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Introduction

Using abundant bulk chemicals to synthesize fine chemicals through a sustainable catalytic strategy is significant in the field of synthetic chemistry.¹ Regarding the efficient activation of inert bulk chemicals, general considerations or solutions are to increase the energy input such as resorting to high temperatures,² noble metal catalysts,³ strong or special oxidants/reductants/acids/ bases,⁴ UV light,⁵ *etc.*⁶ Due to the urgent demand for green and sustainable chemistry,⁷ balancing the requirement of reducing the energy input/consumption and the growing interest of chemists in the activation of inert chemicals is what chemists should imminently address (Fig. 1a).

The nontoxicity and abundance of iron endow iron-catalyzed synthetic chemistry with strong sustainability and potential application prospects in bio-medicine and industrial manufacturing fields.⁸ The low cost and easily available iron(III) salts have been playing versatile roles in the synthetic chemistry community owing to their remarkable Lewis acidity⁹ and redox reactivity. With respect to the redox character of iron(III), it has exhibited the ability to oxidise some active radical species¹⁰ or

Dual ligand-enabled iron and halogen-containing carboxylate-based photocatalysis for chloro/fluoro-polyhaloalkylation of alkenes†

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Herein, we demonstrate a practical dual ligand-enabled iron photocatalysis paradigm—converting all kinds of halogen-containing carboxylates ($C_n X_m COO^-$, X: F, Cl, Br) into $C_n X_m$ radicals for the valuable chloro/ fluoro-polyhaloalkylation of non-activated alkenes with easily available trichloroacetonitrile/Selectfluor as the electrophilic halogenation reagent. The modular *in situ* assembly of the effective iron and $C_n X_m COO^-$ -based light-harvesting species using the two ligands—OMe/CF₃-substituted bipyridine and acetonitrile/trichloroacetonitrile is evidenced by detailed mechanistic studies. The late-stage modification, low loading amount of iron (TON: 257) and feasible gram-scale synthesis show the utility of this protocol. We thus anticipate that the dual ligand-enabled iron photocatalysis paradigm may facilitate activation and transformation of inert bulk chemicals.

electron rich compounds such as anilines¹¹ and phosphines.¹² To achieve iron-mediated oxidative activation of inert compounds (alkane as an example), strong stoichiometric oxidants are usually required to yield high valent iron(rv or v)oxo species that can dominate the activation and transformation of inert alkanes for the generation of value-added alcohols/ ketones, which is a representative mode for iron-catalyzed functionalization of inert compounds (Fig. 1b).¹³ By contrast, the oxidative ability of iron(m) is relatively moderate, hence direct iron(m)-dominated oxidative activation of inert compounds is still rare.

The input of light energy into iron(III) enhances the redox activity to a certain degree.14 Photo-induced iron ligand-to-metal charge transfer (LMCT) possesses excellent ability to produce radical species from abundant chemical feedstocks like alkyl carboxylates (^tBuCOO⁻ as a representative, $E_{1/2}^{ox} = +1.26$ V versus SCE),¹⁵⁻¹⁸ in which the necessary prerequisite is the suitable coordination of the substrate to the iron center. It means that activating those inert compounds with exceedingly weak coordination capacity in this fashion is helpless and doubtful. Very recently, through cooperating iron photocatalysis and redoxactive thiol catalysis, West and Xia elegantly reported direct decarboxylation of F-containing carboxylic acids for hydrofluoroalkylation of alkenes.19 The extremely high oxidative potential of TFA (CF₃COO⁻, $E_{1/2}^{ox} > +2.4$ V versus SCE) and the strong acid system make the corresponding activation mechanism of this special method complicated and confusing (Fig. 1c).

To develop a general strategy for the activation and transformation of inert and weakly coordinating compounds *via* metalbased photocatalysis, we hope to choose abundant and alkalescent halogen-containing carboxylates ($C_n X_m COO^-$, X: F, Cl, Br) as the

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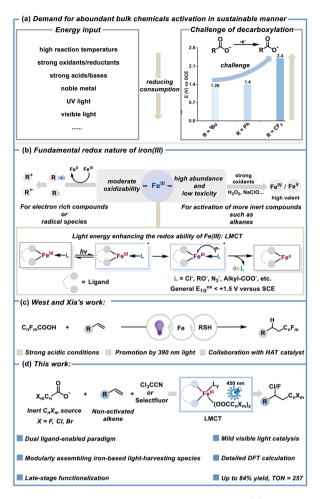


Fig. 1 Paradigms for activation of inert compounds. (a) Pursuing the activation of inert chemicals under milder conditions. (b) The redox character of iron(III). (c) Decarboxylation of C_nF_m COOH promoted by purple light. (d) This work: dual ligand-enabled iron and halogen-containing carboxylate-based photocatalysis for chloro/fluoro-polyhaloalkylation of alkenes.

investigational object to circumvent the strong acid conditions,²⁰ which is conducive to bringing the effect of the external ligand into full play.²¹ Given the fact that the role of the external ligand in 3d metal-based LMCT chemistry is usually intricate but significant, herein, we adopted the dual ligand-enabled strategy to efficiently assemble iron and $C_n X_m COO^-$ -based light-harvesting species for $C_n X_m$ radical production under visible light conditions,²² thus successfully achieving chloro/fluoro-polyhaloalkylation of non-activated alkenes with easily available trichloroacetonitrile/Select-fluor as the electrophilic halogenation reagent (Fig. 1d). OMe/CF₃-substituted bipyridine (bpy) and acetonitrile/trichloroacetonitrile indeed serve as the crucial dual ligands to achieve the desired LMCT between $C_n X_m COO^-$ and the iron(m) center and furnish the decarboxylation of $C_n X_m COO^-$ under visible light rather than UV light.

Results and discussion

To quickly obtain photochemical insights into iron(m) and C_n-X_mCOO⁻-based light-harvesting species, stoichiometric

photolysis experiments of Fe(CF₃COO)₃ were first carried out. As we expected, the high energy light ($\lambda_{max} = 365 \text{ nm}$) irradiation could induce the decomposition of Fe(CF₃COO)₃ to the CF₃ radical, which was captured by a non-activated alkene 1, and the generated radical adduct was further quenched through the halogen atom transfer (XAT) with Cl₃CCN, delivering the chlorotrifluoromethylation product of alkene (Fig. 2a, entry 1).²³ This process failed under visible light ($\lambda_{max} = 450 \text{ nm}$) or darkness conditions (entries 2 and 3). To reduce the energy consumption in the activation of CF₃COO⁻, the external ligands were investigated to achieve the release of the CF₃ radical under visible light conditions. Bpy L1 was incapable of controlling the effective decarboxylation of CF₃COO⁻ to access the chloro-trifluoromethylation of alkene, whereas its derivatives functionalized with electron-donating/withdrawing functional groups (L2 and L3) both showed the potential of being a competent ligand (entries 4-6).

Based on the preliminary results of the above stoichiometric experiments, a hypothetical catalytic cycle is illustrated in Fig. 2b. We proposed that a suitable electrophilic halogenation reagent (VIII or IX) should not only participate in the halogenatom transfer (XAT) with the radical adduct (VII) that was generated from the $C_n X_m$ radical (V) and alkene (VI) for the designed chloro/fluoro-polyhaloalkylation (XIV or XV), but also its concomitant electrophilic carbon/nitrogen radical species (X or XI) generated from the XAT would be responsible for the oxidation of [Fe^{II}/L] species (III) to regenerate the [Fe^{III}/L] complex (I). Notably, the additional effect of the ligand also possibly affected the process of [Fe^{II}/L] to [Fe^{III}/L], indicating that the choosing of the competent ligand is first and foremost for investigating conditions. After our detailed investigations (Fig. S14 and S15[†]), it was found that the chloro-trifluoromethylation and fluoro-chlorodifluoromethylation of alkenes via visible light-induced iron catalysis could be accomplished in good yields and regioselectivity under the regulation of L2 and L4, respectively (Fig. 2c). Regarding the choosing of the halogenation reagent, it was necessary to mention the result that commonly used chlorine sources such as trichloromethane, ethyl chloroacetate analogue, NCS (Nchlorosuccinimide), etc. could not yield the desired chloro-trifluoromethylation product (Fig. S17[†]), and bromo-trifluoromethylation of alkene by this method was infeasible (Fig. S18[†]).

With the optimized conditions in hand, the scope of this protocol was explored in terms of various non-activated alkenes' chloro/fluoro-polyhaloalkylation (Fig. 3). We first examined the reactivity of benzoate-containing terminal olefins. The different alkyl carbon chain structures of olefins did not disturb the reactivity and both provided good yields (4 and 5). Electron-donating or electron-withdrawing substituents in benzoate groups were compatible (6–8), and the substituents installed at *ortho* and *meta* positions also provided satisfactory yields (9–11). The alkene assembled with valuable benzotriazole successfully introduced the Cl and CF₃ groups in 74% yield under our standard conditions (12). Moreover, sulfonate/phenoxy-modified alkenes, simple hexadecane and α -methyl olefin were all well tolerated (13–17). Besides the above-mentioned terminal

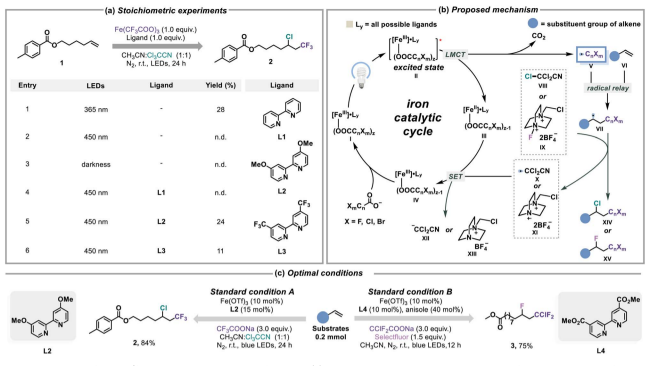


Fig. 2 Investigations of chloro/fluoro-polyhaloalkylation of alkenes. (a) Photo-induced iron mediated chloro-trifluoromethylation of an alkene. (b) Proposed mechanism of photo-induced iron catalyzed chloro/fluoro-polyhaloalkylation of alkenes. (c) Optimal conditions of the chloro/fluoro-polyhaloalkylation of alkenes.

olefins, the successful chloro-trifluoromethylation of the internal olefin enriched the diversity of the product (**18**). Significantly, the synthetic utility of this method in achieving fluoro-polyhaloalkylation²⁴ of non-activated alkenes was elaborated for the divergent synthesis of complex fluorine-containing molecules (**19–25**). Unfortunately, conjugated alkenes were not suitable for this reaction (Fig. S12†).

Furthermore, a wide variety of alkenes derived from pharmaceutical molecules were examined for the late-stage functionalization to extend this practical synthetic platform. The medicine oxaprozin used for rheumatoid arthritis and postoperative analgesia could be modified with halogen-containing alkyl in moderate yield (26). It is noteworthy that the performance of the derivative of piperonylic acid was amazing with 83% yield, whose sensitive ternary ring structure was well preserved (27). Likewise, ciprofibrate as an oral medicine for treating endogenous hypercholesterolemia and hypertriglyceridemia in adults, its derivative could also be achieved chloro-trifluoromethylation smoothly (28). The modification based on amino acid N-Boc-D-proline (29) increased the opportunities to discover late-model biomolecules. Of special note, eugenol possesses antioxidant, anticancer, anti-inflammatory, and antimicrobial activities, and its analogue gave moderate yield (30), despite the existence of the sensitive phenolic hydroxyl group. Besides, the late-stage chloro-trifluoromethylation, chloro-chlorodifluoromethylation, chlorobromodifluoromethylation and chloro-pentafluoroethylation of loxoprofen/febuxostat/flurbiprofen derivatives demonstrated intriguing synthesis potential for new drug research and development (**31–40**). Importantly, the utility and sustainability of this method were further highlighted by the gram-scale synthesis and low loading amount of the iron catalyst (TON = 257).

Based on the encouraging results of this ligand-enabled iron photocatalysis under visible light conditions, we turned our attention to the mechanistic studies. When stoichiometric radical scavenger (TEMPO/BHT) was subjected to the standard conditions (Fig. S30†), the desired chloro-trifluoromethylation process was severely inhibited, in which the radical adduct of the CF₃ radical and BHT was detected by electrospray ionization high-resolution mass spectrometry (ESI-HRMS, Fig. S31[†]). The existence of CF₃ radical species was also evidenced by the designed radical probe experiment (Fig. S32[†]). Theoretically, the production of the $C_n X_m$ radical required the excitation of iron and $C_n X_m COO^-$ -based light-harvesting species, meaning that continuous blue light irradiation should be necessary. The light on/off experiments verified this viewpoint (Fig. 4a). According to the fact that the combination of Fe(OTf)₃ and L2 showed first-order dependence during the kinetic studies (Fig. 4b), we realized that how to assemble iron salt/L2, C_n - $X_m COO^-$ and other potential ligands in this system into ironbased light-harvesting species for $C_n X_m$ radical release is the most crucial factor for the whole catalytic cycle. Because of the weaker coordination ability of OTf⁻ and NO₃⁻ than that of $C_n X_m COO^-$, easily dissociated iron salts like $Fe(OTf)_3$ and Fe(NO₃)₃·9H₂O whose coordination anions prefer to be

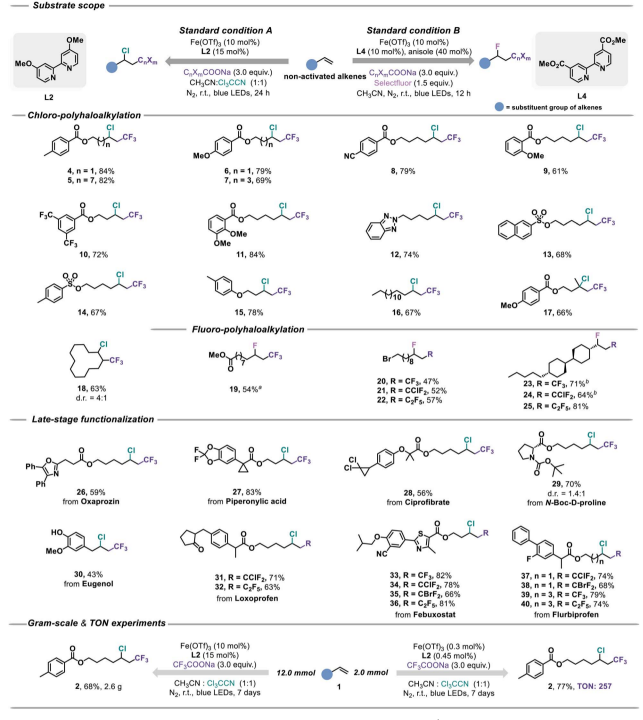
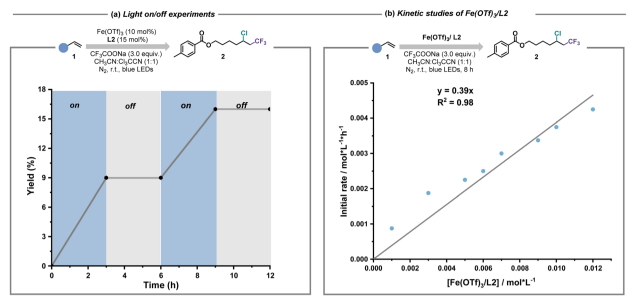


Fig. 3 Chloro/fluoro-polyhaloalkylation of alkenes via iron photocatalysis.^a 24 h reaction time.^b 0.5 mmol Selectfluor was used. Isolated yields.

exchanged by excess $C_n X_m COO^-$, showed good catalytic performance. In contrast, the efficiency of $Fe(acac)_3$ and $Fe(OAc)_3 \cdot nH_2O$ was unsatisfactory, probably resulting from the stagnant ligand exchange between $C_n X_m COO^-$ and $acac^-/OAc^-$ (Fig. 4c). Moreover, the fact that only when $Fe(OTf)_3$, **L2** and $C_n X_m COO^-$ are all present under the irradiation of blue light, can the iron(n) intermediate be observed, was revealed by UV-Vis experiments (Fig. 4d and S29†). To quickly determine the possible structure of iron(m) and $C_n X_m COO^-$ -based light-harvesting species, DFT calculations were carried out (Tables S1 and S2†). As shown in Fig. 5a, the suitable coordination of L2 and CH₃CN to Fe(CF₃COO)₃ (IntA) to form IntB is thermodynamically favorable ($\Delta G = -20.8$ kcal mol⁻¹), which also shows an obvious absorption band at 454 nm and an effective LMCT between CF₃COO⁻ and the iron(m) center in its excited state (Fig. S21–S24†). This result revealed



(c) Investigations of [Fe] -



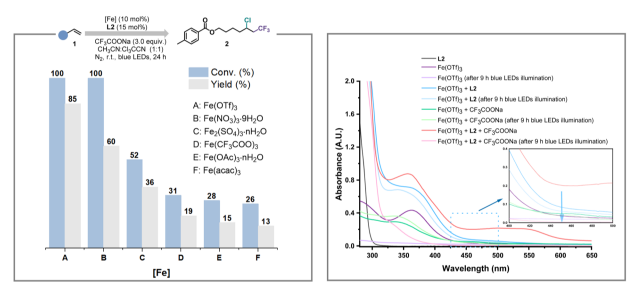


Fig. 4 Mechanistic studies. (a) Necessity of continuous blue light irradiation. (b) The first-order rate dependency on $Fe(OTf)_3/L2$ of the reaction. (c) The comparision of various iron(m) salts. (d) UV-Vis studies on the iron photocatalysis.

that CH_3CN served as the second ligand to construct **IntB**. The ESI-HRMS experiment also identified the real existence of **IntB** under the standard conditions (Fig. S34[†]). Given that some alkenes were equipped with the ester scaffold whose coordination capability in comparison to $C_nX_mCOO^-$ and nitriles should not be ignored, **IntC** as a possible LMCT-light-harvesting species could not be excluded. In contrast, the electron withdrawing inductive effect of the Cl group weakens the structural stability of **IntD**, potentially causing the low efficiency of producing C_nX_m radicals. Furthermore, we compared the performance of alkene **1** and **43** in different solvents such as CH_3CN , dichloromethane (DCM), acetone and CCl_3CN to prove the necessity of the second ligand for the activation of inert $C_nX_mCOO^-$ (Fig. 5b). Due to the fact that there is no

coordination effect between the iron(m) catalyst and DCM/43, the chloro-trifluoromethylation of 43 under the condition of DCM as solvent was inefficient (entry 1). By contrast, replacing DCM with acetone obviously enhanced the yield of **16** (entry 2). The similar reactivities of **43** in CH₃CN and CCl₃CN solvent explained the possibility that **IntD** truly served as alternative iron and $C_n X_m COO^-$ -based light-harvesting species in the absence of acetonitrile (entries 3 and 4). We also studied the relationship between the final yields of **16** and the amounts of adscititious CH₃CN, the desired reactivity for chloro-trifluoromethylation gradually improved, confirming that CH₃CN is the more efficient second ligand than CCl₃CN (Fig. S33†).

۵Me IntB IntD IntA IntC -20.8 ∧G (kcal/mol) = 0.0 -14.8 -7.8 Calculated structur Detected by ESI-HRMS I₁₈F₆FeN₄O₆⁺: 580.0480 found: 580.0475 IntA IntB IntC IntD (c) Stoichiometric CH₃CN as adscititious ligand (b) Identification of CH₃CN/CCI₃CN as the second ligand Fe(OTf)₃ (10 mol%) L2 (15 mol%) Fe(OTf)₃ (10 mol%) CH₃CN (x equiv.) CF₃COONa (3.0 equiv.) L2 (15 mol%) CF₃COONa (3.0 equiv.) DCM:CI₃CCN (1:1) Solvent:Cl₃CCN (1:1) N₂, r.t., blue LEDs, 24 h N₂, r.t., blue LEDs, 24 h 1 or 60 Entry Solvent Yield (%) of 2 Yield (%) of 16 1 DCM 83 23 50 CH₃COCH₃ Yield (%) 2 80 59 37 30 CH₃CN 70 84 3 20 Cl₃CCN 72 65 Substrates ż 10 15 43 x (equiv.)

(a) Iron and C_nX_mCOO⁻-based light-harvesting species

Fig. 5 Identification of iron and $C_n X_m COO^-$ -based light-harvesting species. (a) DFT calculations on iron and $C_n X_m COO^-$ -based light-harvesting species. (b) Experimental evidence for CH₃CN/CCl₃CN as the second ligand. (c) CH₃CN as the second ligand to promote the reaction.

Conclusion

In conclusion, we have developed a general and practical method in which dual ligand-enabled iron photocatalysis was leveraged for the activation of halogen-containing carboxylates under blue light conditions, thus achieving the chloro/fluoro-polyhaloalkylation of non-activated alkenes with broad functional group compatibility. Mechanistic studies revealed that the most likely real iron and $C_n X_m COO^-$ -based light-harvesting species is assembled by the external dual ligand—OMe/CF₃-substituted bipyridine and acetonitrile/trichloroacetonitrile. Extending this strategy to the activation of more inert bulk chemicals is underway in our laboratory.

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

Conceptualization: W. H., Z. Z., K. J., Y. L., D. W., S. L., and L. N.; methodology: W. H., Z. Z., K. J., Y. L., X. Y., D. W., S. L., and L. N.; investigation: W. H., Z. Z., K. J., Y. L., X. Y., D. W., S. L., and L. N.; supervision: Y. L., D. W., S. L., and L. N.; writing—original draft: W. H., Z. Z., K. J., Y. L., D. W., S. L., and L. N.; writing—review & editing: W. H., Z. Z., K. J., Y. L., X. Y., X. J., W. Y., Y. L., D. W., S. L., and L. N.

Conflicts of interest

The authors declare that they have no conflict of interest.

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