditions: 4 (0.2 mmol), 2 (0.5 mmol), fac-Ir(ppy)$_3$ (2 mol %), K$_2$CO$_3$ (0.3 mmol), DMSO (2 mL), 30 W blue LEDs, argon atmosphere, r.t., 6 h, in a sealed tube; isolated yields based on 4 after the chromatographic purification.

Products transformation. To demonstrate the potential application of the protocol, a gram-scale reaction containing 3.0 mmol of oxime ester 1a was performed under the standard conditions to afford the target product 3aa with a satisfactory isolated yield, demonstrating the synthetic practicality and scalability of our photocatalytic trifunctionalizing ipso-spirocyclization (Fig. 2a). Moreover, the synthetic utility of the method was further confirmed by facile derivatization of the formed spirobi[pyrroline] 3aa to yield structurally diverse and valuable molecules (Fig. 2b). Rather surprisingly,
in the presence of protonic acid or trifluoroacetic anhydride, 3aa can be smoothly transformed into the aromatized 1H-pyrrole derivatives (6 and 7) in 75% and 68% yields, respectively. Moreover, [3 + 2] cycloaddition reaction between 3aa and N-hydroxybenzimidoyl chloride allowed access to the 1,2,4-oxadiazoline-fused sipropyrrrole 8. When spirobi[pyrrole] 3aa was treated with N-chlorosuccinimide (NCS) at 80 °C for 24 h, the multi-halogen-substituted spiroaminal 9 was obtained with excellent isolated yield. Delightfully, estrone-derived vinyl azide 2w readily engaged in such photocatalytic transformation with olefinic oxime ester 1a to give the structurally more complex and intriguing compound 3aw, demonstrating the potential application of this methodology (Fig. 2c). Additionally, when 1,1-disubstituted terminal olefinic oxime esters were subjected to the reaction system, the desired 1,5-HAT process did not proceed smoothly and thus no corresponding spirobi[pyrrole] was detected. Interestingly, using vinyl azide 2-azidoallyl diphenylmethyl ether 2x as the coupling partner, the second iminyl radical derived from vinyl azide underwent intramolecular 1,5-HAT (from nitrogen to benzyl carbon atom) and further cyclization to afford a panel of the oxazoline-based pyrrolines 10px–10ux (Fig. 2d). Notably, terminal olefinic oxime ester 1o could also be converted into the expected oxazoline-based pyrroline 10ox albeit with somewhat low yield.
Fig. 2 Gram-scale synthesis and synthetic applications of the reaction.
Mechanistic studies. To rationalize the mechanism of this photocatalytic trifunctionalizing ipso-spirocyclization reaction, several control experiments were carried out. As shown in Scheme 3, upon the addition of electron-transfer scavenger p-dinitrobenzene (DNB) or radical scavenger 2,2,6,6-tetramethyl-piperidinyloxyl (TEMPO) into the model reaction, the formation of the desired product 3aa was completely inhibited (Fig. 3a). And the corresponding 1a-derived TEMPO-trapped adduct was detected through LC−HRMS analysis. When the model reaction was performed in the presence of quintessential radical-trapping reagent 1,1-diphenylethylene (DPE), the yield of 3aa declined slightly from 71% to 43%, and the radical-trapping products (3aa' and 3aa'') were detected through LC−HRMS analysis (Fig. 3a). Subsequently, in the absence of vinyl azide 2a, the hydroimination product 1a' and olefination product 1a'' were obtained in 23% and 32% yield, respectively (Fig. 3b). These results indicate that not only a single electron transfer (SET)/radical process but also the carbon-radical intermediates generated through iminyl radical-triggered intramolecular cyclization might be involved in the present transformation. Additionally, no desired product 3aa was detected when vinyl azide 2a was replaced with 3-phenyl-2H-azirine 2a' generated from the denitrogenative decomposition of vinyl azide under the standard conditions, suggesting that 2H-azirine intermediates might be ruled out in this transformation (Fig. 3c).
On the basis of the above-mentioned results and preceding reports\textsuperscript{43-45}, a photoinduced electron transfer (PET) mechanism was proposed for the trifunctionalizing \textit{ipsos}-spirocyclization. Density functional theory (DFT) calculations were also carried out to further validate the proposed reaction mechanism and the factors governing chemoselectivity at the M06-2X/def2-TZVP/SMD(DMSO)//M06-2X/def2-SVP level of theory\textsuperscript{46-52}. As illustrated in Fig. 4, the oxime ester 1a ($E_{1/2}^{\text{red}} = -1.61$ V vs. SCE, see the Supporting
Information) can undergo exergonic single-electron transfer (SET) process (42.4 kcal mol\(^{-1}\)) with the excited \(*\text{Ir}^{\text{III}} [E_{1/2}^{\text{red}} (\text{Ir}^{\text{IV}}/*\text{Ir}^{\text{III}}) = -1.73 \text{ V vs. SCE}]\) to deliver the first iminyl radical A, which could go through thermodynamically favorable intramolecular 5-\text{exo-trig} cyclization to generate the first pyrroline species B containing an alkyl radical. Alternatively, the 6-\text{endo-trig} cyclization pathway is kinetically less favored (\text{TS}_{6-\text{endo-trig}} = 18.66 \text{ kcal mol}^{-1} \text{ vs. TS}_{5-\text{exo-trig}} = 11.61 \text{ kcal mol}^{-1}). Two possible pathways were expected for the following reaction pathway of B, including intermolecular radical trapping with vinyl azide 2f (black line) and hydrogen atom abstraction with the C-H of solvent dimethyl sulfoxide (red line). DFT calculations determined that the activation free energy of the addition of carbon radical intermediate B to the C=C of 2f was only 5.98 kcal mol\(^{-1}\), which was much lower than that who was found for the pathway via \text{TS} _{\text{B_DMSO}} (25.55 \text{ kcal mol}^{-1}). This radical trapping would create a new C-C bond in α-azide benzyl radical C, which is an exergonic process (23.74 kcal mol\(^{-1}\)). Next, the α-azide benzyl radical C could rapidly release a \text{N}_2 molecule via an almost barrierless transition state (\text{TS} _{\text{C}}, 0.19 \text{ kcal mol}^{-1}) after conformation adjustment to the isomer C(2) and thus result in the formation of the second imine radical D. The iminyl radical D underwent a radical translocation process of 1,5-HAT via transition state \text{TS} _{\text{D_E}} with a moderate energy barrier of 11.05 kcal mol\(^{-1}\) to generate a more stabilized α-amino carbon radical E(1). Then, an intramolecular 5-\text{endo-trig} cyclization with an imine moiety led to the second pyrroline species F containing a benzyl radical with the highest energy barrier 18.95 kcal mol\(^{-1}\) in the whole trifunctionalizing ipso-spirocyclization reaction, indicating that the radical cyclization process is the rate-limiting step in the reaction. Subsequently, the intermediate F (calculated \(E_{1/2}^{\text{ox}} = -1.15 \text{ V vs. SCE}\)) can be easily oxidized by Ir^{\text{IV}} [\(E_{1/2}^{\text{red}} (\text{Ir}^{\text{IV}}/\text{Ir}^{\text{III}}) = 0.77 \text{ V vs. SCE}]\) to the carbon cation G, which could resonate to the protonated product H. Finally, the target spirobi[pyrroline] 3af could be obtained after the deprotonation of compound H with the help of \(p\)-trifluoromethylbenzoate anion or base. Although the reaction pathway of the oxidation of radical intermediate E (calculated \(E_{1/2}^{\text{ox}} = -0.15 \text{ V vs. SCE}\)) to intermediate E_\text{cation} by oxidative Ir^{\text{IV}} complex could be envisioned, the efforts aiming to locate the transition state of
intramolecular ionic cyclization of intermediate E\textit{cation} leading to intermediate H has been failed.

\textbf{a} Proposed mechanism for the nitrogen radical-triggered trifunctionalizing ipso-spirocyclization

\textbf{b} Free energy profiles for the the nitrogen radical-triggered trifunctionalizing ipso-spirocyclization

Fig. 4 Proposed mechanism and free energy profiles for the nitrogen radical-triggered trifunctionalizing ipso-spirocyclization of unactivated alkenes. The values have been given in the unit of kcal mol\textsuperscript{-1} and represent the relative free energies calculated using M06-2X method in DMSO solvent. Note: Two different energy values for one structure refer to conformer changes.

Discussion

In conclusion, we successfully developed a novel, selective, and efficient strategy to realize trifunctionalizing ipso-spirocyclization of unactivated internal and terminal alkenes by photoredox-catalyzed, nitrogen radical-triggered relay cascade involving
1,5-HAT, which would open up a new reaction mode of alkenes. Not only iminyl radical precursor olefinic oxime esters but also amidyl radical precursor olefinic amides are subjected to the developed photocatalytic protocol, allowing a rapid and efficient entry to a diverse array of architecturally new and intriguing spiroaminal frameworks through the simultaneous formation of one C–C bond and two geminal C–N bonds. Furthermore, the resulting spiroaminals can be diversely functionalized through various facile transformations, such as aromatization into 2,5-disubstituted pyrroles, [3 + 2] cyclization, and multi-halogenation reaction. And the applicability of such protocol is also demonstrated by late-stage functionalization of bioactive estro derivative. In addition, the mechanistic studies indicate that this transformation undergoes a nitrogen radical-triggered cyclization-trapping-translocation-cyclization (CTTC) cascade, namely intramolecular 5-exo-trig cyclization, intermolecular radical trapping, 1,5-HAT, and 5-endo-trig cyclization. Further application of this strategy is currently underway in our laboratory.

Methods

General procedure. An oven-dried Schlenk tube (10 mL) equipped with a stirring bar was charged with olefinic oxime ester 1 (0.2 mmol, 1.0 equiv.), K₂CO₃ (0.3 mmol, 1.5 equiv.), and fac-Ir(ppy)₃ (2 mol%). The tube was connected to a vacuum line where it was evacuated and back-filled with Ar for three times. Next, vinyl azide 2 (0.5 mmol, 2.5 equiv.) and DMSO (2 mL) were added under Ar flow. The tube was placed approximately 2 cm from 30 W blue LEDs, and then stirred at room temperature for 6 h. After completion, the reaction was quenched with H₂O (4.5 mL) and extracted with EtOAc. The organic layer was washed with saturated brine and dried over Na₂SO₄. The solvent was removed under reduced pressure, and the resulting residue was purified by column chromatography on silica gel to afford the desired product 3. Note: for olefinic amides 4 (MeCN was used instead of DMSO), the reaction mixture was filtered after completion, concentrated under reduced pressure, and purified by column chromatography on silica gel to afford the desired product 5.

Additional Information

Supplemental Information can be found with this article online. The crystallography data have been deposited at the Cambridge Crystallographic Data Center (CCDC) under accession number CCDC: 2061499 (methyl ester of 3ka) and can be obtained free of charge from
Author contributions
D.Y. and X.D. conceived and designed the study, and wrote the paper. Z.Y.Q., Z.J.Z., L.Y., D.Z. performed the experiments and mechanistic studies. Q.F. and S.P.W. performed the DFT calculations. All authors contributed to the analysis and interpretation of the data.

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