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# Photoinduced copper-catalyzed alkoxycarbonylation of alkyl fluorides†

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As the strongest  $\sigma$  bond in organic chemistry, the functionalization of the carbon–fluoride bond (BDE = 485 kJ mol<sup>-1</sup>) remains a significant challenge. Herein, we developed a carbonylation procedure that utilizes a copper catalyst and visible light irradiation to enable the production of esters from alkyl fluorides and phenols for the first time. From our mechanistic studies, the use of magnesium iodide as an additive was found to be crucial for the success of this protocol which drives the halide exchange and removes fluoride from the reaction.

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Ester compounds are of great importance as they are found in numerous natural products, pharmaceuticals, polymers, synthetic intermediates, etc.<sup>1</sup> Traditionally, esters are mainly synthesized from the esterification between the corresponding carboxylic acid derivatives and alcohols. However, the requirement of strongly alkaline conditions or/and a stoichiometric amount of activators restricts the application of this type of reaction.

On the other hand, using carbon monoxide as a cheap and readily available C1 source, the carbonylation reaction has become a powerful and dominant tool for producing various carbonylated compounds in a highly atom-economical manner.<sup>2</sup> In this area, the carbonylation reaction of organic halides has received considerable attention since 1963.<sup>3</sup> Generally, catalytic methods for carbonylation of organic halides proceed in two possible pathways: $4$  (i) transition metalcatalyzed carbonylation of  $sp^2$ -X bonds and benzylic halides via an ionic pathway (Scheme 1a (1)); Morikawa, Fuchikami, Beller and many other research groups developed various novel protocols for the carbonylation of aryl, allyl, benzyl, and perfluoroalkyl halides with alcohols.<sup>5</sup> The procedure applicable for alkyl halides is still very limited due to the barrier in oxidative addition and fast β-hydride elimination of the alkyl metal species.<sup>6</sup> Recently, (ii) radical carbonylation became a new and alternative way to resolve these problems (Scheme 1a  $(2)$ .<sup>7</sup> In 1988, Watanabe and co-workers developed a transition metal carbonyl complex-catalyzed radical carbonylation of alkyl iodide under visible light irradiation.<sup>8</sup> Subsequently,

Ryu's group reported a visible light-induced Pd-catalyzed radical carbonylation reaction of alkyl iodide with alcohols.<sup>9</sup> In 2016, Alexanian and co-workers developed a Pd-catalyzed radical alkoxycarbonylation of alkyl bromides using the NHC ligand.<sup>10a</sup> More recently, Beller's group disclosed the Rh-catalyzed radical carbonylation of primary alkyl chlorides with sodium iodide as the additive.<sup>10b</sup> Arndtsen and co-workers achieved novel catalytic systems for the carbonylative transformations of organic halides by the combination of a palladium catalyst and visible light.<sup>11</sup> In recent years, our group has also been interested in the metal-catalyzed carbonylation of alkyl halides under mild conditions, and various transition metal catalysts (Ru, Rh, Cu, and Fe) were studied.<sup>12</sup> However, for the carbonylation of alkyl fluorides, which have the highest RESEARCH ARTICLE<br> **(A)** Check for updates<br> **(A)** Check for updates<br>
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<sup>■</sup> unactivated alkyl fluorides ■ dual roles of copper in photochemistry and coupling

Scheme 1 Alkoxycarbonylation of alkyl halides.

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<sup>■</sup> broad scope. >50 examples ■ good functional group compatibility

bond dissociation energy (485 kJ mol<sup>-1</sup>; Scheme 1b),<sup>13,14</sup> no procedure yet has been reported.

Herein, we present the successful alkoxycarbonylation of alkyl fluorides with phenols under light irradiation catalyzed by a copper catalyst (Scheme 1c). The addition of magnesium iodide was found to be crucial for the success of this transformation, which not only generates more reactive species in situ via halogen exchange but also avoids the influence of fluoride ions in the reaction solution on the subsequent radical course. A wide range of valuable esters were produced in high yields with excellent functional group adaptations.

Our initial studies were carried out with 1-fluorooctane (1a) and phenol (2a) as the model substrates to examine the reaction parameters (Table 1 and also see the ESI, Tables S1–S5†). After a series of studies, we were pleased to find that the desired reaction indeed worked smoothly at 80 °C, when employing a catalytic system consisting of  $Cu(CH_3CN)_4PF_6$ (10 mol%) and Xantphos (11 mol%) in the presence of 2.5 equivalents of  $K_2CO_3$  and 1.5 equivalents of MgI<sub>2</sub>, and the desired product 3a was formed in 95% NMR yield (Table 1, entry 1). Control experiments, without light irradiation or the use of blue light, confirmed that the reaction proceeded through a photocatalytic process (Table 1, entries 2 and 3). No reaction occurred at room temperature and the alkyl fluoride remained unreacted (Table 1, entry 4). The use of other metal iodides such as LiI and CaI<sub>2</sub> instead of MgI<sub>2</sub> resulted in no reaction despite the detection of 1-iodooctane (Table 1, entries 5 and 6). Notably, the valence of the copper salt is important

Table 1 Optimization of the reaction conditions<sup>6</sup>  $1a$  $2a$ anisole (1 mL), 320-400 nm  $3a$ 80 °C, 24 h Entry Deviation from standard conditions 3a yield (%) 1 None 95  $(92)^b$ 2 No light 0<br>3 Blue light 0 Blue light 0<br>t.t. 0 4 r.t. 0 5 LiI instead of MgI<sub>2</sub> 0<br>6 CaI<sub>2</sub> instead of MgI<sub>2</sub> 0 6 CaI<sub>2</sub> instead of MgI<sub>2</sub> 0<br>7 No Cu(CH<sub>3</sub>CN)<sub>4</sub>PF<sub>6</sub> 0 7 No Cu(CH<sub>3</sub>CN)<sub>4</sub>PF<sub>6</sub> 0<br>8 CuI instead of Cu(CH<sub>2</sub>CN)<sub>4</sub>PF<sub>6</sub> 15 8 CuI instead of Cu(CH<sub>3</sub>CN)<sub>4</sub>PF<sub>6</sub> 15<br>9 Cu(OAc)<sub>2</sub> instead of Cu(CH<sub>3</sub>CN)<sub>4</sub>PF<sub>6</sub> 0 9 Cu(OAc)<sub>2</sub> instead of Cu(CH<sub>3</sub>CN)<sub>4</sub>PF<sub>6</sub> 0<br>10 BINAP instead of Xantphos <5 BINAP instead of Xantphos 11 11 Niphos instead of Xantphos 16<br>12  $Na_2CO_3$  instead of  $K_2CO_3$   $\leq 5$ 12  $Na_2CO_3$  instead of  $K_2CO_3$   $\leq$  5<br>13  $Cs_3CO_2$  instead of  $K_3CO_2$  33 13  $CS_2CO_3$  instead of  $K_2CO_3$ <br>14 DBU instead of  $K_2CO_3$  0 14 DBU instead of  $K_2CO_3$  0<br>15 CH<sub>2</sub>CN as solvent 0 15 CH<sub>3</sub>CN as solvent 0<br>16 Toluene as solvent 58 Toluene as solvent

"Standard conditions: 1a (0.2 mmol, 2.0 eq.), 2a (0.1 mmol, 1.0 eq.), MgI<sub>2</sub> (0.15 mmol, 1.5 eq.), Cu(CH<sub>3</sub>CN)<sub>4</sub>PF<sub>6</sub> (10 mol%), Xantphos  $(11 \text{ mol\%})$ , K<sub>2</sub>CO<sub>3</sub> (0.25 mmol, 2.5 eq.), CO (40 bar), anisole (1 mL), stirred at 80 °C for 24 h under irradiation (320–400 nm); yields were determined by  ${}^{1}H$  NMR with 1,3,5-trimethoxybenzene as the internal standard. <sup>b</sup> Isolated yield.

for the onset of the reaction as well;  $Cu<sup>1+</sup>$  could initiate the alkoxycarbonylation, whereas no product could be found when  $Cu^{2+}$  was used or in the absence of a copper salt (Table 1, entries 7–9). We then investigated other bidentate phosphine ligands and an obvious effect on the yield of the final product was found, with Xantphos still being the best ligand of choice (Table 1, entries 10 and 11). The effects of bases were tested, and poor yields were obtained with either other inorganic bases or organic bases (Table 1, entries 12–14). Moreover, we also found that the reaction appeared to be sensitive to the properties of solvents; a strongly polar solvent completely inhibited the formation of the targeted ester, and a non-polar or weakly polar solvent ensured the proceeding of the desired alkoxycarbonylation, with anisole being the best solvent (Table 1, entries 15 and 16).

With the optimized reaction conditions in hand, we investigated the substrate scope of this reaction, and the results are shown in Scheme 2. At the first stage, a range of phenols with electron-donating groups or electron-withdrawing groups could serve as effective partners, providing the desired esters in good to excellent yields. Versatile functional groups, such as alkyl (3b–3d, 3o), phenyl (3e), halogen (3f–3i), trifluoromethyl  $(3j)$ , cyano  $(3k)$ , alkoxyl  $(3l)$ , alkylthio  $(3m)$ , alkylsulfonyl  $(3n)$ , and Bpin (3p), were all tolerated well in the reaction system with high yields. Subsequently, the steric hindrance effect was tested. Phenols with different groups substituted at the metaposition were used, and the desired meta-substituted phenol esters were obtained (3q–3s). Phenols bearing ortho-substituents gave the desired esters in moderate to good yields as well irrespective of the steric hindrance effect (3t–3w). Furthermore, phenols possessing di-substituents still reacted well and gave the products in good yields (3x–3z, 4a, and 4b). The natural product sesamol can gave the corresponding product with excellent yield in this reaction (4c). Next, we investigated the reaction with naphthols, and 2-naphthol gave a better yield than 1-naphthol (4d and 4e). Aliphatic alcohol was also tested but was proven to be an unsuitable reaction partner, which gave the corresponding product in only 22% yield (4f). Remarkably, an interesting product was formed when we used 4-hydroxybenzyl alcohol as the substrate. The 4-hydroxybenzyl alcohol reacted with anisole first through electrophilic aromatic substitution, $15$  and then alkoxycarbonylation occurred to give the final product (4g) in 80% yield. Organic Chemistry Frontiers<br>
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> Subsequently, the scope of different types of alkyl fluorides was examined (5a–5u). The alkyl fluoride with a long carbon chain also gave an excellent yield (5a), and functional groups such as ether (5b), thioether (5c), alkenyl (5d), and alkyl groups (5e–5g) were all well tolerated. Phenoxy fluorides with different substituents also gave excellent yields (5h–5k). Meanwhile, heterocycle (5l), phenthio (5m), phensulfonyl (5n), and naphthyloxy (5p) groups were all compatible under the standard conditions to give the desired esters in moderate to good yields. It is noteworthy that the reaction was also extended to the alkoxycarbonylation of secondary and tertiary alkyl fluorides, with the corresponding esters being delivered smoothly (5q and 5r). Importantly, the alkoxycarbonylation



Scheme 2 Reaction scope. Reaction conditions: alkyl fluorides 1 (0.2 mmol, 2eq.), phenols 2 (0.1 mmol, 1.0 eq.), Cu(CH<sub>3</sub>CN)<sub>4</sub>PF<sub>6</sub> (10 mol%), Xantphos (11 mol%), MgI<sub>2</sub> (1.5 eq.), K<sub>2</sub>CO<sub>3</sub> (2.5 eq.), anisole (1 mL), CO (40 bar), 80 °C, irradiation (320–400 nm), 24 h.

can incorporate an ester fragment into natural products, while the alkyl fluoride derivatives from sesamol, estrone, and DLα-tocopherol could also be converted into the corresponding alkoxycarbonylation products in 60–85% yields (5s–5u).

Control studies were also carried out to explore the reaction mechanism. At first, we studied the conversion of alkyl fluoride to alkyl iodide, and the fluoride/iodide exchange reaction

proceeded with an excellent yield at 80 °C (Scheme 3a-I). To gain more insight into the mechanism and to identify the ratedetermining step of the reaction, we studied the kinetic profile of the halogen exchange reaction. As shown in Fig. 1, in the first 2 h, 70% of 1-fluorooctane was converted. Then, the conversion rate gradually slowed down, and conversion was achieved in 85% yield after 24 h. The above results indicate





Fig. 1 Kinetic profile of the halogen exchange reaction between 1-fluorooctane and 1-iodooctane.

that the rate-determining step of our reaction is not halogen exchange. Subsequently, alkyl iodide was tested in this reaction. The reaction of 1b at room temperature or under heating all gave the desired product in good yields (Scheme 3a-II), which means heating is not necessary for the proceeding of alkoxycarbonylation. For a better understanding of the role of magnesium iodide in this reaction, reactions with alkyl iodide and metal fluoride were performed under our standard conditions. The results show that magnesium fluoride has no effect on the carbonylation (Scheme 3a-III). However, when using other fluorides as additives, the reaction gave no desired ester product but converted back to alkyl fluoride (Scheme 3aIV). More specifically, halogen exchange is a reversible reaction, and in the presence of fluoride ions, alkyl iodide will preferentially undergo halogen exchange to the more stable alkyl fluoride rather than carbonylation. Thanks to the ultra-low solubility of magnesium fluoride in anisole, our standard reaction is essentially free of fluoride ions, which is the key to the successful achievement of the reaction with good yields of the desired ester products.

To further shed light on the mechanism, additional mechanistic experiments were conducted. When stoichiometric radical scavengers, 2,2,6,6-tetramethyl-1-piperidinyloxy (TEMPO) and 1,1-diphenylethene (1,1-DPE), were added to the model reaction of 1a and 2a under standard alkoxycarbonylation conditions, the formation of ester 3a was completely suppressed. The adducts of olefin with alkyl radical 6a were detected by GC-MS (Scheme 3b). In the radical inhibition experiments, the yields gradually decreased with the gradual addition of 2,6-di-tert-butyl 4-methylphenol (BHT, a radical scavenger, 0–2 equiv.) (Scheme 3c). Subsequently, in radical clock experiments, reactions using phenol and (1-fluorohex-5 en-3-yl)benzene as the substrates, the corresponding cyclization product 7 was obtained as the major ester (Scheme 3d). All these results imply the existence of radical intermediates in the reaction.

Based on the above results and literature reports, $7-14$  we proposed a mechanism for this reaction (Scheme 4). Initially, the LCu<sup>I</sup>OAr species (I) was generated from the copper salt through ligand exchange with phenol and the ligand.<sup>16</sup> Then the complex I excited to its excited state adduct II under irradiation. Meanwhile, the alkyl fluoride undergoes halogen exchange to generate the corresponding alkyl iodide.



Afterwards, complex II engaged in a single electron transfer reaction with alkyl iodide to form an alkyl radical and complex III. The alkyl radical was then captured by carbon monoxide to afford an acyl radical which will react with complex III to give acyl copper complex IV. Finally, the target ester can be eliminated by reductive elimination and regenerate  $Cu<sup>I</sup>$  species for the next catalytic cycle.

#### **Conclusions**

In summary, we have developed a photoinduced copper-catalyzed alkoxycarbonylation of alkyl fluorides with phenols to synthesize various esters. The remarkable scope and functional group tolerance were evidenced by >50 examples and also the late-stage modification of natural product-related substrates. As the first example of the carbonylation of alkyl fluorides, the procedure provides a new pathway for the functionalization of alkyl fluorides.

### Author contributions

X.-F. W. conceived and designed the project; P. Y. and Y.-H. Z. conducted the experiments; P.Y. analysed and interpreted the experimental data; P. Y. and X.-F. W. prepared the manuscript; P. Y. and X.-F. W. prepared the ESI.†

# Conflicts of interest

There are no conflicts to declare.

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#### References

- 1 (a) R. C. Larock, Comprehensive Organic Transformations: A Guide to Functional Group Preparations, Wiley-VCH, 1999; (b) J. Otera, Esterification: Methods, Reactions, and Applications, Wiley, 2010; (c) C. Liu, S. Tang and A. Lei, Oxidant controlled Pd-catalysed selective oxidation of primary alcohols, Chem. Commun., 2013, 49, 1324–1326; (d) C. Locatelli, F. B. Filippin-Monteiro and T. B. Creczynski-Pasa, Alkyl esters of gallic acid as anticancer agents: A review, Eur. J. Med. Chem., 2013, 60, 233– 239; (e) B. Liu, F. Hu and B.-F. Shi, Recent Advances on Ester Synthesis via Transition-Metal Catalyzed C-H Functionalization, ACS Catal., 2015, 5, 1863–1881.
- 2 (a) B. Gabriele, Carbon Monoxide in Organic Synthesis: Carbonylation Chemistry, Wiley, 2021; (b) J.-B. Peng, F.-P. Wu and X.-F. Wu, First-Row Transition-Metal-Catalyzed Carbonylative Transformations of Carbon Electrophiles, Chem. Rev., 2019, 119, 2090–2127; (c) J.-B. Peng, H.-Q. Geng and X.-F. Wu, The Chemistry of CO: Carbonylation, Chem, 2019, 5, 526–552.
- 3 R. F. Heck and D. S. Breslow, Carboxyalkylation reactions catalyzed by cobalt carbonylate ion, *J. Am. Chem. Soc.*, 1963, 85, 2779–2782.
- 4 B. Lu, M. Xu, X. Qi, M. Jiang, W.-J. Xiao and J.-R. Chen, Switchable Radical Carbonylation by Philicity Regulation, J. Am. Chem. Soc., 2022, 144, 14923–14935.
- 5 (a) J. Tsuji, J. Kiji, S. Imamura and M. Morikawa, Organic Syntheses by Means of Noble Metal Compounds. VIII.1 Catalytic Carbonylation of Allylic Compounds with Palladium Chloride, J. Am. Chem. Soc., 1964, 86, 4350– 4353; (b) U. Hisao, O. Kosukegawa, Y. Ishii, H. Yugari and T. Fuchikami, Carbonylation of 1-perfluoroalkyl-substituted 2-iodoalkanes catalyzed by transition-metal complexes, Tetrahedron Lett., 1989, 30, 4403–4406; (c) W. Mägerlein, M. Beller and A. F. Indolese, Palladium-catalyzed carbonylation of aryl halides - a detailed investigation of the alkoxycarbonylation of 4-bromoacetophenone, J. Mol. Catal. A: Chem., 2000, 156, 213–221; (d) A. M. Trzeciak, W. Wojtków, Z. Ciunik and J. J. Ziółkowski, Low-Pressure Carbonylation of Benzyl Bromide with Palladium Complexes Modified with PNS or  $P(OPh)_{3}$ . Structural Identification of Palladium-Catalyst Intermediate, Catal. Lett., 2001, 77, 245–249.
- 6 (a) A. C. Bissember, A. Levina and G. C. Fu, A Mild, Palladium-Catalyzed Method for the Dehydrohalogenation of Alkyl Bromides: Synthetic and Mechanistic Studies, J. Am. Chem. Soc., 2012, 134, 14232–14237; (b) L. Wu, X. Fang, Q. Liu, R. Jackstell, M. Beller and X.-F. Wu, Palladium-catalyzed carbonylative transformation of C

(sp<sup>3</sup>)-X bonds, ACS Catal., 2014, 4, 2977-2989; (c) X.-W. Gu, Y. Zhang, F. Zhao, H.-J. Ai and X.-F. Wu, Phosphine-catalyzed photo-induced alkoxycarbonylation of alkyl iodides with phenols and 1,4-dioxane through charge-transfer complex, Chin. J. Catal., 2023, 48, 214–223.

- 7 (a) I. Ryu and N. Sonoda, Free-Radical Carbonylations: Then and Now, Angew. Chem., Int. Ed. Engl., 1996, 35, 1050– 1066; (b) I. Ryu, Radical carboxylations of iodoalkanes and saturated alcohols using carbon monoxide, Chem. Soc. Rev., 2001, 30, 16–25; (c) S. Zhao and N. P. Mankad, Metal-catalysed radical carbonylation reactions, Catal. Sci. Technol., 2019, 9, 3603–3613.
- 8 (a) T. Kondo, Y. Tsuji and Y. Watanabe, Photochemical carbonylation of alkyl iodides in the presence of various metal carbonyls, Tetrahedron Lett., 1988, 29, 3833–3836; (b) T. Kondo, Y. Sone, Y. Tsuji and Y. Watanabe, Photo-, electro-, and thermal carbonylation of alkyl iodides in the presence of group 7 and 8-10 metal carbonyl catalysts, J. Organomet. Chem., 1994, 473, 163–173.
- 9 (a) A. Fusano, S. Sumino, S. Nishitani, T. Inouye, K. Morimoto, T. Fukuyama and I. Ryu, Pd/Light-Accelerated Atom-Transfer Carbonylation of Alkyl Iodides: Applications in Multicomponent Coupling Processes Leading to Functionalized Carboxylic Acid Derivatives, Chem. – Eur. J., 2012, 18, 9415–9422; (b) S. Sumino, A. Fusano, T. Fukuyama and I. Ryu, Carbonylation Reactions of Alkyl Iodides through the Interplay of Carbon Radicals and Pd Catalysts, Acc. Chem. Res., 2014, 47, 1563–1574.
- 10 (a) B. T. Sargent and E. J. Alexanian, Palladium-Catalyzed Alkoxycarbonylation of Unactivated Secondary Alkyl Bromides at Low Pressure, J. Am. Chem. Soc., 2016, 138, 7520–7523; (b) P. Wang, Y. Wang, H. Neumann and M. Beller, Rh-catalyzed alkoxycarbonylation of unactivated alkyl chlorides, Chem. Sci., 2022, 13, 13459–13465.
- 11 (a) G. M. Torres, Y. Liu and B. A. Arndtsen, A dual lightdriven palladium catalyst: Breaking the barriers in carbonylation reactions, Science, 2020, 368, 318–323; (b) Y. Liu, C. Zhou, M. Jiang and B. A. Arndtsen, Versatile Palladium-Catalyzed Approach to Acyl Fluorides and Carbonylations by Combining Visible Light- and Ligand-Driven Operations, J. Am. Chem. Soc., 2022, 144, 9413–9420; (c) K. E. Chami, Y. Liu, M. A. Belahouane, Y. Ma, P.-L. Lagueux-Tremblay and B. A. Arndtsen, A Visible Light Driven Nickel Carbonylation Catalyst: The Synthesis of Acid Chlorides from Alkyl Halides, Angew. Chem., Int. Ed., 2023, 62, e202213297.
- 12 (a) H.-J. Ai, Y. Yuan and X.-F. Wu, Ruthenium pincer complex-catalyzed heterocycle compatible alkoxycarbonylation of alkyl iodides: substrate keeps the catalyst active, Chem. Sci., 2022, 13, 2481–2486; (b) X. Qi, R. Zhou, H.-J. Ai and X.-F. Wu, HMF and furfural: Promising platform molecules in rhodium-catalyzed carbonylation reactions for the synthesis of furfuryl esters and tertiary amides, J. Catal., 2020, 381, 215–221; (c) Y. Li and X.-F. Wu, Copper/iron cocatalyzed alkoxycarbonylation of unactivated alkyl bromides, Commun. Chem., 2018, 1, 39; (d) F. Zhao, P. Russo,

R. Mancuso, B. Gabriele and X.-F. Wu, Copper-catalyzed carbonylative coupling of alkyl iodides with phenols for the synthesis of esters, J. Catal., 2022, 413, 907–912.

- 13 (a) H. Yamamoto, T. Hiyama, K. Kanie, T. Kusumoto, Y. Morizawa and M. Shimzu, Organofluorine Compounds: Chemistry and Applications, Springer Berlin Heidelberg, 2000; (b) H. Amii and K. Uneyama, C-F Bond Activation in Organic Synthesis, Chem. Rev., 2009, 109, 2119–2183; (c) T. Ahrens, J. Kohlmann, M. Ahrens and T. Braun, Functionalization of Fluorinated Molecules by Transition-Metal-Mediated C-F Bond Activation To Access Fluorinated Building Blocks, Chem. Rev., 2015, 115, 931–972; (d) T. Iwasaki and N. Kambe, Cross- and Multi-Coupling Reactions Using Monofluoroalkanes, Chem. Rec., 2023, 23, e202300033.
- 14 (a) J. Choi, D. Y. Wang, S. Kundu, Y. Choliy, T. J. Emge, K. Krogh-Jespersen and A. S. Goldman, Net Oxidative Addition of  $C(sp^3)$ -F Bonds to Iridium via Initial C-H Bond Activation, Science, 2011, 332, 1545–1548; (b) C. B. Caputo, L. J. Hounjet, R. Dobrovetsky and D. W. Stephan, Lewis Acidity of Organofluorophosphonium Salts: Hydrodefluorination by a Saturated Acceptor, Science, 2013, 341, 1374–1377; (c) W. Gu, M. R. Haneline, C. Douvris and O. V. Ozerov, Carbon–Carbon Coupling of C(sp<sup>3</sup>)–F Bonds Using Alumenium Catalysis, J. Am. Chem. Soc., 2009, 131, 11203–11212; (d) M. Ahrens, G. Scholz, T. Braun and E. Kemnitz, Catalytic Hydrodefluorination of Fluoromethanes at Room Temperature by Silylium-ion-like Surface Species, Angew. Chem., Int. Ed., 2013, 52, 5328–5332; (e) C. Douvris and O. V. Ozerov, Hydrodefluorination of Perfluoroalkyl Groups Using Silylium-Carborane Catalysts, Science, 2008, 321, 1188– 1190; (f) C. Douvris, C. M. Nagaraja, C.-H. Chen, B. M. Foxman and O. V. Ozerov, Hydrodefluorination and Other Hydrodehalogenation of Aliphatic Carbon-Halogen Bonds Using Silylium Catalysis, J. Am. Chem. Soc., 2010, 132, 4946–4953. Organic Chemistry Frontiers Week on 06 X-2022. 2021. This article is calibrical controllar on distribution and the state of the st
	- 15 (a) S. F. R. Taylor, J. Sá and C. Hardacre, Friedel-Crafts Alkylation of Aromatics with Benzyl Alcohol over Gold-Modified Silica, ChemCatChem, 2011, 3, 119–121; (b) M. Rueping and B. J. Nachtsheim, A review of new developments in the Friedel-Crafts alkylation – From green chemistry to asymmetric catalysis, Beilstein J. Org. Chem., 2010, 6, 6, DOI: [10.3762/bjoc.6.6](https://doi.org/10.3762/bjoc.6.6).
	- 16 (a) Y. Jia, Z. Zhang, G.-M. Yu, X. Jiang, L.-Q. Lu and W.-J. Xiao, Visible Light Induced Copper-Catalyzed Enantioselective Deaminative Arylation of Amino Acid Derivatives Assisted by Phenol, Angew. Chem., Int. Ed., 2023, 62, e202312102; (b) J.-P. Yue, J.-C. Xu, H.-T. Luo, X.-W. Chen, H.-X. Song, Y. Deng, L. Yuan, J.-H. Ye and D.-G. Yu, Metallaphotoredox-enabled aminocarboxylation of alkenes with  $CO<sub>2</sub>$ , Nat. Catal., 2023, 6, 959-968; (c) C. Chen, J. C. Peters and G. C. Fu, Photoinduced copper-catalysed asymmetric amidation via ligand cooperativity, Nature, 2021, 596, 250–256; (d) C. Chen and G. C. Fu, Copper-catalysed enantioconvergent alkylation of oxygen nucleophiles, Nature, 2023, 618, 301–307.