Photoinduced copper-catalyzed asymmetric C-O cross-coupling

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Construction of carbon-heteroatom bonds is one of the most active areas of research in organic chemistry, because the function of organic molecules is often derived from the presence of heteroatoms. Whereas considerable advances have recently been achieved in radical-involved catalytic asymmetric C-N bond formation, there has been little progress in the corresponding C-O bond-forming processes. Here we describe a photoinduced copper-catalyzed cross-coupling of readily oxime esters and 1,3-dienes to generate diversely substituted allylic esters with high regio- and enantioselectivity (>75 examples; up to 95% ee). The reaction proceeds at room temperature under excitation by purple light-emitting diodes and features the use of a single, earth-abundant copper-based chiral catalyst as both the photoredox catalyst for radical generation and the source of asymmetric induction in C-O coupling. Combined experimental and DFT computational studies suggest the formation of π-allyl copper complex from redox-active oxime esters as bifunctional reagents and 1,3-dienes through a radical-polar crossover process.

The scaffold of organic compounds mainly consists of carbon-carbon bonds, but their functions are often derived from the presence of various heteroatoms. In medicinal chemists’ toolbox, the assembly of carbon-heteroatom bonds accounts for almost half of all top ten major categories of chemical reactions employed in the drug discovery$^{1,2}$. Not surprisingly, the development of more efficient and sustainable methods for the construction of carbon-heteroatom bonds, particularly the C-N and C-O bonds, continues to attract considerable modern synthetic efforts. With the evolvement of radical chemistry and transition-metal catalysis, radical-involved cross-coupling are emerging as a particularly straightforward strategy for construction of various carbon-heteroatom bonds$^{3-6}$. The remarkable findings of Fu and Peters demonstrate that photoinduced asymmetric copper catalysis provides a robust and reliable method for enantioselective C-N coupling of racemic tertiary alkyl chlorides with amines under mild conditions (Fig. 1)$^{7,8}$. Subsequent works of Liu$^{9,11}$ and Liu$^{12}$ showed that certain chiral ligands (phosphoric acid, bisoxazolines) and directing groups substantially improved the copper’s catalytic activity, and opened up a much broader substrate scope of radical precursors and N-nucleophiles. Notably, MacMillan’s$^{13}$ organocatalytic photoredox-based approach, Zhang’s Co(II)-based metalloradical catalysis$^{14}$ and Nagib’s$^{15}$...
radical relay chaperone strategy also provided a range of complementary strategies for direct enantioselective N-centered radical-mediated sp³ C-N bond formation. Despite these considerable advances, there has been little progress in corresponding radical-involved C-O bond-forming processes. With several recent noteworthy exceptions of alkene-based intramolecular enantioselective C-O bond-forming radical cylications from the groups of Buchwald\(^\text{16,17}\) and Liu\(^\text{18}\), the development of new catalytic systems for radical-involved catalytic asymmetric intermolecular C-O coupling remains an unsolved and long-standing challenge\(^\text{19,20}\).

<table>
<thead>
<tr>
<th></th>
<th>sp³ C-N cross-coupling</th>
<th>sp³ C-O cross-coupling</th>
</tr>
</thead>
<tbody>
<tr>
<td>intermolecular</td>
<td>sp³ C-radical + amines(^8,9)</td>
<td>sp³ C-radical + alcohols(^19)</td>
</tr>
<tr>
<td></td>
<td>sp³ C-radical + azides(^12)</td>
<td>only a single example with enamides rarely explored</td>
</tr>
<tr>
<td></td>
<td>N-radical + sp³ C-H(^13)</td>
<td></td>
</tr>
<tr>
<td>intramolecular</td>
<td>sp³ C-radical + thioureas(^9-11)</td>
<td>sp³ C-radical + carboxylic acids(^16,17)</td>
</tr>
<tr>
<td></td>
<td>sp³ C-radical + imidate(^15)</td>
<td>sp³ C-radical + oximes(^18)</td>
</tr>
<tr>
<td></td>
<td>metalloradical + sp³ C-H(^14)</td>
<td></td>
</tr>
</tbody>
</table>

**Fig. 1 | Qualitative overview on the radical-involved sp³ C-N and sp³ C-O cross-coupling.**

The copper-catalyzed asymmetric allylic substitution provides a less expensive and potentially equally efficient approach for construction of valuable enantiopure allylic compounds compared to the dominant Pd, Rh, and Ir-based catalytic systems\(^\text{21,22}\). The key intermediates—the \([\sigma+\pi]\)-allyl copper(III) or \(\pi\)-allyl copper(III) complexes—are mostly formed from nucleophilic organometallic species and electrophilic allylic systems having a leaving group through ionic mechanisms (Fig. 2a)\(^23\). Despite powerful, however, the use of highly reactive and moisture sensitive reagents (e.g., organomagnesium, organolithium, organoaluminum, organozinc, or the milder organoboron reagents) and prefuctionalized allylic systems considerably limited the substrate range. Notably, application of this strategy to the heteroatom nucleophiles as coupling partners has not been realized in copper-catalyzed asymmetric allylic alkylation. Recent advances in copper-catalyzed photoredox and radical chemistry have shown that photoreexcited copper catalysts can not only generate highly reactive radical species from various redox-active precursors but also trap these open-shell species to enable ensuing asymmetric coupling\(^24-26\). Liu has shown that copper(II)-bound N-centered radical could cleavage the allylic sp³ C-H bond of alkenes in a site-specific manner via hydrogen atom transfer to generate allylic radicals\(^27\). This method allows site- and enantioselective allylic C-H cyanation. Quite recently, the groups of Glorius\(^28-32\), Gevorgyan\(^33\), and Shi\(^34\) demonstrated that alkyl radicals, which are generated from N-hydroxyphthalimide esters or alkyl halides under photoredox catalysis, can be sequentially trapped by 1,3-dienes and palladium, chromium, or titanium catalyst to generate the corresponding electrophilic or nucleophilic \(\pi\)-allylmethal complexes. Employing these intermediates, they achieved a range of aminoaalkylation and dialkylation reactions of 1,3-dienes. However, in most of these transformations, the control of both regio- and stereoselectivity of these radical transformations remains a formidable challenge. Drawing inspiration from these breakthroughs and based on our long standing interest in photoinduced copper catalysis\(^35\), we questioned whether \(\pi\)-allyl-copper complexes could also be formed by a radical pathway to surmount the aforementioned limitations often encountered in traditional copper-catalyzed ionic asymmetric allylic substitution (Fig. 2b). Specifically, we hypothesized that a photorexcited Cu(I) complex might reduce a redox-active radical precursor via a single electron transfer (SET) process to generate a radical species and a Cu(II) species. Then, interception of such a radical species by a feedstock 1,3-diene would give a new allyl radical that might be readily trapped by the initially formed Cu(II) species to generate
π-allylcopper(III)/nucleophile complex in the presence of a nucleophilic reagent. Finally, this high-valent Cu(III) species is prone to undergo reductive elimination to form a new C-O chemical bond with concomitant formation of Cu(I) catalyst. To date, exploration of this strategy in the development of an asymmetric allylic sp³ C-O coupling, particularly the intermolecular variant, has not yet been documented.

Given their readily availability from ketones and carboxylic acids as well as susceptibility to SET reduction, oxime esters have been widely used as cyanoalkyl radical precursors in various radical cross-coupling reactions. Based on our design plan (Fig. 2b), we hypothesized that π-allylcopper(III) complex might be generated by combining oxime esters and 1,3-dienes under photoinduced copper catalysis. However, at the outset of this study, we were not sure oxime esters can serve as bifunctional reagents as both radical sources and O-nucleophile, or whether the regio- and stereocontrol can be realized by using a chiral ligand in such a radical-polar crossover process. Here we describe a radical strategy for formation of π-allylcopper(III) complex and achievement of its highly regio- and enantioselective asymmetric C-O cross-coupling under photoinduced copper catalysis (Fig. 2c). The protocol lies at the intersection of several significant fields of modern chemical catalysis. The obtained enantioenriched 1,2-cyanoalkylesterification products of 1,3-dienes are important building blocks in both synthetic and medicinal chemistry.

**Fig. 2 | Strategic development for formation of π-allylcopper complexes and new reaction design.**

- **a** Previous ionic pathway for generation of allylcopper complexes (well established)

- **b** Radical pathway for generation of allylcopper complexes (unexplored)

- **c** Conceptual design and proof

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and enantioselectivity, leading to many valuable chiral allylic esters (>75 examples).

Results and discussion

Reaction development and optimization. On the basis of our conceptual design, we began the study by examining cross-coupling reaction of 1-phenylbutadiene 1a and oxime ester 2a under visible-light-induced copper catalysis (Table 1). The major challenge was to control both regio- (1,2- vs 1,4-adduct) and enantioselectivity. After extensive screening of reaction parameters, we discovered that irradiation of the cross-coupling partners with purple LEDs at room temperature for 2 hours in the presence of Cu(CH$_3$CN)$_4$PF$_6$ (1.0 mol%) and a chiral bisoxazoline ligand L1 (1.2 mol%) gave the best results, affording the desired 1,2-adduct 3aa in 79% isolated yield with 94% enantiomeric excess (ee) (entry 1) (Supplementary Tables S1-5). Notably, the competitive side product 1,4-adduct 4aa was not detected. The reaction efficiency appeared to be sensitive to the solvents and copper salts. For instance, the use of THF and DMF led to significant decrease of yield (entries 2 and 3). While employment of Cu(OTf)$_2$ gave comparable results, the reaction did not work when using CuI, with 2a being fully recovered (entries 4 and 5). The substitution patterns of the ligands have also played an essential role. Compared to ligand L1, ligands L2-L4 resulted in substantially lower reaction efficiency and enantioselectivity (entries 6-8). Interestingly, the reaction that was performed under irradiation of blue LEDs ($\lambda_{\text{max}} = 460$ nm) proceeded sluggishly, with 3aa being obtained in only 5% yield while with 91% ee (entry 9).

Table 1 | Effect of variations in the reaction parameters and conditions

<table>
<thead>
<tr>
<th>Entry</th>
<th>Variation$^a$</th>
<th>Yield (%)$^b$</th>
<th>ee (%)$^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>None</td>
<td>88 (79)$^d$</td>
<td>94</td>
</tr>
<tr>
<td>2</td>
<td>Use of THF, 16 h</td>
<td>21 (33)$^d$</td>
<td>86</td>
</tr>
<tr>
<td>3</td>
<td>Use of DMF, 16 h</td>
<td>0 (99)$^f$</td>
<td>-</td>
</tr>
<tr>
<td>4</td>
<td>Use of Cu$^+$</td>
<td>0 (99)$^f$</td>
<td>-</td>
</tr>
<tr>
<td>5</td>
<td>Use of Cu(OTf)$_2$</td>
<td>81</td>
<td>94</td>
</tr>
<tr>
<td>6</td>
<td>Ligand (R, R)-L2</td>
<td>15 (16)$^f$</td>
<td>9</td>
</tr>
<tr>
<td>7</td>
<td>Ligand (R, R)-L3</td>
<td>61</td>
<td>88</td>
</tr>
<tr>
<td>8</td>
<td>Ligand (S, S)-L4</td>
<td>69</td>
<td>-82</td>
</tr>
<tr>
<td>9</td>
<td>Blue LEDs</td>
<td>5 (63)$^f$</td>
<td>91</td>
</tr>
<tr>
<td>10</td>
<td>No light</td>
<td>0 (99)$^f$</td>
<td>-</td>
</tr>
<tr>
<td>11</td>
<td>No ligand (R, R)-L1</td>
<td>0 (22)$^f$</td>
<td>-</td>
</tr>
<tr>
<td>12</td>
<td>No Cu(CH$_3$CN)$_4$PF$_6$</td>
<td>0 (99)$^f$</td>
<td>-</td>
</tr>
<tr>
<td>13</td>
<td>Added water (0.1 equiv.)</td>
<td>84</td>
<td>94</td>
</tr>
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</table>

$^a$Reaction conditions: 1a (0.3 mmol), 2a (0.2 mmol), Cu(CH$_3$CN)$_4$PF$_6$ (1.0 mol%), ligand L1 (1.2 mol%), 1,2-dichloroethane (DCE) (4.0 mL), 2 x 20 W purple LEDs ($\lambda_{\text{max}} = 390$ nm), at room temperature. $^b$Yields were determined by GC analysis with 1,3,5-trimethoxybenzene as the internal standard. $^c$Determined by HPLC analysis on a chiral stationary phase. $^d$Isolated yield. $^e$Recovery of 2a.

Control experiments established that each component is critical to the desired cross-coupling reaction.
(entries 10-12). Remarkably, without ligand L1, the majority of oxime ester 2a was consumed, but only leading to a messy mixture. Our concern that allylic radical initially formed upon radical addition might isomerize to oxidation labile benzylic radical to lead to 1,4-adduct appears to be unfounded, as evidenced by our observation of exclusive formation of 1,2-adduct. From a practical point of view, it is worthy of noting that Cu(CH$_3$CN)$_2$PF$_6$ and chiral bisoxazoline ligand are commercially available and the protocol is not moisture-sensitive (entry 13). Until now, the oxime esters have been mainly used a source of cyanoalkyl radical source with generation of a stoichiometric carboxylic anion as byproduct. By contrast, this protocol allows the waste carboxylic anions to be used as O-nucleophiles, leading to atom-economic incorporation of the oxime esters.

**Reaction scope.** With the optimized conditions in hand, we examined the substrate scope with respect to the two coupling partners on a 0.2 mmol scale. First, the scope of oxime esters was tested by variation of the O-acyl moiety (Table 2a). Aside from 2a, aryl carboxylic acid-derived oxime esters 2b-h bearing various netural (H), electron-donating (e.g., Me, t-Bu, OMe) or electron-withdrawing (e.g., Cl, Br, I, CO$_2$Me) groups at the para-position of the aromatic ring are all capable partners for the cross-coupling. The desired products 3ab-ah were obtained in generally good yields (62-78%) with excellent enantioselectivity (91-93% ee). Notably, the halogen atoms (Cl, Br, I) may be subjected to further synthetic manipulations (in particular transition metal-catalyzed cross-coupling reactions) at the halogenated positions (products 3ae-ag). Moreover, the reaction of oxime esters 1i-k with functional groups at the meta- or ortho-position also reacted well, affording 3ai-ak in 73-80% yields with 91-94% ee, indicating that the steric hindrance did not affect the transformation. Reactions with substrates having a fused aromatic ring (2l) or heteroaromatic ring (2m-p) were also amenable to this process, giving the corresponding coupled products 3al-ap in good yields with consistently excellent enantioselectivity (89-94% ee). Then, we proceeded to examine the generality of this asymmetric cross-coupling by using a range of aliphatic carboxylic acid-derived oxime esters. For instance, oxime esters 2q-w derived from cyclobutanone and primary, secondary (cyclohexyl), tertiary acids (e.g., t-butyl and adamantyl acids) as well as α,β-unsaturated acids were all well accommodated under the current catalytic system. Thus, the expected allylic ester products 3aq-aw were isolated in good yields ranging from 54% to 94% with 90-94% ee.

Markedly, as shown in Table 2b, oxime esters 2x-z, and 2aa that are derived from drugs Diclofenac, Febuxostat, Gemfibrozil, and Ciprofibrate were also converted into their corresponding 1,2-cyanoalkylacyloxylation products 3ax, 3ay, 3az, and 3aaa with good yields and satisfactory enantioselectivity, respectively. Naturally occurring acid and amino acid derived oxime esters 2ab-ac were also suitable for the coupling, providing the expected products with good yields and stereoselectivity. These results showed the robustness and utility of the developed reaction.

Finally, we turned our attention to investigating the effect of variation of oxime moiety on the reaction efficiency and enantioselectivity by reacting with 1a (Table 2c). The reaction of 3,3-disubstituted oxime esters 5a and 5b proceeded smoothly to afford the expected products 6aa and 6ab in high yields with 93% and 92% ee, respectively. Sterically more demanding 5c can also be well tolerated to give 6ac with modest results. 2-Methyl substituted oxime ester 5d also participated in the reaction efficiently to produce a good yield of 6ad with high ee values, but with 1:1 d.r. caused by initial non-enantioselective addition of cyanoalkyl radical. Notably, benzocyclobutenone-derived oxime ester 5e also proved to be suitable for the reaction, leading to product 6ae in 75% yield with 92% ee.

**Table 2** | Scope of the oxime esters in photoinduced copper-catalyzed asymmetric C-O cross-coupling

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**Reaction conditions:**

1a (0.3 mmol), 2 (0.2 mmol), Cu(CH$_3$CN)$_2$PF$_6$ (1.0 mol%), ligand (R,R)-L1 (1.2 mol%),

DCE (0.5 M), rt, purple LEDs, 2-24 h.

### a variation of O-acyl moiety

| 3aa | R = Me, 79%, 94% ee | 3ai | R = Me, 73%, 92% ee |
| 3ab | R = H, 71%, 93% ee | 3aj | R = Me, 78%, 92% ee |
| 3ac | R = i-Bu, 62%, 93% ee | 3ak | R = Me, 74%, 91% ee |
| 3ad | R = OMe, 65%, 92% ee | 3al | R = Me, 80%, 94% ee |
| 3ae | R = Cl, 77%, 92% ee | 3am | R = Me, 73%, 91% ee |
| 3af | R = Br, 78%, 92% ee | 3an | R = Me, 75%, 92% ee |
| 3ag | R = I, 72%, 92% ee | 3ao | R = Me, 70%, 94% ee |
| 3ah | R = CO$_2$Me, 74%, 91% ee | 3ap | R = Me, 68%, 94% ee |

### b drug and natural product derivatives

| 3aq | R = Ph, 80%, 94% ee |
| 3ar | R = Me, 78%, 90% ee |
| 3as | R = Me, 94%, 93% ee |
| 3at | R = Me, 85%, 94% ee |
| 3au | R = Me, 68%, 94% ee |
| 3av | R = Me, 62%, 89% ee |

3ax

86%, 79% ee
(from Diclofenac)

3ay

70%, 84% ee
(from Febuxostat)

3ax

76%, 94% ee
(from Gemfibrozil)

3aaa

79%, 94% ee
(from Ciprofibrate)

3aab

84%, 93% ee
(from (E)-Oclade-tri-enoic acid)

3aac

81%, 93% ee
(from Linoleic acid)

3aad

70%, 90% de
(from L-Proline)

### c variation of oxime moiety

| 6aa | R = Me, 87%, 93% ee |
| 6ab | R = Me, 91%, 92% ee |
| 6ac | R = Me, 75%, 92% ee |

6ad, 70% ee (1:1 d.r.)

93% ee, 94% ee

6ae, 78% ee

85% ee (E/Z = 17:1)

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\(^a\)Reaction conditions: 1a (0.3 mmol), 2 (0.2 mmol), Cu(CH$_3$CN)$_2$PF$_6$ (1.0 mol%), ligand L1 (1.2 mol%).
1,2-dichloroethane (DCE) (4.0 mL), 2x20 W purple LEDs (λ<sub>max</sub> = 390 nm), at room temperature; isolated yields. 9With Cu(CH<sub>3</sub>CN)<sub>4</sub>PF<sub>6</sub> (2.0 mol%), ligand L1 (2.4 mol%), and 3.0 equiv. of 1a.

Next, a representative set of 1,3-dienes were evaluated to further investigate the generality of this cross-coupling (Table 3). First, it was found that the 1-aryl and heteroaryl substituted 1,3-dienes were all competent reaction partners. For example, substrates 1b-g bearing either alkyl or electron-withdrawing groups (e.g., F, Cl, Br, CF<sub>3</sub>) at the para-position of the phenyl ring were well tolerated, furnishing the corresponding allylic esters 7ba-g in 60-73% yields with 87-92% ee. Again, as shown in the reactions of 1h-m, the variation of the substitution pattern and steric hindrance of the aromatic ring has little effect on both the reaction efficiency and enantioselectivity. As such, the expected products 7ha-ma were isolated in 70-80% yields with 87-95% ee, regardless of the electronic properties of the substituent. The reaction of 1-(2-naphthyl)-substituted 1,3-diene 1n also proceeded smoothly to produce 7na in 53% yield with 88% ee. Additionally, 1,3-dienes 1o and 1p with 3-furanyl or 3-thiophenyl were well tolerated, furnishing the relative products 7oa and 7pa in good yields with 88% and 91% ee, respectively. Notably, as shown in the cases of 1q-s, 1,1-disubstituted 1,3-dienes and those bearing a methyl group at C2 or C4 position were also proven to be suitable coupling partners. The expected products 7ra and 7sa were obtained with good yields and satisfactory enantioselectivity, while 7qa was isolated with moderate yield due to low conversion. The 1:1 d.r. of 7sa was caused by the initial non-enantioselective radical addition of the cyanoalkyl radical.

Challenging alkyl-substituted 1,3-dienes such as 1t and 1u could also participate in the desired cross-coupling. In both cases, however, mixtures of 1,2- and 1,4-adducts were isolated in good yields but with moderate regio- and enantioselectivity, probably because of the sterically similarity at C-2 and C-4 positions of such type of 1,3-dienes. We are currently investigating modifications to the chiral ligand, which will provide better stereocontrol for these alkyl-substituted 1,3-dienes. X-ray crystallographic analysis of product 7fh, produced in the reaction of oxime ester 1f and 2h with 77% yield and 92% ee, confirmed the (R)-configuration of the newly formed absolute stereochemistry. Thus, all other cross-coupling products were tentatively assigned by analogy with 7fh. Remarkably, biologically relevant molecule, Febuxostat-derived 1,3-diene 1v, could also couple with 2a efficiently to give the desired product 7va in 52% yield with 84% ee.

Table 3 | Scope of 1,3-dienes in photoinduced copper-catalyzed asymmetric C-O cross-coupling
According to our design plan, an allylic radical might be involved during the formation of the π-allylcopper complex, which was supposed to be formed upon addition of cyanoalkyl radical to 1,3-diene. Thus, we envisioned that if 2-styrenyl substituted cycloketoxime esters could undergo SET-reduction-mediated ring-opening to give allylic radical under the current catalytic system; then an asymmetric C-O cross-coupling might also occur between such allyl radical species and the leaving
group-carboxylic anion. As such, the reaction would provide an atom-economic process for catalytic asymmetric fragmentation/rearrangement of cycloketoxime esters. As highlighted in Table 4, the reaction indeed worked well and the standard catalytic system showed broad substrate scope. A range of aryl carboxylic acid-derived cycloketoxime esters 8a-g with an electron-donating or electron-withdrawing substituent at the para-, meta-, or ortho-position of the phenyl ring all underwent the desired reaction smoothly. The corresponding products 9a-g were obtained with generally good yields (73-90%) and excellent enantioselectivity (90-93% ee). Moreover, substrates 8h and 8i containing (hetero)arene annulated systems and 8j with 2-thiophenyl group were also suitable for the reaction, with the relative products 9h-j being formed with comparable results. As shown in the synthesis of 9j-n, primary (8k) and secondary (8l), as well as sterically very demanding carboxylic acid-derived cycloketoxime esters (8m and 8n) can also be well tolerated without loss of reaction efficiency and enantioselectivity (91-93% ee). We also briefly examined the effect of the R² group on the reaction. As shown in the reactions of 8o-s with R² being aryl or heteroaryl group, though the change of their electronic properties has some effect on the reaction efficiency, the expected products 9o-s could still be obtained with high to excellent ee values.

Table 4 | Preliminary attempts to achieve photoinduced copper-catalyzed ring-opening asymmetric C-O coupling of cycloketoxime esters

<table>
<thead>
<tr>
<th>Reaction conditions: 8 (0.2 mmol), Cu(CH₃CN)₄PF₆ (2.0 mol%), ligand (R, R)-L1 (2.4 mol%), DCE (0.5 M), rt, purple LEDs, 0.5-4 h</th>
<th>Cu(CH₃CN)₄PF₆ (2.0 mol%), ligand (R, R)-L1 (2.4 mol%), DCE (0.5 M), rt, purple LEDs, 0.5-4 h</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>R² = Ph</td>
<td>R² = Me</td>
</tr>
<tr>
<td>R¹ = R = H, 76%, 91% ee</td>
<td>R¹ = R = Me, 90%, 92% ee</td>
</tr>
<tr>
<td>9a</td>
<td>9b</td>
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<tr>
<td>9c</td>
<td>9d</td>
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<td>9e</td>
<td>9f</td>
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<td>9j</td>
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<td>9k</td>
<td>9l</td>
</tr>
<tr>
<td>9m</td>
<td>9n</td>
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<tr>
<td>82%, 92% ee</td>
<td>81%, 91% ee</td>
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<tr>
<td>75%, 90% ee</td>
<td>76%, 91% ee</td>
</tr>
<tr>
<td>83%, 92% ee</td>
<td>80%, 93% ee</td>
</tr>
<tr>
<td>88%, 93% ee</td>
<td>71%, 91% ee</td>
</tr>
</tbody>
</table>

Synthetic applications. A 5.0 mmol scale reaction of 2a also worked well to give comparable results (3aa, 1.22 g, 73% yield, 93% ee), indicating the scalability of the process (Fig. 3a). Moreover, the alkene moiety of product 3aa allows further synthetic manipulation. For example, the carbon-carbon double bond in 3aa could undergo hydrogenation easily to give 10 in good yield without loss of enantiomeric selectivity (Fig. 3b). Irradiation of 3aa with blue LEDs in the presence of photocatalyst Ir[dp(F,C,F,ppy)₂(db-bpy)]PF₆ enabled isomerization of (E)-3aa to its (Z)-form 11 without a deleterious effect on the enantiopurity (Fig. 3c).
**Mechanistic investigation.** First, we obtained UV-Vis absorption spectra of each reaction component, and confirmed that Cu(CH$_3$CN)$_4$PF$_6$/L1 complex was the only visible-light-absorbing species. Stern-Volmer studies of Cu(CH$_3$CN)$_4$PF$_6$/L1 with oxime ester 2a supported that a photoexcited Cu(I) complex could reduce 2a by SET (see Supplementary Fig. S7). A study of the effect of the enantiopurity of ligand L1 on the enantioselectivity of the product 3aa demonstrated an obvious linear effect (see Supplementary Fig. S8). This observation suggested that the active catalyst species for the chiral induction in C-O cross-coupling was consistent with a 1:1 ratio of the copper catalyst and chiral bisoxazoline ligand L1.

To gain some more insights into the mechanism, we then carried out a series of control experiments (Fig. 4). When commonly used radical scavengers, 2,2,6,6-tetramethyl-1-piperidinyloxy (TEMPO) and PhSeSePh, were added to the model reaction of 1a and 2a under the standard conditions, the formation of desired product 3aa was completely inhibited in both cases. And the radical trapping adducts 12 and 13 could be detected by high-resolution mass spectrometry (HRMS) or isolated in quantitative yield (Fig. 4a). These results suggest the possible involvement of cyanoalkyl radical 2a-B that should be formed from the iminyl radical 2a-A through its β-C-C bond cleavage.$^{43}$ In addition, when the radical clock substrate 14 was reacted with 2a, the cyclopropyl group was completely opened to give product 15 in modest yield with 83% ee (Fig. 4b). Thus, this radical clock experiment also provides further evidence of a radical process. To understand the behavior of the O-nucleophile-carboxylic anion-formed upon SET reduction of oxime ester, crossover experiments with two different oxime esters were performed (Fig. 4c). When equimolar amounts of oxime esters 2h and 5a were employed, four cross-coupling products were isolated with similar yields and enantioselectivity. In particular, the formation of formal three-component cross-coupled products 3aa and 16 implied that copper-catalyzed C-O bond-forming step should occur through out-of-cage coupling of radical intermediate and carboxylic anion.

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**Fig. 3 | Synthetic utility.** a, Gram-scale synthetic capability. b, Pd/C-catalyzed reduction of the carbon-carbon double bond. c, Photoinduced isomerization of carbon-carbon double bond.
To further obtain a deeper understanding of such photoinduced copper-catalyzed asymmetric C-O cross-coupling, several elementary steps involving regioselectivity and stereoselectivity were evaluated through density functional theory (DFT) calculations at the SMD\(^{46}\)(DCE)/M06\(^{47}\)/[6-311++G(d,p)]/SDD\(^{48}\) (Cu) level. According to our previous study\(^{43}\), the photoexcited state *Cu(I) complex can be oxidatively quenched by oxime ester (2a) via a SET process to generate a radical anion (2a') and a ground-state Cu(II) cation. As shown in Fig. 5a, the resultant radical anion 2a' undergoes...
three key elementary steps, N–O bond cleavage, β–C–C bond cleavage, and radical addition to 1a, to provide the allyl radical (1a-A). TS1, TS2 and TS3 are the corresponding transition states for these three steps with the Gibbs activation energies (ΔG°*) of 7.1, 8.5 and 5.1 kcal mol⁻¹, respectively. In TS1, the N–O bond is elongated to 1.62 Å from 1.39 Å in 2a’, which indicates the breaking of N–O bond is in progress. When the N–O bond breaks, the spin densities of the resultant N–O cleaved intermediate 2a” is mainly localized on N with 0.94. Then, the iminyl radical (2a-A) is dissociated from 2a” with a negative Gibbs free energy change (ΔG°*) of –2.4 kcal mol⁻¹. In TS2, β–C–C bond of 2a-A is activated to 1.99 Å from 1.54 Å in 2a-A, which means the cyanoalkyl radical (2a-B) is forming. When the β–C–C bond is cleaved, the spin density becomes mainly localized on the terminal C4 of 2a-B with 1.04. In TS3, the C1’–C4 distance is shortened to 2.47 Å, which indicates the C1’–C4 bond is gradually forming. When 2a-B attacks the terminal position of 1-phenylbutadiene (1a), the spin densities become mainly localized on the C2’ and C4’ of radical intermediate 1a-A with 0.50 and 0.57, respectively. Overall, the allyl radical formation occurs with a small energy barrier of 8.5 kcal mol⁻¹ and a negative ΔG°* value of –87.5 kcal mol⁻¹. Subsequently, as shown in Fig. 5b, two possible C–O bond-forming reductive eliminations have been investigated: one is the C2’–O reductive elimination from the stable π-allylcopper complex D-Iₘ to provide the 1,2-adduct (R)-3aa (path a), whereas the other is the C4’–O reductive elimination from π-benzylcopper complex D-IIₘ to obtain 1,4-adduct (R)-4aa (path b). TS4aₘ and TS4bₘ are the corresponding transition states for these two steps with the ΔG°* values of 6.7 and 9.6 kcal mol⁻¹, respectively. The C2’–O distance is shortened to 2.03 Å in TS4aₘ from 2.82 Å in D-I, while the C4’–O distance is shortened to 2.08 Å in TS4bₘ from 3.08 Å in D-IIₘ, which indicates the bond formations of C2’–O and C4’–O are in progress. The former path a occurs much more favorably by 2.9 kcal/mol than the latter path b due to the stable π-allylcopper interaction. Thus, the 1,2-adduct is a major product both in thermodynamics and kinetics, which is in line with the regioselectivity observed experimentally. In order to disclose the origin of high enantioselectivity, two favorable enantio-determining C2’–O reductive eliminations leading to (R)-3aa and (S)-3aa have been located, as shown in Fig. 5c. Starting from Cu(II) benzoate complex C, the allyl radical 1a-A is electrophilically captured by the Cu(II) center from the Re-face and Si-face to generate R-type and S-type carbon-centered chiral Cu(III) intermediates D-Iₘ and D-Iₙ, respectively. D-Iₘ is more stable than D-Iₙ by 5.1 kcal mol⁻¹. Then, the C2’–O reductive eliminations of D-Iₘ and D-Iₙ occurs through TS4aₘ and TS4aₙ to provide the asymmetric C–O cross-coupling products (R)-3aa and (S)-3aa. TS4aₘ leading to R-configured product is lower than TS4aₙ leading to S-configured product by 4.2 kcal mol⁻¹, which is qualitatively in agreement with the observed 94% ee ((R)-3aa) (Fig. 5d). To understand well the reason why the R-configured product shows excellent enantioselectivity, further structural analysis from the perspective of the steric effect suggests that the steric hindrance between the aryl group of chiral ligand and the carboxylic group of substrate may be responsible for determining the enantioselectivity of R-configured product.
Fig. 5 | Results of density functional theory (DFT) calculations (all values are in kilocalories per mole). a, The process of allyl radical formation. b, Formation of the allycopper complex and the regioselectivity of the C-O cross-coupling reaction. c, The enantioselectivity of the C-O cross-coupling reaction. d, Optimized structures of enantio-determining transition states.

To shed more light on the intermediacy of radical intermediates, EPR experiments were also performed (Fig. 6a). It was found that a solution of $2a$ and $\text{L1/Cu(I)(CH}_3\text{CN)}_2\text{PF}_6$ in DCE show obvious Cu$^\text{II}$ signals under visible light irradiation (Fig. 6a (i)). An iminyl radical trapping adduct $2a$-$\text{A-I}$ was also probed by adding radical trap 5,5-dimethyl-1-pyrroline-N-oxide (DMPO) (Fig. 6a (ii)). These results indicate the involvement of iminyl radical and its formation can only occur through a photoinduced copper(I)-catalyzed SET process.

Based on these mechanistic studies, we proposed a potential mechanism for this transformation with the model reaction as an example (Fig. 6b). First, irradiation of a $\text{L1Cu(I)X}$ complex (A) by purple light could lead to its excited state [L1Cu(I)X]* (B), which would then engage in a single electron transfer (SET) process with redox-active oxime ester $2a$ to generate an iminyl radical $2a$-$\text{A}$ and $\text{L1Cu(II)}$ species with release of carboxylic anion ($R_1^1\text{CO}_2^-$). Meanwhile, coordination of the waste carboxylic anion ($R_1^1\text{CO}_2^-$) with the $\text{L1Cu(II)}$ species could result in a copper-nucleophile complex (C). Next, iminyl radical $2A$-$\text{a}$ undergoes facile β-C-C bond cleavage to form cyanoalkyl radical $2a$-$\text{B}$ that adds to the terminal position of diene $1\text{a}$ to furnish radical species $1\text{a}$-$\text{A}$. At this stage, radical $1\text{a}$-$\text{A}$ might associate with Cu(II) complex C to form π-allyl Cu(III) complex D (specifically energetically favorable D-I$^\text{R}$ as shown in Fig. 5$^{50}$). However,
**Conclusion**

In conclusion, we have developed a photoinduced copper-catalyzed intermolecular asymmetric C-O cross-coupling that enables overall transformation of readily available oxime esters and 1,3-dienes to the corresponding diversely substituted chiral allylic esters with high regio- and enantioselectivity. This redox-neutral reaction proceeds at room temperature under visible light irradiation, and features the use
of a copper-based chiral catalyst as both photoredox catalyst for radical generation and the source of asymmetric induction in C-O cross-coupling. Key to the success of this transformation is the establishment of a general strategy for formation of π-allylcopper complex through a radical pathway under visible light irradiation. This strategy provides an alternative to the in situ-generation of π-allylcopper(III) complexes that are typically from nuleophilic organometallic reagents and pre-functionalized allylic systems. Overall, considering that photochemistry can generate various reactive intermediates that would otherwise be difficult to produce under thermal conditions, many exciting extensions of this radical strategy to other related catalytic asymmetric cross-couplings can be envisaged.

Methods

**General procedure for the synthesis of products 3 and 7.** In a flame-dried 10.0 mL Schlenk tube equipped with a magnetic stirrer bar was charged sequentially with Cu(CH$_3$CN)$_4$PF$_6$ (0.75 mg, 0.002 mmol, 1.0 mol%) and L1 (0.86 mg, 0.0024 mmol, 1.2 mol%), followed by the addition of 1,2-dichloroethane (4.0 mL). Then the mixture was stirred at room temperature for 30 min. To the resulting mixture were added 1 (0.30 mmol, 1.5 equiv.) and 2 (0.20 mmol, 1.0 equiv.) under Ar. The mixture was stirred at a distance of ~1 cm from two 20 W Kessil purple LEDs at rt for about 2-24 h until the reaction was completed, as monitored by TLC analysis. The residue was purified by acidic or neutral alumina chromatography with petroleum ether and ethyl acetate (PE/EA) (10:1, v/v) to afford 3 or 7.

**Computational method.** All density functional theory (DFT) calculations were carried out with the Gaussian 16 package. Molecular geometries of all studied systems were optimized at M06[6-31G(d)/LanL2DZ(Cu)] level. Frequency calculations were performed to confirm all the stationary points as local minimum (zero imaginary frequencies) or transition states (one imaginary frequency). Intrinsic reaction coordinate (IRC) calculations were conducted to verify that each transition state actually connects the two corresponding minimum. In addition, the single-point energies of all stationary points were performed at the M06[6-311++G(d,p)/SDD(Cu)] level. The single-point calculations were considered the solvent effects with the SMD model in 1,2-dichloroethane (DCE) solution.

**Preprint.** A previous version of this work was published as a preprint.

**Data availability**

Data relating to the characterization data of materials and products, general methods, optimization studies, experimental procedures, mechanistic studies, mass spectrometry, high-performance liquid chromatography and NMR spectra, computational studies are available in the Supplementary Information.

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**Author contributions**

J.C., P.-Z.W., G.Q.L., H.Q., B.Z., and X.-D.H. are responsible for the plan and implementation of the experimental work. Y.J.L and W.G. are responsible for implementation of the DFT calculations and related writing. J.-R.C. and W.-J.X. supervised the project and wrote the manuscript with revisions provided by the other authors. All authors discussed the results and commented on the manuscript.

**Competing interests**

The authors declare no competing interests.