The development of catalytic carboacylation of simple olefins, which would enable the rapid construction of ketones with high levels of complexity and diversity, is very challenging. To date, the vast majority of alkene carboacylation reactions are typically restricted to single- and two-component methodologies. Here we describe a three-component carboacylation of alkenes via the merger of radical chemistry with nickel catalysis. This reaction manifold utilizes a radical relay strategy involving radical addition to an alkene followed by alkyl radical capture by an acyl-nickel complex to forge two vicinal C−C bonds under mild conditions. Excellent chemoselectivity and regioselectivity have been achieved by utilizing a pendant weakly chelating group. This versatile protocol allows for facile access to a wide range of important β-fluoroalkyl ketones from simple starting materials.
Ketones are important structural motifs prevalent in pharmaceuticals, agrochemicals and natural products, as well as versatile intermediates for numerous valuable transformations. Transition metal-catalyzed cross coupling with acyl electrophiles has emerged as an efficient platform for ketones synthesis, providing an alternative to classical nucleophilic carbonyl addition and Friedel–Crafts acylation. Particularly, catalytic carboacylation of alkenes in the presence of transition metal catalysts has attracted considerable attention in synthetic chemistry, due to the fact that: (i) alkenes are abundant and ubiquitous building blocks in chemical and material industries; (ii) this protocol installs multiple C–C bonds across an olefin in one step, allowing for rapid access to complex ketones from simple starting materials.

To date, the vast majority of alkene carboacylation reactions proceed via intramolecular insertion of an acyl-metal intermediate into an olefin. As a result, this reaction is typically restricted to single-component variants, with very few exceptions of two-component reactions. The development of fully intermolecular, three-component carboacylation of simple olefins, which would enable the rapid construction of ketone products with high levels of complexity and diversity, is highly desired and remains elusive. One main challenge to this approach is the propensity for decarbonylation from an acyl-metal species, which must intercept the alkene. Another challenge associated with a three-component variant is achieving regioselectivity and chemoselectivity. A number of elegant methods have been reported to control regioselectivity in transition metal-catalyzed three-component dicarbofunctionalization of alkenes. For instance, previous studies have demonstrated that employment of activated substrates can facilitate regiocontrol. More recently, utilization of directing groups to achieve highly regioselective dicarbofunctionalization of alkenes has also been reported. Nevertheless, the capability to achieve selective alkene functionalization in the presence of multiple double bonds remains a longstanding challenge.

Over the last decade, nickel-catalyzed cross-coupling has emerged as a powerful tool to forge C–C bonds in chemical synthesis. Particularly, the merger of radical chemistry with nickel catalysis has enabled the invention of numerous useful transformations. We recently questioned whether radical-based nickel catalysis might offer an alternative pathway to facilitate the three-component carboacylation of alkenes. Specifically, we envisioned that a radical relay strategy, involving radical addition to an alkene followed by alkyl radical capture by an acyl-nickel species, would bypass the challenging acyl-metal/alkene capture required in a conventional two-electron mode. Furthermore, we anticipated that a pendant chelating group could facilitate the capture of the alkyl radical by the nickel species, while also controlling chemoselectivity and regioselectivity via coordination. This strategy described was also validated in seminal work by Nevada and coworkers in which an allicyclic acetate could serve to stabilize a putative alkynickel species after radical recombination. Given the increasing importance of fluoroalkyl moieties in the areas of medicinal, agrochemical, and material chemistry, as well as elegant progress in radical fluoroalkylation of alkenes, we sought to explore the carboacylation of alkenes with fluoroalkyl precursors. Herein, we report selective, three-component carboacylation of alkenes with acyl chlorides via Ni-catalyzed radical relay (Fig. 1). This method takes advantage of Ni-catalyzed reductive coupling of electrophiles under mild conditions, delivering valuable β-fluoroalkyl carbonyls that are not easily accessible by other methods.

**Results**

**Optimization study.** We first sought to employ carbonyls, ubiquitous intermediates in organic synthesis, as chelating groups. Evaluation of this nickel-catalyzed intermolecular reductive carboacylation strategy was first examined with allyl hexanoate, tert-butylbenzoyl chloride, and perfluorobutyl iodide in the presence of nickel catalysts and reductants. We were delighted to find that 93% yield of the desired carboacylation product could be obtained with Mn dust as stoichiometric reductant in the presence of catalytic amounts of NiCl₂·glyme and 4,4-di-tert-butyl-2,2′-dipryridyl (dbbpY) at 25°C (See Supplementary Table 1). Control experiments have demonstrated that both the nickel catalyst and Mn dust are essential for the desired transformation to proceed, while moderate yield of 3 can be obtained in the absence of ligand, indicating the chelating ability of the pendant ester group (See Supplementary Table 1).

**Substrate scope.** With optimal reaction conditions in hand, we explored the generality of this carboacylation with respect to the alkene fragment and found that a variety of readily available alkenes are viable partners for this transformation (Fig. 2a). Alkenes tethered with different chelating groups, including esters, carbonates, carbamates, sulfonates, and phosphates, undergo the desired reductive coupling with moderate to excellent efficiency (products 1–9, 42–93% yields). Interestingly, electron-rich aryl rings were capable of guiding the desired transformations, exemplified by the reaction of aryl vinyl ethers to furnish α-oxo-β-fluoroalkyl ketones in high yields (products 10 and 11, 63% and 76% yields, respectively). The mild reaction conditions allow for good compatibility with a wide range of important functional groups including aryl bromides and chlorides, providing a versatile platform for further synthetic manipulations (products 13, 74% yield). Gratifyingly, more complex partners derived from naturally occurring molecules were also successfully employed, demonstrating the potential applicability of this methodology in late-stage functionalization. For example, derivatives of borneol...
and estrone functioned as competent alkene partners, furnishing each of the desired coupling products with high efficiency (products 14 and 15, 77% and 67% yields, respectively). Notably, acyclic internal alkenes, such as (E)-but-2-en-1-yl benzoate, also participated in this Ni-catalyzed difunctionalization manifold with moderate efficiency (see Supplementary Fig. 5).

Next, we evaluated the scope of the acyl chloride component in this protocol. As revealed in Fig. 2b, aromatic acyl chlorides containing electron-neutral, -donating, and -withdrawing groups proceeded smoothly under the optimal conditions, delivering the product ketones with moderate to excellent efficiency (products 16–26, 49–83% yields). Moreover, ortho substitution on the aryl ring was tolerated, albeit with somewhat diminished efficiency (product 25, 54% yield). Notably, heteroaromatic acyl chlorides such as thiophene and furan were also viable substrates, furnishing the corresponding heteroaryl ketones in moderate yields (products 27 and 28, 72% and 58% yields, respectively).

Additionally, aliphatic acyl chlorides demonstrated promising levels of efficiency, as exemplified by dialkyl ketone product 29 (42% yield).

Finally, we examined this three-component functionalization protocol with varied fluoroalkyl precursors (Fig. 2c). A series of perfluoroalkyl iodides, including ethyl iododifluoroacetate, can be readily employed with excellent levels of efficiency (products 30–35, 53–91% yields), providing a simple and efficient way to incorporate perfluoroalkyl substituents into complex molecules. Importantly, this efficient alkene carboacylation strategy employs a 1:1 ratio of alkene and fluoroalkyl iodide at room temperature, without the need for excess amounts of perfluoroalkyl iodides in all cases (Fig. 2). Gratifyingly, trifluoromethiodide (CF3I) was also reactive, affording the desired trifluoromethylacetylation product in promising levels of efficiency (29% yield) (see Supplementary Fig. 6). Additionally, electron-deficient tertiary alkyl bromides were also suitable coupling partners under slightly
In 4:1 CH$_3$CN/DME [0.05 M], 41% of alkene recovered. 48 h.

Fig. 3 Chemoselectivity guided by pendant chelating groups. a Reactivity tuned by the chelating group. b Chemoselectivity in the presence of multiple double bonds.

modified conditions. For example, the reaction of ethyl 2-bromo-2-methylpropanoate gave the desired carboxylated product in 36% yield (see Supplementary Fig. 7).

The effect of the weakly coordinating group in chelation was evaluated by testing a set of different substrates shown in Fig. 3a. In cases of substrates which form six- or seven-membered chelate rings, the reactions proceeded with excellent efficiency (2 and 36, 89% and 88% yields, respectively). A slightly decreased efficiency was observed in the case of the substrate proceeding via an eight-membered chelate ring (37, 57% yield). It is worth noting, however, that the reaction of vinyl benzoate 38, bearing a potential five-membered chelate ring, is also hampered by starting material decomposition under the standard conditions. Moreover, alkynes bearing strained or non-chelating groups were ineffective, further highlighting the importance of the directing groups (39 and 40).

Intrigued by these results, we examined the chemoselectivity and regioselectivity with substrates bearing multiple double bonds. As shown in Fig. 3b, our reductive protocol can be used for selective functionalization of alkenes through the directing effect. Exclusive site selectivity was observed for carboacylation of terminal alkenes that could form six-membered chelate rings, leaving the non-chelating alkenes untouched (products 41–46, 43–68% yields). Notably, for substrates in which two terminal alkenes are present, excellent site selectivity was observed at olefins guided by the tethered chelating group with a more favorable chelate geometry (products 41, 43% yield), further highlighting the powerful selectivity feature of this catalytic system. Additionally, olefins derived from naturally occurring molecules, including valencene, coumarin, dehydroisoandrosterone, and cholesterone, could also be employed to furnish the desired products with moderate efficiency and excellent selectivity (products 43–46, 47–68% yields), demonstrating the inherent value of our carboacylation protocol in late-stage functionalization.

To further demonstrate the synthetic benefit of our nickel-catalyzed reductive carboacylation strategy, the difunctionalized products were converted to several useful synthetic functionalities (Fig. 4). The acetyl group of compound 3 can be readily removed under acidic conditions to give alcohol 47 in 95% yield. Selective reduction of aryl ketones with NaBH$_4$ led to the formation of secondary alcohol 48 in 80% yield. Furthermore, nucleophilic substitution of 3 with benzylamine afforded γ-fluoroalkylated amine 49 in 70% yield. γ-Fluoroalkylated thioether 50 was obtained in 56% yields through a β-elimination/Michael addition reaction with 4-methoxy-a-toluenethiol.

Proposed mechanism and mechanistic studies. Our proposed mechanism for this Ni-catalyzed reductive coupling is outlined in Fig. 5a. Oxidative addition of the active Ni(0) species I to acyl chloride affords Ni$^{II}$ complex II$^{70}$. Concurrently, Ni(0)- or Ni$^{II}$-mediated single-electron reduction of C$_4$F$_9$I generates the electrophilic C$_4$F$_9$ radical, and subsequent radical addition to the alkene coupling partner would deliver alkyl radical species III. At this juncture, we expected that Ni$^{III}$ complex II could be intercepted by nucleophilic alkyl radical III to yield Ni$^{II}$ adduct IV$^{51,52}$, which would undergo reductive elimination to deliver the final product and Ni$^{II}$ species V. Single-electron reduction of Ni$^{II}$ V ($E_{\text{red}}$ [Ni$^{II}$/Ni$^{III}$] = −1.2 V vs SCE in DMF$^{71}$) by Mn dust ($E_{\text{red}}$ = −1.4 V vs SCE in MeCN) would regenerate Ni(0) species I and complete the catalytic cycle. We expected that the chelating group would affect the feasibility and stability of the Ni$^{III}$ complex, thus influencing the reactivity of this reaction. Alternatively, alkyl radical III could be captured by Ni(0) to form Ni$^{II}$ complex VII$^{72}$, followed by oxidative addition of acyl chloride to deliver the crucial Ni$^{III}$ adduct IV.

We have conducted several preliminary mechanistic experiments to elucidate the proposed mechanism (Fig. 5b–d). Addition
starting materials will allow it to enjoy extensive application in the area of organic chemistry.

**Methods**

**General procedure for the carboacylation reaction.** To a flame-dried 8 mL reaction vial was charged with NiCl₂·glyme (0.02 mmol, 10 mol%), 4,4-di-tert-butyl-2,2′-dipyridyl (0.04 mmol, 20 mol%), and Mn (0.6 mmol, 3.0 equiv.). The vial was capped. After it was evacuated and backfilled nitrogen three times, CH₂CN [0.1 M] was added via a syringe, followed by the addition of acyl chloride (0.3 mmol, 1.5 equiv.). The reaction mixture was allowed to stir for approximately 1 min before fluoroalkyl iodide (0.2 mmol, 1.0 equiv., if liquid) and alkene (0.2 mmol, 1.0 equiv. if liquid) were added. The reaction mixture was allowed to stir at 1500 rpm for 20 h at 25 °C. The reaction was quenched with 1 N HCl, extracted with ethyl acetate (EA) three times. The combined organic layers were dried with MgSO₄, filtered, and concentrated in vacuo. The crude material was purified by flash chromatography to afford the product. See Supplementary Methods for further experimental details.

**Data availability.** The authors declare that all the data supporting the findings of this work are available within the article and its Supplementary Information files or from the corresponding author upon request.

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Author contributions
X.Z. and H.Y.T. contributed equally to this work. L.C. conceived and designed the project. L.C., X.Z., H.Y.T., L.G., S.Z., and F.L.Q. designed the experiments. X.Z., H.Y.T., L.G., and S.Z. performed the experiments and analyzed the data. L.C. prepared the manuscript.

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