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# Copper-catalyzed oxidative carbon–heteroatom bond formation: a recent update

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This review updates recent advances in Cu-catalyzed (anaerobic) oxidative carbon–heteroatom bond formation on  $\text{sp}^3$ - and  $\text{sp}^2$ -C–H bonds as well as alkenes, classified according to the types of stoichiometric oxidants.

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# 1. Introduction

Oxidative molecular transformation that incorporates heteroatom units into carbon-based organic scaffolds is one of the most fundamental and important synthetic transformations, enhancing the molecular complexity. Therefore, development of new oxidative synthetic methodologies that convert readily available substrates in lower oxidation states into highly functionalized (oxidized) molecules in a chemo- and stereo-selective manner is a long-standing goal in chemical synthesis. In this context, direct installation of carbon–heteroatom bonds on ubiquitous C-H bonds (both  $sp^3$ - and  $sp^2$ -hybridized) is of great significance to streamline the multi-step molecular REVIEW ARTICLE<br>
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transformations needed for the synthesis of target functional molecules. However, it is challenging to functionalize such C–H bonds selectively unless otherwise adjacent activating groups are installed because of the inherent inert property and the ubiquitous nature of the C–H bonds. On the other hand, oxidative difunctionalization of alkenes provided another efficient method to address highly oxidized molecular complexity through installing two distinct functional groups in a one-pot fashion. Thus, development of chemo-, regio-, and stereoselective difunctionalization of alkenes is the major concern to be addressed in chemical synthesis.

Transition-metals are capable of realizing various state-ofthe-art processes for C–H oxidation<sup>1</sup> and oxidative difunctionalization of alkenes. $<sup>2</sup>$  For the catalysis of choice, ubiquitous first</sup> row transition metals have recently attracted much attention not only as alternatives to precious late transition metals, but also for exploration of their unprecedented catalytic processes.<sup>3</sup>



Xu Zhu

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Shunsuke Chiba

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Scheme 1 Cu-Catalyzed oxidative carbon–heteroatom bond formation on C–H bonds and alkenes.

Among such first row transition metals, copper complexes exhibit unique and versatile reactivity and good functional group tolerance.<sup>4</sup> A broad range of oxidation states of copper complexes (mainly from  $Cu^{0}$  to  $Cu^{III}$  applied in chemical synthesis) $5,6$  enables the promotion of redox reactions in either a single-electron or a two-electron-transfer fashion (or both in the sequential processes), depending on the reaction conditions and types of substrates used. A variety of stoichiometric terminal oxidants have been devised and applied for realizing the catalytic turnover in Cu-mediated oxidative molecular transformation and/or serving as the sources of heteroatoms introduced into the products (Scheme 1). Published on 11 February 2016. Downloaded on 12 February 2016. Downloaded on 12 February 2022 9:12:45 AM. This article is licensed under a common common and the common common and the common common common and the common co

This review focuses on recent advances in copper-catalyzed oxidative carbon–heteroatom bond forming reactions on C–H bonds as well as alkenes. These reactions are classified according to the different types of stoichiometric terminal oxidants employed in the processes. Among the available oxidants in copper-catalyzed oxidative molecular transformation, molecular oxygen  $(O_2)$  has been extensively employed as the terminal oxidant for catalytic turnover, enabling a variety of oxidative reactions. As these achievements are reviewed elsewhere in detail, $\overline{7}$  this review will exclude copper-catalyzed aerobic reactions.

# 2. With I(III) reagents

#### 2.1.  $PhI(OAc)<sub>2</sub>$

The combination of Cu complexes and  $PhI(OAc)$ <sub>2</sub> could generate higher oxidation state Cu<sup>n</sup> species ( $n =$  II or III), which mediate unprecedented single-electron-oxidation processes. Chang reported benzylic/allylic sp<sup>3</sup>-C-H oxygenation with N-hydroxyphthalimide (NHPI) under CuCl-catalyzed-PhI(OAc)<sub>2</sub>mediated reaction conditions (Scheme 2).<sup>8</sup> The radical mechanism is proposed, where the oxidatively formed phthalimide N-oxyl (PINO) radical undergoes H-radical abstraction from the substrates (Sub-H) to generate the corresponding C-radicals (Sub-). Their subsequent recombination with the PINO radical affords the products. The role of  $PhI(OAc)_2$  is to maintain the catalytic cycle by re-oxidizing lower valent Cu species. Interestingly, a radical-clock substrate, cyclopropylmethylbenzene was coupled with the PINO radical keeping the cyclopropyl moiety intact, indicating kinetically faster radical recombination or alternative organometallic mechanism involved.





**Scheme 2** Cu-Catalyzed-PhI(OAc)<sub>2</sub>-mediated sp<sup>3</sup>-C-H oxygenation with **NHPI** 



Scheme 3 Aliphatic C–H amination with amidines under Cu-catalyzed-PhI(OAc)<sub>2</sub>-mediated reaction conditions.

Chiba reported intramolecular aliphatic C–H amination of  $N$ -alkylamidines under Cu(OAc)<sub>2</sub>-catalyzed-PhI(OAc)<sub>2</sub>-mediated reaction conditions (Scheme  $3$ ).<sup>9</sup> The reaction is initiated by generation of amidinyl radical 2 probably through formation of  $Cu(m)$ -amidine intermediate 1 followed by homolysis of the N–Cu bond. A subsequent 1,5-H radical shift<sup>10</sup> of the amidinyl radical affords the corresponding C-radical 3, further singleelectron-oxidation of which with  $Cu(II)$  or  $Cu(III)$  species generates a carbocation and subsequent C–N bond formation to furnish dihydroimidazole. The radical recombination mechanism is not ruled out for C–N bond formation.

The Cu–PhI(OAc)<sub>2</sub> system is capable of oxidizing the aromatic sp<sup>2</sup>-C-H bond with the assistance of appropriate ortho-directing groups. For example, ortho-C–H amination of aniline derivatives was developed using the picolinamide directing group under Cu-catalyzed-PhI(OAc)<sub>2</sub>-mediated reaction conditions (Scheme 4).<sup>11</sup> The single-electron-oxidation of the benzene ring in chelate complex 4 followed by morpholine transfer to cation radical 5 is proposed for the C–H amination.

Various modes of oxidative functionalization of alkenes have been realized using Cu-catalyzed-PhI(OAc)<sub>2</sub>-mediated reaction systems, in which the acetate moiety on  $PhI(OAc)_2$  is incorporated during the process. The mechanisms for alkene functionalization



**Scheme 4** Cu-Catalyzed-PhI(OAc)<sub>2</sub>-mediated directed sp<sup>2</sup>-C-H amination.



Scheme 5 Cu-Catalyzed-PhI(OAc)<sub>2</sub>-mediated aminoacetoxylation of alkenes.

vary with the substrates used. Blakey disclosed intramolecular Cu-catalyzed-PhI(OAc)<sub>2</sub>-mediated aminoacetoxylation of alkenylsulfonamides for synthesis of nitrogen heterocycles (Scheme 5).<sup>12</sup> The mode of cyclization (either *endo* or *exo*) depends on the alkene substituents. The process is proposed to be initiated by electrophilic activation of alkenes by amide– $Cu(m)$  species 6. Subsequently, acetoxy-cupration of alkenes takes place to afford metallacycle intermediate 7, in which more substituted carbon is preferentially acetoxylated. Reductive elimination of the C–N bond finally furnishes the heterocyclic products.

Interestingly, the reactions of alkenylureas under similar reaction conditions give oxyacetoxylation products (Scheme 6).<sup>13</sup>

On the other hand,  $Cu(OAc)_{2}$ -catalyzed-PhI(OAc)<sub>2</sub>-mediated reactions of N-allylamidines afford acetoxymethyl dihydroimidazoles *via* aminoacetoxylation of alkenes (Scheme 7).<sup>14</sup> When  $2,2'$ -methylene bis[ $(4R,5S)$ -4,5-diphenyl-2-oxazoline] 1 is employed instead of 1,10-phenanthroline, chirality induction of 48% ee is observed in aminoacetoxylation. This observation implicates that the process might involve aminocupration of alkenes by the putative amidinyl copper $(m)$  intermediate 8 *via* an organometallic pathway to form an organocopper $(m)$ 



Scheme 6 Cu-Catalyzed-PhI(OAc)<sub>2</sub>-mediated oxyacetoxylation of alkenylureas.



Scheme 7 Cu-Catalyzed-PhI(OAc)<sub>2</sub>-mediated aminoacetoxylation of alkenylamidines.

intermediate that is unlike the aliphatic CH amination of N-alkylamidines involving free radical intermediates (Scheme 3). Finally, nucleophilic displacement of organocopper $(m)$  moiety 9 with an acetate ion gives the final product.

Recently, Buchwald reported Cu-catalyzed enantioselective synthesis of functionalized lactones from alkenylcarboxylic acids through oxyfunctionalization of alkenes as the key step (Scheme 8).<sup>15</sup> For example, oxyazidation is enabled using a catalytic amount of Cu(MeCN)<sub>4</sub>PF<sub>6</sub> with TMSN<sub>3</sub> and PhI(OAc)<sub>2</sub> (Scheme 8). The azido radical and higher valent  $Cu(II)$  species are initially formed by the redox reaction between  $TMSN_3$ ,  $Cu(1)$ complex, and  $PhI(OAc)<sub>2</sub>$ . The resulting azido radical then adds onto alkenes to generate tertiary radical 10 that recombines with the intramolecular Cu<sup>II</sup>-carboxylate moiety in a stereoselective fashion with the chiral Box ligand. Finally, C–O reductive elimination from metallacycle 11 gives the lactone product with regeneration of the  $Cu(i)$  catalyst.



Scheme 8 Cu-Catalyzed-PhI(OAc)<sub>2</sub>-mediated asymmetric oxyazidation of alkenes.



Scheme 9 Cu-Catalyzed oxy-arylation and -vinylation of allylamides with iodonium salts.

#### 2.2. Diaryliodonium salts  $[Ar<sub>2</sub>IX]$

Treatment of Cu(I) complexes with diaryliodonium salts results in formation of aryl-Cu $(m)$  species like 12 in Scheme 9 that could be used for electrophilic activation of alkenes to induce their carbo-functionalization. For example, Gaunt reported that the reaction of N-allylamides could afford oxyarylation products in a diastereoselective fashion through the transient carbocation 13 (Scheme 9). $^{16}$  Similarly, oxyvinylation was enabled by using vinyl(aryl)iodonium salts. The enantioselective variant was also devised using chiral Cu–bisoxazoline complexes as the catalyst.<sup>17</sup>

#### 2.3. The Togni reagents

It has been shown that single-electron-reduction of 1-trifluoromethyl-1,2-benziodoxol-3(1H)-one (known as Togni reagent II) by Cu(I) complexes generates CF<sub>3</sub> radical 14 along with Cu(II) 2-iodobenzoate 15 (Scheme 10). This reductive process could be combined with the subsequent oxidative alkene difunctionalization through addition of the  $CF_3$  radical onto alkenes followed by recombination of the resulting C-radical 16 with external heteroatom sources 17, furnishing the final product



Scheme 10 Cu-Catalyzed trifluoromethylation of alkenes with the Togni reagent.



Scheme 11 Cu-Catalyzed intermolecular oxytrifluoromethylation





and  $Cu(i)$  species. The overall process is thus able to have a catalytic turnover.

For example, intermolecular oxytrifluoromethylation was realized by radical recombination with the  $Cu(II)$  2-iodobenzoate derived from the Togni reagent (Scheme  $11$ ).<sup>18</sup>

Intramolecular C–O bond formation upon trifluoromethylation was enabled by using alkenylcarboxylic acids (Scheme  $12-i$ )<sup>19</sup> and alkenyloximes (Scheme 12-ii), delivering the corresponding trifluoromethylated lactones and isoxazolines, respectively.

Similarly, trifluoromethylazidation (Scheme 13-i)<sup>20</sup> and trifluoromethylthiocyanation (Scheme  $13$ -ii)<sup>21</sup> were reported using  $TMSN<sub>3</sub>$  and  $TMSNCS$ , respectively, as the external heteroatom sources. In the case of trifluoromethylazidation, use of 3,3-dimethyl-1,2-benziodoxole (known as Togni reagent I) provided better yields of the desired azidation products as the reaction with 1,2-benziodoxol-3-one generated 2-iodobenzoyloxylaiton product as the side product.



Scheme 13 Cu-Catalyzed trifluoromethylazidation and trifluoromethylthiocyanation of alkenes.



#### 3. With peroxides

Various types of peroxides have been employed as the stoichiometric oxidants and often as the sources of oxygen functionality in Cu-catalyzed oxidative molecular transformation.<sup>4d</sup> In principle, reduction of peroxides by lower valent  $Cu(I)$  species provides the corresponding higher valent  $Cu(II)$  alkoxide and highly reactive alkoxy radical (Scheme 14) that cooperate synergistically to mediate subsequent Kharasch–Sosnovsky<sup>22</sup> type oxidative functionalization of the substrates. The resulting alkoxy radical could further oxidize  $Cu(1)$  species to give another  $Cu(II)$  alkoxide (path-ii) or undergo H-radical abstraction from the substrates to form the C-radical (path-iii).

#### 3.1. With di-t-butylperoxide (t-BuOOt-Bu)

Warren reported seminal studies on aliphatic C–H amination with simple amines using the well-defined  $Cu(I)$   $\beta$ -diketiminate complex and t-BuOOt-Bu as well as their detailed mechanistic studies by kinetic, spectroscopic, and structural analyses of possible intermediates (Scheme 15).<sup>1e,23</sup> It is conceivable that  $Cu(II)$  amide complex 18 formed by the alkoxy-amide exchange (path-i) undergoes C–H bond amination of alkanes via aliphatic H-radical abstraction and subsequent C–N bond formation with the resulting C-radical (path-ii) along with generation of  $Cu(i)$  species that maintains the catalytic cycle with  $t$ -BuOO $t$ -Bu.

More recently, Hartwig disclosed Cu-catalyzed aliphatic C–H amidation and imidation using t-BuOOt-Bu as the stoichiometric oxidant (Scheme  $16$ ).<sup>24</sup> The reactions prefer to oxidize secondary C–H bonds than primary ones, while tertiary C–H



Scheme 15 Cu-Catalyzed-t-BuOOt-Bu-mediated aliphatic C–H amination.



bonds are interestingly the least reactive. The stoichiometric reaction analyses using the isolated well-defined copper amidate complexes implicated that the C–H amidation is enabled by H-radical abstraction with the t-butoxy radical (path-iii) and radical recombination of the resulting C-radical with transient  $Cu(II)$ amidate complexes (path-iv). Analogous ligand-free Cu-catalyzed aliphatic C-H amidation and imidation with  $t$ -BuOO $t$ -Bu were developed independently by Huang and Yu/Cheng.<sup>25</sup>

This type of Cu-catalyzed-t-BuOOt-Bu-mediated aliphatic C–H oxidation strategy could be further applied for synthesis of tertiary carbamates using isocyanates as the amide source (Scheme 17).<sup>26</sup> The reaction of Cu(I) species with  $t$ -BuOO $t$ -Bu and isocyanate generates  $Cu(n)$ -amide complex 19, which is coupled with the C-radical derived from alkanes via H-radical abstraction by the transient  $t$ -butoxy radical, affording tertiary carbamates.

Intramolecular benzylic C–H alkoxylation of aromatic heterocycles having a hydroxyalkyl tether was devised under Cu-catalyzed $t$ -BuOO $t$ -Bu-mediated reaction conditions (Scheme 18).<sup>27</sup> The intermolecular variant also worked in the same system, while the yields of the C–H alkoxylation products were moderate.



Scheme 17 Cu-Catalyzed-t-BuOOt-Bu-mediated synthesis of tertiary carbamates with isocyanates as the amide source.



Zhu recently reported carboetherification of aryl alkenes with solvent amounts of acetonitrile and alcohol under Cu-catalyzed $t$ -BuOO $t$ -Bu-mediated reaction conditions (Scheme 19).<sup>28</sup> The radical clock experiment implicated that benzylic radical intermediate 20 is formed by the addition of acetonitrile through either carbocupration followed by homolysis of the resulting C–Cu bond or addition of the a-cyanomethyl radical. The final product is delivered through formation of the C–O bond via radical recombination of the benzylic radical intermediate with alcohol.

#### 3.2. With t-butylhydroperoxide (t-BuOOH: TBHP)

t-Butylhydroperoxide (TBHP) exhibits reactivity analogous with that of di-t-butylperoxide for Cu-catalyzed aliphatic C–H oxidation with various heteroatom sources such as amides via radical intermediates (Scheme 20). For example, Guerra recently reported that Cu–Al mixed oxide could be utilized as a heterogeneous catalyst for Kharasch–Sosnovsky type allylic C–H oxygenation with TBHP as the stoichiometric oxidant.<sup>29,30</sup>

In addition, TBHP uniquely serves as an oxygen source for enhancing the oxidation state of the substrates in aliphatic C–H oxidation. Patel developed Cu-catalyzed-TBHP-mediated synthesis of esters from aldehydes and alkylarenes such as toluene (Scheme 21-i), $31$  in which Cu–alkoxides formed *via* benzylic oxygenation with the TBHP couple with aldehydes to deliver esters.



Toluene could serve as a benzoate precursor in synthesis of esters via C-H oxygenation of cyclic ethers (Scheme 21-ii). $32,33$ 

As a mechanistically distinct example of aliphatic C–H oxidation, Loh developed unique a-amination of aldehydes with secondary amines under Cu-catalyzed-TBHP-mediated reaction conditions that provides  $\alpha$ -amino acetals as the products (Scheme 22). $34$  The process is composed of a multi-step sequence including two successive single-electron-oxidations of enamine intermediate 21 derived from condensation of aldehydes and amines. The resulting  $\alpha$ -methoxy aziridinium ion 22 undergoes ring-opening with methanolysis, giving a-amino acetal. The detailed roles of CuI and TBHP in the single-electron-transfer processes are not certain.

The Cu-TBHP system is also capable of oxidizing  $sp^2$ -C-H bonds. Jiao disclosed *ortho-azidation* of anilines with TMSN<sub>3</sub> under Cu-catalyzed-TBHP-mediated reaction conditions at ambient temperature (Scheme  $23$ ).<sup>35</sup> The single-electron-oxidation of the benzene ring by higher valent  $Cu(II)$  species 23 adjacent to the primary amine moiety followed by azido ion transfer in cation radical 24 is proposed for the C–H azidation. This is a rare example of the directed ortho C–H functionalization of anilines.

Benzaldimines 25 derived from condensation of anilines and aldehydes could be used for analogous ortho-azidation and the resulting 2-azidoarylimines 26 undergo subsequent denitrogenative cyclization to give benzimidazoles (Scheme  $24$ ).<sup>36</sup> A 2-pyridyl group has also been utilized as the directing group in aromatic C–H oxidation under Cu-catalyzed-TBHP-mediated reaction conditions.<sup>37</sup>

Bolm developed Cu-catalyzed oxidative N-acylation of sulfoximines with aldehydes (Scheme  $25$ ).<sup>38</sup> The *t*-butoxy radical derived







Scheme 22  $Cu$ -Catalyzed  $\alpha$ -amination of aldehydes





Scheme 24 Cu-Catalyzed-t-BuOOH-mediated aromatic ortho-azidation.



from decomposition of TBHP by Cu(I) complexes (Scheme 25-i) could abstract the H-radical from aldehydes to generate the corresponding acyl radicals 27 (Scheme 25-ii). The transient acyl radicals 27 undergo radical recombination with sulfoximines mediated by higher valent  $Cu(II)$  species to give the products along with re-generation of lower valent Cu(I) species (Scheme 25-iii).

In the presence of  $Cu(i)$  complexes, TBHP can serve as the source of the t-butoxy radical that mediates H-radical abstraction to give the C-radical, whereas TBHP itself can also be introduced as the new oxygen functionality via the C–O bond forming process. Loh demonstrated Cu-catalyzed threecomponent coupling of alkenes, aliphatic alcohols, and TBHP for construction of the corresponding carbooxygenation products. In this process,  $\alpha$ -hydroxy radicals 28 generated from aliphatic alcohols add to alkenes to give secondary radicals 29 that recombine with TBHP mediated by  $Cu(II)$ species (Scheme 26).<sup>39</sup> Patel reported analogous Cu-catalyzed



Scheme 26 Cu-Catalyzed carbooxygenation of alkenes



Scheme 27 Cu-Catalyzed aromatic C–H oxygenation.

three-component coupling of electron-deficient alkenes, cycloalkanes, and TBHP.<sup>40</sup>

#### 3.3. With benzoyl peroxide (BPO)

While benzoyl peroxide (BPO) has rarely been utilized as the partner with Cu-catalyzed oxidative molecular transformation, Martin recently reported Cu-catalyzed intramolecular aromatic C–H oxygenation of 2-arylbenzoic acids specifically mediated by BPO as the terminal oxidant (Scheme  $27$ ).<sup>41</sup> The reaction could not be facilitated by TBHP. Together with treatment of the biaryl lactones with LiOH for hydrolysis, the overall process is considered as formal aromatic C–H hydroxylation.

## 4. With O-benzoyl-N,Ndialkylhydroxylamines

Among various types of hydroxyl amine derivatives, O-benzoyl- $N$ , $N$ -dialkylhydroxylamines (BzO–NR<sub>2</sub>) have been of particular use for electrophilic amination of organocopper $(i)$  species. For example, Hirano/Miura reported direct aromatic C–H amination of electron-deficient arenes with BzO–NE $t_2$  by the  $Cu(OAc)<sub>2</sub>$ -1,10-phen catalytic system in the presence of LiOt-Bu (Scheme  $28$ ).<sup>42</sup> The process is composed of aromatic C-H cupration for the formation of  $\arg l-Cu(i)$  species 30 and subsequent electrophilic amination with  $BzO-NEt<sub>2</sub>$ . The mechanism of electrophilic amination was previously investigated by Johnson and suggested as the  $S_N2$  mechanism.<sup>43</sup> This strategy was applied for C2-amination of quinoline-N-oxides by Li/Wu.<sup>44</sup>

Hirano/Miura demonstrated the combination of borylcupration of alkenes with subsequent electrophilic amination of the resulting alkyl-Cu species with BzO–NR<sub>2</sub>, offering elegant aminoboration of alkenes in stereo- and regio-selective manners. For example, the reactions of arylalkenes such as *trans*- $\beta$ -methylstyrene with bis(pinacolato)diboron (pinB-Bpin) and BzO–NBn<sub>2</sub> under the CuCl-dppbz catalytic system in the presence of



Scheme 28 Cu-Catalyzed aromatic C–H amination with hydroxylamines.





LiOt-Bu resulted in the formation of aminoboration products (Scheme 29).<sup>45</sup> As boryl-cupration of alkenes takes place in synselective and regiospecific fashions to form organocopper intermediate 31 and the subsequent electrophilic amination proceeds with retention of the configuration of the organocopper moiety, the overall stereochemical outcome of the process is regioselective syn-aminoboration of alkenes.

This Cu-catalyzed aminoboration of alkenes was capable of functionalizing bicyclic alkenes (Scheme 30). $46$  The 1,2-aminoborane product from norbornadiene could be further converted into the corresponding diastereomerically pure cyclopentane derivative via hydroxylation of the C–B bond by sodium perborate followed by ring-opening cross metathesis with ethylene.

As for the aminoboration of non-activated terminal alkenes, its regioselectivity could be controlled by switching the ligands



Scheme 30 Cu-Catalyzed aminoboration of norbornadiene and further molecular transformation.



on the Cu( $I$ ) catalysts (Scheme 31).<sup>47</sup> Namely, the CuCl–xantphos system installs the amine moiety at the internal carbon, while the CuBr-N-heterocyclic carbene (IPrCuBr) complex induces the amination at the terminal carbon.

Buchwald, Hirano/Miura, and Hartwig independently reported formal hydroamination of alkenes by the sequence of Cu-catalyzed hydrocupration of alkenes with hydrosilanes and electrophilic amination of the resulting alkylcopper species 32 with BzO–NR<sub>2</sub>. Various types of alkenes such as aryl alkenes,  $48$ unactivated 1,1-disubstituted alkenes,<sup>49</sup> unactivated internal alkenes (Scheme 32),<sup>50</sup> oxa/aza-bicyclic alkenes,<sup>51</sup> and alkenylsilanes $52$  have been employed for regio- and enantioselective hydroamination, in which choice of ligands on copper catalysts is crucial to control the reactions  $(i.e.$  to prevent the side reactions such as hydride reduction of hydroxyl amines).

Wang recently developed Cu-catalyzed diamination of alkenes using alkenyl O-Me-hydroxamic acids and BzO–NR<sub>2</sub> that afforded functionalized nitrogen heterocycles (Scheme 33).<sup>53</sup> The process is composed of intramolecular aminocupration of alkenes and subsequent electrophilic amination. As a deuterium-labeling



Scheme 32 Cu-Catalyzed enantioselective formal hydroamination of non-activated internal alkenes.



Scheme 33 Cu-Catalyzed diamination of alkenyl O-Me hydroxamic acids with O-benzoyl hydroxylamines.

experiment on the terminal alkenyl carbon revealed that the reaction does not retain the original stereochemistry of alkenes, the radical intermediates are supposed to be involved prior to the electrophilic amination.

### 5. With oxaziridines

Highly strained three-membered ring oxaziridines work with copper complexes to facilitate aminooxygenation of alkenes. Yoon developed CuCl<sub>2</sub>-catalyzed aminooxygenation of alkenes with N-sulfonyloxaziridines that could be dramatically facilitated by chloride additives such as  $Bu_4NCl$  (Scheme 34).<sup>54</sup> The detailed mechanistic investigation elucidated that  $Cu(II)$ –oxaziridine complex 33 undergoes C–O bond forming radical addition onto alkenes to generate C-radical intermediate 34 tethered with a  $Cu(m)$  sulfonamide moiety. Subsequent radical recombination forms the C–N bond and regenerates  $Cu(II)$  species that can maintain catalytic turnover further. The presence of the radical intermediate was proved by a radical clock experiment. Open Sec. Rev. Sec. 2013. Continues are the model of the second under the second of the second of the second under the second under a creative Commons are the second under the second under the second under the second unde

This aminooxygenation strategy with N-sulfonyl oxaziridines was capable of functionalizing indoles (Scheme 35).<sup>55</sup> It is worthy to note that the resulting aminal product derived from N-acyltryptamine was readily transformed to 3-aminopyrroloindoline by base treatment.



Scheme 34 CuCl<sub>2</sub>-catalyzed aminooxygenation of alkenes with N-sulfonyloxaziridines.



Scheme 35 Cu-Catalyzed aminooxygenation of indoles with N-sulfonyloxaziridines.

The  $Cu(<sub>II</sub>)$ -oxaziridine complexes also undergo the remote H-radical abstraction to enable aliphatic C–H amination (Scheme 36).<sup>56</sup> The reaction of N-sulfonyl oxaziridines having an alkyl tether under the CuCl<sub>2</sub>–LiCl catalytic system provides intramolecular C–H amination products via 1,6-H-radical abstraction by  $Cu(II)$ -oxaziridine complex 35 followed by subsequent radical recombination of the resulting C-radical 36 to form the C–N bond. The resulting hemiaminal product could be converted into cyclic enamide by acid treatment.

On the other hand, Aubé recently reported Cu(I)-catalyzed allylic  $sp<sup>3</sup>$  C–H oxygenation with *N*-alkyl oxaziridines (Scheme 37).<sup>57</sup> This method could oxidize the allylic position via the sequence of (1) formation of aminyl radical  $37$  having a Cu(II)-alkoxide tether through single-electron-reduction of N-alkyl oxaziridines by the  $Cu(1)$  complex; (2) generation of the allylic radical by a 1,5-H radical shift to form C-radical 38 and subsequent radical recombination with the Cu $(n)$ –alkoxide moiety to form cyclic hemiaminal 39 with regeneration of  $Cu(i)$  species; (3) hydrolysis to form the final product, y-hydroxy ketone.

#### 6. With diaziridinone derivatives

Similar to oxaziridines, N,N-di-t-butyldiaziridinone shows straindriven oxidative reactivity towards Cu(I) complexes, enabling catalytic diamination of various types of alkenes.<sup> $4b$ </sup> Shi revealed



Scheme 36 Cu(II)-catalyzed C-H amination with N-sulfonyloxaziridines.



that  $N$ , $N$ -di-t-butyldiaziridinone oxidizes Cu( $i$ ) complexes to form an equilibrium mixture of four-membered  $Cu(m)$  species 40 and  $Cu(II)$ –N radical species 41 (Scheme 38).

Of particular interest is that the regioselectivity in diamination of conjugated dienes could be switched by the choice of Cu(I) catalyst systems and electronic nature of dienes with the different reaction mechanisms.<sup>58</sup> Namely, conjugated dienes and N,N-di-t-butyldiaziridinone in the presence of a catalytic amount of CuBr generally undergo diamination of internal alkenes, whereas terminal alkenes could be functionalized under the CuCl–phosphine ligand catalytic system (Scheme 39).

Detailed mechanistic investigation suggested that internal diamination proceeds with the four-membered  $Cu(m)$  species 42 via (1) coordination and migratory insertion to dienes; (2) C–N reductive elimination that renders the overall process cis-diamination (Scheme 40-i). On the other hand, diamination of terminal alkenes involves  $Cu(II)$ –N radical species 43



Scheme 38 The reaction of di-t-butyldiaziridinone with Cu(I) complexes.



Scheme 39 Regioselectivity in Cu(I)-catalyzed diamination of conjugated dienes with N,N-di-t-butyldiaziridinone.



that initiates radical C–N bond formation to the sterically less hindered terminal carbon to generate an allyl radical. The second C–N bond formation is enabled by the radical recombination of the allyl radical with an  $N-Cu(n)$  moiety to afford the diamination product along with regeneration of the Cu(I) catalyst (Scheme 40-ii).

This stepwise radical-mediated diamination of alkenes with  $N$ , $N$ -di-t-butyldiaziridinone by the Cu( $I$ )–phosphine ligand catalytic system is amenable to functionalize not only conjugated dienes but also 1,1-disubstituted alkenes.<sup>59</sup> Asymmetric terminal diamination of conjugated dienes was also developed by the CuCl-chiral phosphine<sup>60</sup> and Cu( $I$ )-chiral phosphate systems.<sup>61</sup> Analogous to N,N-di-t-butyldiaziridinone, N,N-di-tbutylthiadiaziridine  $1,1$ -dioxide<sup>62</sup> and  $N,N$ -di-t-butyl-3-(cyanimino)-diaziridine<sup>63</sup> could be utilized for catalytic radical diamination of conjugated alkenes under the CuCl–phosphine ligand systems (Scheme 41).

In addition to diamination of alkenes, N,N-di-t-butyldiaziridinone could be utilized for Cu(I)-catalyzed  $\alpha$ -amination of esters (Scheme 42). $64$  The reaction of esters with N,N-di-tbutyldiaziridinone under the CuCl-n-Bu<sub>3</sub>P catalytic system provides the corresponding hydantoins. The proposed mechanism involves  $\alpha$ -cupration of esters by the transient Cu( $\pi$ )–N radical species or four-membered  $Cu(m)$  species derived from N,N-di-t-butyldiaziridinone and CuCl. The resulting  $\alpha$ -cupro(III)-esters 44 undergo C–N reductive elimination that is followed by cyclization to afford hydantoins.



Scheme 41 The reaction of di-t-butyldiaziridinone with Cu(I) complexes.



Scheme 42 Synthesis of hydantoins from esters and N,N-di-t-butyldiaziridinone via a-amination.

# 7. With fluoroamine derivatives

#### 7.1. With N-fluorobenzenesulfonimide (NFSI)

A highly reactive oxidant, N-fluorobenzenesulfonimide (NFSI) reacts readily with  $Cu(I)$  complexes to afford  $Cu(III)$ -imide species 45 that is in equilibrium with  $Cu(II)$ -stabilized sulfonimide radical 46 (Scheme 43). Thus, this nitrogen-centered radical derived from NFSI and  $Cu(1)$  complexes could initiate aminofunctionalization of alkenes.

For example, Zhang reported regioselective aminofluorination of styrenes with NFSI under CuBr/bathocuproine-catalyzed reaction conditions (Scheme  $44$ ).<sup>65</sup> The use of bis(pinacolato)diborone  $(B_2pin_2)$  and AgF as additives was crucial to facilitate the aminofluorination. The DFT calculations suggested that the C–F bond formation is likely enabled by F-radical abstraction from NFSI. This process concurrently generates the sulfonimide radical that can maintain the radical chain for aminofluorination.

In place of fluorine incorporation, cyano, amido, and azido moieties could be installed by Cu-catalyzed radical aminofunctionalization with NFSI. Xiong/Li/Zhang revealed that the reaction of styrene with NFSI and TMSCN under the CuBr-1, 10-phen catalytic system gives an aminocyanation product (Scheme 45-i).<sup>66</sup> Similarly, aminoazidation was developed by Studer using  $TMSN_3$  (Scheme 45-ii).<sup>67</sup> The C–CN and C–N<sub>3</sub> bond formation is mediated via radical recombination presumably through formation of organo-Cu $(m)$  intermediate 47 followed by ligand exchange with TMSCN or  $TMSN<sub>3</sub>$  to afford another organo-Cu( $\text{m}$ ) species 48 and subsequent C–CN or C–N<sub>3</sub> reductive elimination. The driving force of the ligand exchange could



Scheme 43 The reaction of NFSI with Cu(I) complexes for aminofunctionalization of alkenes.







Scheme 45 Cu(I)-catalyzed aminocyanation and diamination of styrene with NFSI

be preferential elimination of TMSF due to the strong affinity between Si and F atoms.

Interestingly, when the Cu-catalyzed reaction of styrene with NFSI was conducted in the presence of  $PhB(OH)_2$  in acetonitrile, a diamination product was formed through incorporation of acetonitrile by the Ritter-type reaction (Scheme 46). $67$ 

On the other hand, the Cu $(i)$ -catalyzed reactions of aliphatic alkenes with NFSI provide non-stabilized secondary radicals 49, which undergo intramolecular radical addition to the phenylsulfonyl moiety to afford sultams (Scheme 47).<sup>68</sup>

The combination of the Cu $(i)$ -catalyst and NFSI is also capable of functionalizing benzylic sp<sup>3</sup> C–H bonds (Scheme 48-i)<sup>69</sup> as well as  $sp^2$  C-H bonds (Scheme 48-ii)<sup>70</sup> on 5-membered-aromatic heterocycles such as thiophene and furan through the radical mechanism with the transient sulfonimide radical.



Scheme  $46$  Cu(OTf)<sub>2</sub>-catalyzed diamination of styrene with NFSI and acetonitrile.



Scheme 47 Synthesis of sultams by the Cu(I)-catalyzed reaction of aliphatic alkenes with NFSI.



Scheme 48 Cu(I)-catalyzed C-H amination with NFSI.

#### 7.2. With Selectfluor $\mathbf{B}$  and its derivatives (F-TEDA-X)

Selectfluor<sup>®</sup> [1-chloromethyl-4-fluoro-diazoniabicyclo[2.2.2]octane bis(tetrafluoroborate] and its derivatives (known as F-TEDA-X reagents, where X stands for their counter anions) have been used as versatile fluorination reagents in organic synthesis, $71$  while these reagents could be utilized as strong oxidants of  $Cu(II)$  and  $Cu(i)$  complexes to generate highly reactive  $Cu(m)$  species that can abstract the H-radical from  $sp<sup>3</sup>$  hybridized carbons. The combination of Cu catalysts and F-TEDA-X reagents is thus capable of oxidizing unactivated aliphatic C–H bonds. For example, Baran developed Cu-catalyzed Ritter-type aliphatic C–H amination with acetonitrile in the presence of F-TEDA-PF<sub>6</sub> (Scheme 49).<sup>72,73</sup> The C–H amination is likely enabled by a stepwise sequence involving (1) H-radical abstraction; (2) SET oxidation of the resulting C-radical 50 to carbocation 51; (3) Ritter-type amination by solvent acetonitrile. Nitrilium ion 52 is finally hydrolyzed to give acetamide products.

Interestingly, substrates having hydroxyl or carbonyl groups rendered the C–H amination process more chemo-selective and efficient presumably by their chelation effect. For example, the



Scheme 49 CuBr<sub>2</sub>-catalyzed C-H amination of cyclooctane with  $F-TEDA-PF<sub>6</sub>$ .



Scheme 50 CuBr<sub>2</sub>-catalyzed C–H amination of menthol with F-TEDA-PF<sub>6</sub>.

CuBr<sub>2</sub>-catalyzed reaction of  $(-)$ -menthol with F-TEDA-PF<sub>6</sub> afforded the chemo-selective C–H amination product dihydrooxazine in very high yield through intramolecular trap of the transient nitrilium ion 53 by the hydroxyl group (Scheme 50). Dihydrooxazine moiety 54 could be easily hydrolyzed into the corresponding 1,3-aminoalcohol.

A similar directing effect in chemo-selective aliphatic C–H oxygenation was observed in the reactions of N-(2-alkylphenyl) benzamides in the presence of  $Cu(OTf)_2$  as the catalyst and F-TEDA-BF<sub>4</sub> (Selectfluor<sup>®</sup>) for synthesis of  $4H-3,1$ -benzoxazines through *ortho-aliphatic* C–H oxygenation (Scheme 51).<sup>74</sup> The reactions selectively functionalize the ortho-alkyl group presumably via H-radical abstraction by the amide–Cu chelate intermediate,



Scheme 51 Cu(OTf)<sub>2</sub>-catalyzed C-H oxygenation with F-TEDA-BF<sub>4</sub>.

whereby aliphatic C–H bonds in the other positions (e.g. metamethyl group) are kept intact.

# 8. With 2,2,6,6-tetramethyl-1 piperidinyloxy (TEMPO)

A persistent radical, TEMPO (2,2,6,6-tetramethyl-1-piperidinyloxy), has been utilized for Cu-catalyzed oxidative C–O bond forming reactions. The unique feature of TEMPO is that it works as an oxidant of  $Cu(I)$  species to generate  $Cu(I)$  species and its reduced form TEMPO-H could be reoxidized by molecular  $O_2$  to regenerate TEMPO (Scheme 52).75

Chemler developed Cu-catalyzed enantioselective intra/ intermolecular aminooxygenation of alkenyl N-sulfonamides with TEMPO for synthesis of indoline and pyrrolidine derivatives (Scheme 53), in which TEMPO plays two roles as the external oxygen source and stoichiometric oxidant to realize catalytic turnover.<sup>76</sup> The detailed mechanistic analyses in experimental and theoretical manners revealed that the process is initiated by concerted syn-aminocupration of alkenes by  $Cu(n)$ -amido species 55 to construct heterocyclic frameworks having organocopper $(\text{II})$  moiety 56.<sup>77</sup> Subsequent  $C-Cu(n)$  bond homolysis results in formation of C-radical 57 that is trapped by TEMPO to afford the aminooxygenation product. The resulting lower valent  $Cu(I)$  species is reoxidized to the  $Cu(II)$  complex by TEMPO. Open Sec Rev Warterby sliphatic C-H bonds in the order positions (sg *metre* (continued on the complete on the complete on 22, 5.6 **- tetramethyl-1-**<br> **B C-MU C-2, A S-ACCE (FRAPCO)**<br> **B Properties**<br> **EVERY COMPOUTE T** 



Scheme 52 Oxidation of Cu(I) species to Cu(II) species by TEMPO



Scheme 53 Cu-Catalyzed aminooxygenation of alkenes with TEMPO.





In some cases especially when N-pentenylsulfonamides were employed, the use of molecular oxygen as an atmosphere could make the aminooxygenation process more efficient (Scheme 54). The TEMPO loading could be reduced to 1.5 equiv. as molecular oxygen serves as an oxidant to reoxidize TEMPO-H to TEMPO.

# 9. With metallic oxidants

Mild and cost-economical metallic oxidants have been employed as the terminal stoichiometric oxidants to regenerate higher valent active Cu species to realize catalytic turnover for the Cu-mediated oxidative molecular transformation.

#### 9.1.  $MnO<sub>2</sub>$

Hirano/Miura reported Cu(OAc)<sub>2</sub>-catalyzed intramolecular aromatic C–H amination of biaryl-2-picolinamide for synthesis of carbazoles with  $MnO<sub>2</sub>$  as the stoichiometric terminal oxidant to realize the catalytic turnover (Scheme  $55$ ).<sup>78</sup> The reaction is initiated by aromatic C–H cupration by copper $\left[\text{II}\right]$ – picolinamide chelate complex 58 to afford organo-Cu $(n)$  intermediate 59. Further redox disproportionation with  $Cu(OAc)_{2}$ forms copper $(m)$  intermediate 60 and subsequent C-N bond



Scheme 55  $Cu$ -Catalyzed-MnO<sub>2</sub>-mediated aromatic C–H amination for synthesis of carbazoles.



Scheme 56 Cu-Catalyzed-MnO<sub>2</sub>-mediated aromatic C–H amination for synthesis of indolines.

reductive elimination establishes formation of the carbazole product along with generation of a lower valent  $Cu(I)$  species that is oxidized by  $MnO<sub>2</sub>$  to regenerate the Cu( $\pi$ ) catalyst.

The analogous Cu-catalyzed-MnO<sub>2</sub>-mediated C-H amination strategy with picolinamides was applied for synthesis of indolines (Scheme 56).<sup>79</sup>

Using  $MnO<sub>2</sub>$  as the terminal oxidant, Chemler developed a series of  $Cu(<sub>II</sub>)$ -catalyzed amino- and oxy-functionalization of alkenes with alkenylsulfonamides, -anilines, and -alcohols for synthesis of the corresponding heterocycles. As shown in Scheme 57, the process is composed of a multi-step sequence involving (i) amino- or oxy-cupration of alkenes to form fivemembered ring organocopper $(n)$  intermediates 61; (ii) C–Cu $(n)$ bond homolysis to generate C-radical  $62$  and  $Cu(1)$  species; (iii) radical recombination with various internal/external carbon or heteroatom sources to provide difunctionalized final products; (iv) regeneration of the higher valent  $Cu(II)$  species by oxidation of the lower valent Cu( $i$ ) species by MnO<sub>2</sub>.

For example, treatment of N-aryl-2-allylaniline with 20 mol% of Cu(OTf)<sub>2</sub> in the presence of Cs<sub>2</sub>CO<sub>3</sub> and MnO<sub>2</sub> afforded indoline derivative through intramolecular carboamination of alkenes (Scheme 58).<sup>80</sup> In this process, the resulting C-radical 63 was added directly to the intramolecular benzene ring to construct the new C–C bond (Scheme 58-i). Enantioselective carboamination of  $N$ -pentenyl $(p$ -tolyl)sulfonamides was enabled by using the  $(R,R)$ -Ph-Box ligand for the Cu(OTf)<sub>2</sub> catalyst, delivering optically active bicyclic sultams (Scheme 58-ii).



Scheme 57  $Cu$ -Catalyzed-MnO<sub>2</sub>-mediated oxy- and amino-functionalization of alkenes.



Scheme 58 Cu-Catalyzed-MnO<sub>2</sub>-mediated carboamination of alkenes.

On the other hand, the reactions of N-mesyl-4-pentenylamines having the geminal diaryl moiety at the C2 position under the  $Cu(OTf)<sub>2</sub>(R,R)-Ph-Box$  catalytic system in the presence of MnO<sub>2</sub> provided 6-azabicyclo[3.2.1]octane in high enantioselectivity through carboamination of alkenes (Scheme 58-iii).<sup>81</sup>

The transient C-radicals generated via aminocupration of N-sulfonyl alkenylamines could undergo an iodine transfer reaction with isopropyl iodide to form the corresponding 2-iodomethyl indolines and pyrrolidines (Scheme  $59$ ).<sup>82</sup> Similarly, chlorination and bromination reactions were achieved in moderate yields using 1,1-dichloroethylene and (2,2-dibromo-1-methylcyclopropyl)benzene.

Cu-Catalyzed diamination of N-sulfonyl-alkenylamines was developed using tosylamide  $(TsNH<sub>2</sub>)$  as the external nitrogen source (Scheme  $60$ ).<sup>83</sup> In this case, the second C-N bond formation was enabled by radical recombination of C-radical 64 with TsNH<sub>2</sub> and Cu( $\pi$ ) species.

The Cu $(OTf)_{2}$ -catalyzed-MnO<sub>2</sub>-mediated reaction conditions were amenable to carboetherification of alkenyl alcohols, in which the second C–C bond formation was possible both in intra- and intermolecular fashions (Scheme  $61$ ).<sup>84</sup> Construction of 6-oxabicyclo[3.2.1]octanes was carried out using 4-pentenylalcohol with a geminal diaryl moiety at the C2 position via oxycupration of alkenes followed by intramolecular radical



**Scheme 59** Cu-Catalyzed-MnO<sub>2</sub>-mediated-aminohalogenation of alkenes.





Scheme 61 Cu-Catalyzed-MnO<sub>2</sub>-mediated carboetherification of alkenes.



Scheme 62 Cu-Catalyzed-MnO<sub>2</sub>-mediated intermolecular amination of 1,1-disubstituted alkenes.

cyclization on the aryl group (Scheme 61-i). Intermolecular C–C bond formation was also realized with aryl alkenes, in which the transient C-radical underwent the oxidative Heck-type coupling with aryl alkenes to deliver 2-allyltetrahydrofurans (Scheme 61-ii).

Chemler also reported Cu-catalyzed intermolecular amination of 1,1-disubstited alkenes with N-arylsulfonamides in the presence of MnO<sub>2</sub> as a terminal oxidant (Scheme 62).<sup>85</sup> The reactions uniquely afforded N-aryl enamide products in an anti-Markovnikov fashion.

#### 9.2.  $Ag_2CO_3$

 $Ag_2CO_3$  has been specifically employed as the stoichiometric oxidant for Cu-catalyzed aliphatic and aromatic C–H oxidation. Kuninobu/Kanai reported synthesis of b-lactams by Cu-catalyzed intramolecular sp<sup>3</sup> C–H amidation of  $N$ -(8-quinolinyl)amides in the presence of  $Ag_2CO_3$  (Scheme 63).<sup>86,87</sup> Installation of the N-8-quinolinyl moiety is crucial on the amide substituent to allow it to serve as the bidentate directing group. The C–H functionalization is mediated by the transient amide– $Cu(m)$ complex 65 through concerted metalation–deprotonation on sp<sup>3</sup> -C–H bonds with the acetate or carbonate counter ions (the reaction with the carbonate counter ion is shown below) on the copper to provide metallacycle intermediate 66. Finally, C-N bond forming reductive elimination affords  $\beta$ -lactam along with a lower valent  $Cu<sup>I</sup>$  species that reacts further with amide and  $Ag_2CO_3$  to regenerate the amide–Cu( $\text{m}$ ) complex to maintain the catalytic turnover.

The  $Cu(OAc)<sub>2</sub>-Ag<sub>2</sub>CO<sub>3</sub>$  reaction system was also utilized for aromatic C–H functionalization of benzamides having a 2-pyridylmethyl moiety on the amide nitrogen through directed



Scheme 63 Cu-Catalyzed-Ag<sub>2</sub>CO<sub>3</sub>-mediated directed sp<sup>3</sup> C–H amidation.



Scheme 64 Cu-Catalyzed-Ag<sub>2</sub>CO<sub>3</sub>-mediated directed sp<sup>2</sup> C–H sulfonylation.



Scheme 65 Cu-Catalyzed-Ag<sub>2</sub>CO<sub>3</sub>-mediated sp<sup>2</sup> C–H amidation.

## 10. Information of the oxidants: their commercial availability and preparation methods

Among the terminal oxidants for the Cu-catalyzed oxidative carbon–heteroatom bond formation discussed in this review, the price of the commercially available ones from Sigma-Aldrich is summarized in Table 1. Togni reagent  $II^{90}$  (entry 4) and peroxides (entries 5–7) are potentially explosive, so the reactions with these reagents should need special care with proper protecting shields.

The typical preparation methods of non-commercialized oxidants such as vinyl(aryl)iodonium triflate (Section 2.2.), O-benzoyl-N,N-dialkylhydroxylamines (Section 4), oxaziridines (Section 5), and diaziridinones (Section 6) are illustrated in Schemes 66–69, respectively.

Vinyl(aryl)iodonium triflate is readily prepared from the sequential treatment of the corresponding alkenylboronic acids with  $BF_3 \cdot OEt_2$ , 2-iodotoluene diacetate, and TfOH (Scheme 66).<sup>91</sup>

Table 1 A price list of commercially available oxidants

				<b>View Article Online</b>
<b>Review Article</b>				<b>Chem Soc Rev</b>
Cu(OAc) <sub>2</sub> (10 mol%) O Me Me O Me Me PhSO <sub>2</sub> Na (2 equiv)		Table 1 A price list of commercially available oxidants		
$Ag2CO3$ (2 equiv)	Oxidant Entry		Price $(USS)^{a}$	CAS no.
Н $SO2$ Pr	$Phi(OAc)$ <sub>2</sub> $\mathbf{1}$		$21.6/5$ g	3240-34-4
CICH <sub>2</sub> CH <sub>2</sub> CI, 120 °C	$\overline{2}$ Ph <sub>2</sub> IOTf		$114/1$ g	66003-76-7
72%	3	Togni reagent I	66.4/250 mg	887144-97-0
റ Me Me	4	Togni reagent II	50.6/250 mg	887144-94-7
	5 $t$ -BuOO $t$ -Bu		48.1/250 mL	110-05-4
	6	$t$ -BuOOH (5–6 M in decane)	151/100 mL	75-91-2
	7 (PhCO <sub>2</sub> ) <sub>2</sub>		$43.5/50$ g	94-36-0
67 SO <sub>2</sub> Ph	8 Selectfluor <sup>®</sup>	$(PhSO2)2NF (NFSI)$	$77.2/5$ g	133745-75-2
Scheme 64 Cu-Catalyzed-Ag <sub>2</sub> CO <sub>3</sub> -mediated directed sp <sup>2</sup> C-H sulfonylation.	9 <b>TEMPO</b> 10		$34.1/5$ g	140681-55-6
	11	MnO <sub>2</sub> (>99%)	$44.9/5$ g 49.4/100 g	2564-83-2 1313-13-9
	12	$Ag_2CO_3$ (>99%)	$34.9/5$ g	534-16-7
Cu(OAc) <sub>2</sub> (10 mol%)				
$Ag2CO3$ (3.0 equiv)		$a$ http://www.sigmaaldrich.com/united-states.html.		
benzene, 120 °C, 24 h		Preparation of O-benzoyl-N,N-dialkylhydroxylamines is con-		
		ducted either by benzoylation of N,N-dialkylhydroxylamines with		
93%				
		reactions of dibenzoyl peroxide with the corresponding secondary		
$\Omega$		amines (Scheme 67-ii). <sup>92</sup> O-Benzoyl-N,N-dialkylhydroxylamines		
Ċы OAc		should be stored in the freezer.		
68				
		1) BF <sub>3</sub> •OEt <sub>2</sub> (1.2 equiv), 0 °C		
		2) 2-MeC <sub>6</sub> H <sub>4</sub> I(OAc) <sub>2</sub> (1.2 equiv), 0 °C		
	$B(OH)_2$	3) TfOH (1.2 equiv), 0 °C	Me	-OTf
	Ph	CH <sub>2</sub> Cl <sub>2</sub>	Ph	
				55%
				after recrystallization
Intermolecular $sp^2$ C-H amidation/amination of quinoline		Scheme 66 Preparation of N-noxyl-3,3-dimethyloxaziridine.		
<b>Scheme 65</b> Cu-Catalyzed-Ag <sub>2</sub> CO <sub>3</sub> -mediated sp <sup>2</sup> C-H amidation. concerted metalation-deprotonation. Shi demonstrated Cu-catalyzed ortho-aromatic C-H sulfonylation of benzamides using sodium sulfinate as the sulfonylation reagent (Scheme 64). <sup>88</sup> The C-S bond forming reductive elimination from the transient $Cu(m)$ metallacycle 67 furnishes the sulfonylation product. N-oxides was also reported under the $Cu(OAc)2-Ag2CO3$ reaction system (Scheme 65). <sup>89</sup> Various lactams/cyclic amines are incor-				benzoyl chloride (Scheme 67-i) or by nucleophilic substitution
porated into the key organocopper $(m)$ intermediate 68 prior to its	(i)			



Scheme 66 Preparation of N-noxyl-3,3-dimethyloxaziridine



Scheme 67 Preparation of O-benzoyl-N,N-dialkylhydroxylamines.



Scheme 68 Preparation of N-noxyl-3,3-dimethyloxaziridine



Oxaziridines are synthesized by oxidation of the corresponding imines with  $Oxone^{\circledR}$ . The scalable procedure for synthesis of N-nosyl-3,3-dimethyloxaziridine reported by Yoon is shown in Scheme  $68.^{54,93}$ 

The synthetic procedure of N,N-di-t-butyldiaziridinone includes a sequence of preparation of di-t-butylurea and oxidative intramolecular N–N bond formation (Scheme 69). $94$  N,N-di-tbutyldiaziridinone should be stored in the dark.

# 11. Conclusions

This review highlighted up-to-date developments in coppercatalyzed (anaerobic) oxidative formation of carbon–heteroatom bonds on C–H bonds and alkenes. Various combinations of copper catalysts and readily available stoichiometric oxidants have been devised to enable unique and unprecedented oxidative molecular transformations. Unlike other transition metals, the reaction modes enabled by the copper species are multifarious. For example, many of the carbon–heteroatom bond-forming processes in Cu-catalyzed oxidative molecular transformations include organocopper (C–Cu) species as the key intermediates. The chemical reactivity of the organocopper species is uniquely diverse depending on the oxidation state of the copper moiety. Higher valent  $C-Cu$ <sup>III</sup> species undergo substitution reactions with heteroatom nucleophiles or reductive elimination of the C–heteroatom bond, whereas lower valent  $C-Cu<sup>T</sup>$  species exhibit a nucleophilic character to react with heteroatom electrophiles such as O-benzoyl-N,N-dialkylhydroxylamines. On the other hand,  $C-Cu^{\text{II}}$  species undergo homolysis to generate C-radicals that could be further functionalized with radical trapping reagents such as TEMPO or heteroatom nucleophiles through radical recombination pathways with the aid of Cu<sup>II</sup> species. More challenges and opportunities still remain for elucidation of the detailed reaction mechanisms and identification of active Cu-species that control the course of the reaction and improve catalytic turnovers and overall process efficiency. We anticipate that copper complexes have inexhaustible potential as catalysts to enhance our synthetic capability further.

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