A mechanistic investigation of the photoinduced, copper-mediated cross-coupling of an aryl thiol with an aryl halide†

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Photoinduced, copper-catalyzed cross-coupling can offer a complementary approach to thermal (non-photoinduced) methods for generating C–X (X = C, N, O, S, etc.) bonds. In this report, we describe the first detailed mechanistic investigation of one of the processes that we have developed, specifically, the (stoichiometric) coupling of a copper–thiolate with an aryl iodide. In particular, we focus on the chemistry of a discrete [CuI(SAr)2] complex (Ar = 2,6-dimethylphenyl), applying a range of techniques, including ESI-MS, cyclic voltammetry, transient luminescence spectroscopy, optical spectroscopy, DFT calculations, Stern–Volmer analysis, EPR spectroscopy, actinometry, and reactivity studies. The available data are consistent with the viability of a pathway in which photoexcited [CuI(SAr)2] acts as an electron donor to an aryl iodide to afford an aryl radical, which then reacts in cage with the newly generated copper(II)–thiolate to furnish the cross-coupling product in a non-chain process.

Introduction

The utility of cross-coupling chemistry has continued to expand at a rapid rate as novel or underexplored reaction pathways are exploited to achieve important new families of bond constructions. We have recently reported that, in the presence of light and a simple copper catalyst, coupling reactions of a variety of nucleophiles (nitrogen, sulfur, oxygen, and carbon) with aryl or alkyl electrophiles can be accomplished under mild conditions (–40 to 30 °C; eqn (1)).

We have suggested the outline of a possible pathway for these processes (Fig. 1, illustrated for C–S coupling), recognizing that the course of the cross-coupling is likely to vary with different coupling partners and reaction conditions. We have been interested in mechanistic similarities and dichotomies with photoredox catalysis, a mode of reactivity that has been the focus of great interest in recent years. For example, we have hypothesized that, distinct from a classical photoredox catalyst wherein a particular metal complex serves exclusively as an electron donor/acceptor, in our processes the copper complex may play a role both in electron transfer and in the key bond-forming step (e.g., C–S bond construction in Fig. 1). Furthermore, the mechanism depicted in Fig. 1 is not a radical-chain process; although non-chain pathways have frequently been invoked in earlier studies of photoredox catalysis, Yoon has recently concluded that, for three representative and mechanistically distinct transformations, the photoredox
catalyst serves to initiate a chain reaction. In this report, we describe our first study focused primarily on the mechanism of a photoinduced, copper-mediated cross-coupling, specifically, an investigation of the stoichiometric coupling of an aryl iodide with a copper–thiolate (eqn (2)).

\[
\text{Ar}^\text{I} - \text{I} \quad \text{[Cu\text{((SAr)}_2])^+} \\
\text{1.0 equiv} \\
\text{hv (100-watt Hg lamp)} \\
\text{CH}_3\text{CN, 0 °C} \\
\rightarrow \text{Ar}^\text{I} - \text{SAr} \quad \text{(2)}
\]

**Results and discussion**

**Background**

The photoinduced coupling of aryl thiols with aryl halides in liquid ammonia, in the absence of a catalyst, through an SRN1 mechanism is well-established through the work of Bunnett.\(^{12-14}\) In our initial report, we observed that a model photoinduced cross-coupling proceeds significantly more rapidly in the presence of a copper catalyst than in its absence (eqn (3)).

\[
\text{Ar}^\text{I} - \text{SAr} \\
\text{10% CuI} \\
\text{hv (100-watt Hg lamp)} \\
\text{1.0 equiv NaOt-Bu} \\
\text{CH}_3\text{CN, 0 °C, 5 h} \\
\rightarrow \text{85% (10% without CuI)} \quad \text{(3)}
\]

Under our reported conditions, the reaction mixture is heterogeneous, with a substantial portion of the NaSPh present as a solid. We have determined that, when the same partners are coupled at much lower concentration in a homogeneous solution, the rates of product formation can be similar in the presence and in the absence of CuI.\(^{15}\) Thus, a copper-mediated pathway and a copper-free pathway for C–S bond formation are possible, and which one is dominant can depend on the relative concentration in solution of sodium versus copper thiolates (the latter are generally more soluble in CH\(_3\)CN). In the present investigation, we seek to gain insight into the copper-mediated pathway.

In the mechanistic framework that we have previously described (Fig. 1), irradiation of a copper(I)–thiolate complex (A) leads to a photoexcited state (B). Electron transfer from B to the aryl halide furnishes a copper(II)–thiolate complex (C) and an aryl radical. Radical recombination then forms the C–S bond of the thioether, either directly\(^{16}\) or through a copper(II)–halide intermediate (D). Displacement of the halide of complex D by thiolate then regenerates copper(I)–thiolate complex A.

We have also considered a variety of other mechanisms, including the three illustrated in Fig. 2–4. The pathway depicted in Fig. 2 produces an aryl radical and a copper(II)–thiolate as a key step.

\[
\begin{align*}
\text{A} & \xrightarrow{\text{hv}} \text{B} \\
\text{B} & \xrightarrow{\text{Ar}^\text{I} - \text{SAr}} \text{C} \\
\text{C} & \xrightarrow{\text{Ar}^\text{I} - \text{X}} \text{D} \\
\end{align*}
\]

**Fig. 2** An alternative mechanism: coupling of an aryl radical with a copper(II)–thiolate as a key step.

Another mechanism under consideration largely follows the SRN1 pathway for copper-free C–S coupling reactions,\(^{12}\) the difference being that a photoexcited copper(II)–thiolate (B), rather than a photoexcited copper-free thiolate, serves as the initiating electron donor to the aryl halide, thereby generating a radical anion (F) that can participate in a chain reaction to form the thioether (Fig. 3).

\[
\begin{align*}
\text{A} & \xrightarrow{\text{hv}} \text{B} \\
\text{B} & \xrightarrow{\text{[L}_n\text{Cu\text{((SAr)}_2)]^+}} \text{C} \\
\end{align*}
\]

**Fig. 3** An alternative mechanism: SRN1 (radical chain process).

We have also considered mechanisms that do not involve an organic radical as an intermediate. For example, in the pathway depicted in Fig. 4, photoexcited complex B reacts with the aryl halide to cleave the C–X bond in a concerted process without the intermediacy of an aryl radical.\(^{19}\) Reductive elimination of the resulting copper(II) complex (G) leads to the thioether product.
(Ar=SAr) and copper(i)-halide adduct D. Ligand exchange then completes the catalytic cycle by regenerating copper(i)-thiolate A.

Previously reported mechanistic observations

In our original report, we described cyclization/stereochemistry data (eqn (4)) that are more readily accommodated by a radical/electron-transfer pathway (Fig. 1–3) than by a concerted pathway (Fig. 4) for C–X bond cleavage. Furthermore, in a relative-reactivity study (eqn (5)), we determined that the aryl halide that is more easily reduced via electron transfer (4-chlorobenzonitrile; −2.03 V vs. SCE in DMF21) is more reactive than the one with the weaker C–X bond (1-bromonaphthalene; −2.17 V vs. SCE in DMF21); this contrasts with thermal (non-photoinduced) copper-catalyzed S arylation, wherein essentially exclusive coupling of the aryl bromide is proposed to result from concerted oxidative addition.19,22

Our efforts to isolate a mononuclear [CuI(SPh)2]− complex (e.g., A in Fig. 1–4), which we had detected in an ESI-MS study of a C–S coupling reaction, led instead to a copper(i)-thiolate cluster, [CuI(SPh)2][Na(12-crown-4)2]. This cluster did, however, serve as a suitable stoichiometric coupling partner with an aryl iodide, as well as an effective (pre)catalyst for a photoinduced C–S cross-coupling.

Although these observations are consistent with our initial working hypothesis for the mechanism of photoinduced, copper-catalyzed C–S cross-couplings (Fig. 1), we concluded that a more detailed investigation was warranted.

Synthesis and characterization of a monomeric copper(i)-thiolate model complex

A copper(i)-thiolate (A) is the starting point in each of the pathways illustrated in Fig. 1–4. For ease of analysis in the present investigation, we sought a model system of simple speciation (monomeric). As demonstrated by Tshuva, the use of a hindered arylthiolate (2,6-dimethylthiophenolate; SAr, Ar = 2,6-dimethylphenyl) can avoid the formation of a cluster;23 furthermore, we had reported in our initial study that this arylthiolate serves as a suitable coupling partner in photoinduced C–S cross-couplings.24 Reaction of mesitylcopper(i), 2,6-dimethylthiophenol, and NaOrBu in CH3CN, followed by the addition of 12-crown-4, provided the desired sodium salt, [CuI(SAr)2][Na(12-crown-4)2] (1; “[CuI(SAr)2]Na”; eqn (6) and Fig. 5).

Our available data are consistent with the suggestion that this copper(i)-thiolate is a monomer in solution, as in the solid state. On the basis of diffusion-ordered NMR spectroscopy (DOSY), we estimate the hydrodynamic radii of the anion and the cation to be 4.2 and 4.4 Å, respectively, which are comparable to the corresponding computed radii of 4.5 and 4.8 Å. Furthermore, the molar conductivity for complex 1 in acetonitrile, 128.5 S cm2 mol−1, falls within the range (120–160 S cm2 mol−1) for other coordination compounds that are 1 : 1 electrolytes.24

NaSAr (Ar = 2,6-dimethylphenyl) is significantly more soluble in CH3CN than is NaSPh; consequently, for the photoinduced coupling of NaSAr with Ph–I under our standard conditions, the rates of reaction in the absence and in the presence of CuI are similar, in contrast to our observations with NaSPh (eqn (3)). Nevertheless, we have determined that [CuI(SAr)2]Na (1) couples at 0 °C with Ph–I in 56% yield, thereby substantiating the viability of photoinduced copper-mediated S arylation with this model complex (eqn (7)).

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In an ESI-MS study of the coupling of ArSH with Ph–I under our standard copper-catalyzed cross-coupling conditions (eqn (3)), we have detected an anion with a molecular weight of 337.2, which corresponds to that of [CuI(SAr)3]2–; under these conditions, we do not observe [CuI(SAr)2]2–, despite the large excess of thiolate relative to copper. Furthermore, 1H NMR and optical absorption spectra of complex 1 in the presence of excess thiolate, as well as DFT calculations, indicate that formation of [CuI(SAr)3]2– is unfavorable. Collectively, our data suggest that complex 1 exists as a two-coordinate monomer in solution, even in the presence of excess thiolate.

**Electrochemistry**

We have examined through electrochemistry the redox behavior of [CuI(SAr)2]Na (1) and of NaSAr (Fig. 6). The cyclic voltammogram of complex 1 shows an irreversible oxidative feature at $E_p = -0.18$ V vs. SCE that is also irreversible at $-20$ °C and at scan rates up to $1.5$ V s$^{-1}$ at $25$ °C. Following oxidation of 1, an irreversible feature is observed at $-1.85$ V vs. SCE, which corresponds to the reduction of bis(2,6-dimethylphenyl) disulfide (ArS–SAr), presumably formed from complex 1 upon electrochemical oxidation (oxidation of 1 with [FeCp2][PF6] also leads to the formation of ArS–SAr).

![Cyclic voltammograms of [CuI(SAr)2]Na (1) top and of NaSAr bottom. Conditions: scan rate: 100 mV s$^{-1}$; supporting electrolyte: 0.08 M (n-Bu)4N[B(C6F5)4]; working electrode: glassy carbon; reference electrode: Ag/AgNO3 (0.1 mM)/acetonitrile; auxiliary electrode: platinum wire; temperature: 25 °C.](Image 1)

To gain insight into whether free thiolate, generated either by simple ligand dissociation or through irradiation, might play a significant role in stoichiometric reactions of complex 1, we have monitored by cyclic voltammetry a solution of 1 (0.020 M) and [(n-Bu)4N][B(C6F5)4] as electrolyte in acetonitrile. The cyclic voltammogram is unchanged over 15 minutes of irradiation with a Hg lamp, suggesting that irradiation of complex 1 does not lead to the release of a detectable amount of free thiolate.

**Photophysical study of [CuI(SAr)2]Na (1)**

Complex 1 absorbs strongly in the ultraviolet region (top of Fig. 7), although only weakly at 365 nm ($\epsilon_{365} = 3$ M$^{-1}$ cm$^{-1}$), a prominent emission band for the 100 watt Hg lamp used in our photoinduced C–S couplings. The complex luminesces upon excitation at 355 nm with a lifetime of $\sim 7$ µs in acetonitrile, as determined by transient luminescence spectroscopy (bottom of Fig. 7). The lifetime of the emissive state does not change as a function of the observation wavelength, consistent with a single species being the source of luminescence. While the lack of a reversible Cu$^1$/Cu$^{$II$}$ redox couple precludes a true evaluation of the excited-state reduction potential for complex 1, we estimate this potential to be $-2.5$ to $-2.7$ V on the basis of the first ground-state oxidative feature ($E_p = -0.18$ V vs. SCE) (see Electrochemistry) and an approximate $E^0$ of 2.3–2.5 eV.$^{27,28}$ These data suggest that the excited state of complex 1 is sufficiently long-lived and reducing to engage in electron transfer with electrophiles such as aryl iodoles (Ph–I: $-1.91$ V vs. SCE in DMF$^{21}$).

![Top: Optical spectrum of [CuI(SAr)2]Na (1) ($\lambda_{max} = 258$ nm, $\epsilon = 2.3 \times 10^4$ cm$^{-1}$ M$^{-1}$); inset: excitation spectrum at 675 nm emission (dashed line) and emission spectrum at 353 nm excitation (solid line); in acetonitrile at 25 °C. Bottom: Time-resolved decay of the luminescence intensity of 1; in acetonitrile at 25 °C (25 μM); Nd:YAG laser at 355 nm excitation; observation wavelength: 675 nm.](Image 2)

To gain insight into the predicted electronic structure of the excited state of complex 1, we have performed time-dependent DFT calculations.$^{29}$ These calculations indicate that the lowest energy singlet state ($\lambda_{scale} = 325$ nm) consists of a transition from the HOMO (Cu–S antibonding) to the arene $\pi^*$ (Fig. 8). The population of a high-energy arene $\pi^*$ orbital in the excited state is consistent with 1 being a potent photoreductant.
Stern–Volmer kinetic analysis

The mechanisms outlined in Fig. 1–3 begin with electron transfer from a photoexcited copper(I)–thiolate [B] to the aryl halide. We have conducted a Stern–Volmer kinetic analysis of this elementary step, specifically, the reaction of the excited state of [CuI(SAr)2]Na (1) with Ph–I (reduction potentials: [CuI(SAr)2]−, ~−2.6 V; Ph–I: −1.91 V vs. SCE in DMF31), and we have determined that the rate constant for quenching is $8 \times 10^5$ M$^{-1}$ s$^{-1}$. As expected, an increase in the concentration of Ph–I leads to a decrease in the lifetime of the excited state (Fig. 9). The observed quenching results from electron transfer, not energy transfer; the emission spectrum of complex 1 exhibits no overlap with the absorption spectrum of Ph–I.30,31

\[ \Phi = \text{quantum yield} = 0.08 \]
\[ Q = \text{Stern–Volmer quenching fraction} \]
\[ = \frac{k_Q[\text{Ph–I}]}{\tau_0^{-1} + k_Q[\text{Ph–I}]} \]
\[ = \frac{8 \times 10^8 \text{M}^{-1} \text{s}^{-1}}{1.91 \text{V} \text{SCE in DMF}} = 0.10 \]
\[ \theta = \text{Volmer quenching fraction} = \frac{\Phi}{Q} = 0.8 \]

Consideration of a radical chain mechanism: quantum yield and chain length

We have established that the quantum yield (\(\Phi\)) for the stoichiometric coupling of [CuI(SAr)2]Na (1) with Ph–I when irradiated at 365 nm is 0.08(2),32 a value that can be accommodated either by a non-chain mechanism or by a chain mechanism with rapid chain termination. By dividing the quantum yield by the Stern–Volmer quenching fraction (\(Q\)), we have determined the chain length (the number of molecules of product formed per photoinduced electron-transfer event) for the C–S coupling of complex 1 with Ph–I to be 0.8 (eqn (8)).

This suggests that this cross-coupling proceeds via a non-chain pathway, as a chain mechanism would be expected to furnish more than one molecule of product from each photoinduced electron transfer. In contrast, Yoon concluded on the basis of a similar analysis that three representative reactions that involve photoredox catalysis proceed through a chain pathway.9

Viability of coupling an aryl radical with a copper–thiolate

The mechanistic observations described above are consistent with the suggestion that an aryl radical is generated under our cross-coupling conditions. This intermediate could subsequently form a C–S bond by reacting with species such as a Cu(II)–thiolate (Fig. 1) or a Cu(I)–thiolate (Fig. 2). We sought insight into the viability of such couplings by exploring reactions of an aryldiazonium salt, which can readily be converted into an aryl radical via one-electron reduction.34

When [CuI(SAr)2]Na (1; \(E_p = −0.18 \text{ V vs. SCE}\)) and 4-methoxyphenyldiazonium tetrafluoroborate (\(E^o = 0.14 \text{ V vs. SCE}\)) are allowed to react in CD$_3$CN at −20 °C for 30 minutes, no coupling is evident by $^1$H NMR spectroscopy (Table 1, entry 1). However, upon warming the mixture to room temperature for 30 minutes, C–S bond formation proceeds in 57% yield (entry 2). One possible pathway for this transformation begins with electron transfer from complex 1 to the aryldiazonium salt to afford a copper(II)–thiolate and Ar$^+$–N$_2$, which loses N$_2$ to generate an aryl radical that couples with the copper(II)–thiolate to form the C–S bond (eqn (9)).34

<table>
<thead>
<tr>
<th>Entry</th>
<th>Conditions</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>−20 °C, 30 min</td>
<td>&lt;2$^a$</td>
</tr>
<tr>
<td>2</td>
<td>r.t., 30 min</td>
<td>57</td>
</tr>
<tr>
<td>3</td>
<td>1.1 equiv. FeCp$_2$$^*$, −20 °C, 30 min</td>
<td>22</td>
</tr>
<tr>
<td>4</td>
<td>r.t., 30 min, 1.1 equiv. [FeCp$_2$$^*$]BF$_4$</td>
<td>56</td>
</tr>
</tbody>
</table>

$^a$ Run in CD$_3$CN; unreacted diazonium salt: >90%.
To assess the viability of the coupling of an aryl radical with a copper(i)–thiolate, we sought a reductant that would reduce the arylidiazonium salt and thereby generate an aryl radical under conditions in which copper(i)–thiolate 1 would not ([CD3CN, −20 °C; Table 1, entry 1). We determined that, in the presence of decamethylferrocene (FeCp2; E° = −0.12 V vs. SCE), the arylidiazonium salt is completely consumed within 30 minutes at −20 °C, furnishing a mixture of compounds that includes a 22% yield of the C–S coupling product (entry 3). The low yield of the diarylsulfide indicates that under these conditions an aryl radical reacts inefficiently, at best, with a copper(i)–thiolate to form a C–S bond; control experiments suggest the alternative possibility that at least some of the cross-coupling product may be formed from reaction of the aryl radical with a small amount of copper(II)–thiolate that is generated through a redox equilibrium between Cu(I)FeIII and Cu(II)FeII as the ferrocenium ion is formed.15 When the coupling illustrated in entry 2 is conducted in the presence of [FeCp2][BF4] (entry 4), the yield of the diarylsulfide is essentially unchanged (56%; entry 2 versus entry 4). This result indicates that the ferrocenium ion that is produced in entry 3 is not responsible for the diminished yield in that reaction.

**Rate of capture of an aryl radical by a copper–thiolate; in-cage versus out-of-cage coupling**

To obtain insight into the rate of capture of the aryl radical intermediate, we have determined the ratio of uncyclized/cyclized products for C–S couplings of several aryl iodides that have previously been employed in radical-clock studies (Table 2).16,19 Our data indicate that capture of the aryl radical by a copper–thiolate occurs competitively with a cyclization process that has a first-order rate constant of $\sim 4 \times 10^6$ s$^{-1}$ in benzene.

We have examined the relationship between the amount of [Cu(SAr)$_2$]Na (1) and the ratio of uncyclized/cyclized products, and we have determined that the product ratio remains essentially constant as we alter the quantity of complex 1 or the overall concentration (eqn (10)). These observations can be accommodated by the mechanism illustrated in Fig. 1, if C–S bond formation occurs between the aryl radical and copper(n)–thiolate C within the solvent cage (i.e., a single copper complex serves first as the electron donor and then as the source of SAr).

In contrast, for the mechanism illustrated in Fig. 2, the cyclized/uncyclized product ratio should depend on parameters such as stoichiometry and concentration, since C–S bond formation requires the aryl radical to leave the solvent cage and to encounter a copper(i)–thiolate (i.e., one copper complex serves as the electron donor and a different copper complex provides the SAR group).40,41

**Table 2 Reaction of an aryl radical: cyclization versus capture by a copper–thiolate**

<table>
<thead>
<tr>
<th>Cu(SAr)$_2$Na</th>
<th>1.0 equiv</th>
<th>hv (100-watt Hg lamp)</th>
<th>CH$_2$CN, 0 °C, 5 h</th>
<th>Products</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ar$^+$–I</td>
<td>Products$^a$</td>
<td>k (s$^{-1}$)$^b$</td>
<td></td>
<td></td>
</tr>
<tr>
<td><img src="image1" alt="Uncyclized" /></td>
<td>1 : &gt;20</td>
<td>9.6 x 10$^9$ (DMSO)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><img src="image2" alt="Cyclized" /></td>
<td>1 : 5</td>
<td>4.2 x 10$^8$ (benzene)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><img src="image3" alt="Uncyclized" /></td>
<td>20 : 1</td>
<td>6.0 x 10$^5$ (water)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

$^a$ Ratio determined through GC analysis with dodecane as an internal standard (average of two experiments). $^b$ Rate of cyclization of the aryl radical (20–25 °C).
copper(n)-thiolate complexes. Indeed, photolysis of a solution of [CuI(SAr)2]Na (1) in the presence of excess Ph−I and NaSAr in propionitrile : butyronitrile (1 : 1) at −78 °C results in a blue solution, the EPR spectrum of which is consistent with the presence of some amount of a copper(n)-thiolate radical (Fig. 10).

The four-line hyperfine coupling is consistent with an I = 3/2 paramagnetic copper complex with a single metal center. The spectrum shows modest g anisotropy compared to other copper(n)-thiolate complexes, which is consistent with a highly covalent Cu–S interaction. This suggests that significant radical character resides in sulfur p orbitals, and DFT calculations support this assessment (see below).

Optical spectroscopy can serve as an additional technique for characterizing the putative copper(n)-thiolate. Upon irradiating complex 1 in the presence of Ph−I and NaSAr in propionitrile at −78 °C, a feature at 582 nm is observed (Fig. 11), which is consistent with the blue color of the reaction mixture. This feature is near the range found for sulfur-rich copper(II) proteins (593 to 610 nm).

While the above data provide strong evidence for the generation of a copper(n)-thiolate radical upon photolysis of a mixture of complex 1, Ph−I, and NaSAr, they do not identify the specific paramagnetic copper species, and further characterization is complicated by its instability even at −78 °C. The presence not only of Ph−I, but also of NaSAr, is required for detection of this copper(n)-thiolate radical by optical and by EPR spectroscopy. In view of the need for exogenous thiolate, we hypothesize that a copper(n) tris(thiolate), [CuII(SAr)3], may be formed, e.g., via electron transfer from [CuI(SAr)2]− to the aryl halide to form CuII(SAr)2, followed by trapping by NaSAr (eqn (11)). DFT calculations suggest that binding of an arylthiolate to CuII(SAr)2 is exergonic by ~4 kcal mol−1.

Alternatively, the copper(n) tris(thiolate), [CuII(SAr)3]−, could be generated by electron transfer to Ph−I from the excited state of NaSAr, followed by coupling of the thyl radical with [CuI(SAr)2]− (eqn (12)). Our observations to date do not allow us to definitively distinguish between these two pathways for the formation of putative [CuII(SAr)3]−.

DFT calculations of [CuII(SAr)3]− and of CuII(SAr)2 predict that significant spin density would reside on the thiolate ligands for either compound, which suggests that C–S bond formation could occur through direct reaction of the aryl radical with the copper-bound thiolate ([CuII(SAr)2]− : Cu 0.23e−, 3S 0.57e−; CuII(SAr)2: Cu 0.14e−, 2S 0.63e−; Fig. 12).

Conclusions

In this report, we describe the first detailed mechanistic investigation of one of the photoinduced, copper-mediated cross-couplings that we have developed, specifically, the coupling of a thiol with an aryl iodide. Due to the existence of a parallel, copper-free C–S bond-forming pathway, we have focused our attention on understanding the stoichiometric chemistry of a key proposed intermediate, [CuII(SAr)3]− (Ar = 2,6-dimethylphenyl); our observations to date are consistent with the viability of the elementary steps outlined in Fig. 1 (A → D).

We have established that [CuI(SAr)2]Na (1) is a two-coordinate monomer both in the solid state and in solution, and we have detected [CuI(SAr)3]− through ESI-MS under cross-coupling conditions. Complex 1 undergoes excitation upon irradiation at 365 nm (a prominent emission band of a Hg lamp), and it luminesces with a lifetime of ~7 μs; we estimate...
its excited-state reduction potential to be $\sim$2.6 V. Through a Stern–Volmer study, we have determined that the excited state is effectively quenched by Ph–I, as expected on the basis of its reduction potential; correspondingly, complex 1 reacts with Ph–I upon irradiation to afford the C–S coupling product. By employing actinometry, we have established that the chain length for the coupling of complex 1 with Ph–I is 0.8, indicating a non-chain mechanism. Our EPR and optical spectroscopy data suggest that a copper(II)–thiolate is formed when complex 1 is irradiated in the presence of Ph–I and NaSAr. Furthermore, through the use of an aryldiazonium salt, we have independently generated an aryl radical in the presence of copper(I)– and copper(III)–thiolates, and we have provided evidence that C–S bond formation is more efficient in the case of a copper(II)–thiolate. Finally, with the aid of radical clocks, we have established that C–S bond formation likely occurs via an in-cage mechanism in which a single copper complex serves both as an electron donor ($\text{Cu}^3 \rightarrow \text{Cu}^1$) and a source of SAr (copper(II)–thiolate). Thus, the available data support the viability of the mechanism illustrated in Fig. 1 (A $\rightarrow$ D), a mechanism that is distinct from most applications of photoredox catalysts in organic synthesis; other C–S coupling pathways, for example involving initial photooxidation of the aryl halide by a photoexcited copper-free thiolate, may also be operative. Our current efforts are directed at evaluating the degree to which the mechanism illustrated in Fig. 1, or alternative mechanisms, applies to other photoinduced, copper-catalyzed cross-couplings.

Acknowledgements

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Notes and references


5 For the sake of simplicity in Fig. 1–4, the copper complexes are shown without charges, and all processes are depicted as being irreversible. Halides (X) may be bound as inner- sphere or as outer-sphere ligands.


11 Although not explicitly shown, capture of [L$_n$Cu$_m$(SAr)]X by Ar$^+$ could generate a Cu(m) intermediate, [L$_n$Cu$^{(m)}$(Ar$^+$)(SAr)]X, which could then reductively eliminate to afford Ar$^+$SAr and complex D.


For additional information, see the ESI† For other classes of nucleophiles that we have examined, the rate in the presence of copper is significantly greater than in the absence of copper. For example, see ref. 2 and 3.

For early examples of copper-mediated C–S coupling processes that have been suggested to proceed without the formation of a Cu(n) intermediate, see: (a) C. L. Jenkins and J. K. Kochi, J. Org. Chem., 1971, 36, 3095; (b) C. L. Jenkins and J. K. Kochi, J. Am. Chem. Soc., 1972, 94, 856.

Copper[0] “complex” E could be copper metal or a formal copper[0] complex that bears a redox-active ligand.


A concerted pathway for C–X bond cleavage is generally invoked in thermal (non-photoinduced) copper-catalyzed cross-couplings. For example, see: (a) C. Chen, Z. Weng and J. F. Hartwig, Organometallics, 2012, 31, 8031; (b) M. Font, T. Parella, M. Costas and X. Ribas, Organometallics, 2012, 31, 7976.

A step-wise ET process whereby \([\text{L}_2\text{Cu}^+\text{(SAr})]X\) is generated and then captured by \(\text{Ar}^+\) to generate G is also plausible. See also ref. 11.


For a mechanism wherein C–X bond cleavage occurs through halogen-atom transfer, 1-bromonaphthalene would generally be expected to be more reactive than 4-chlorobenzonitrile.


These workers have crystallographically characterized \([\text{Cu}^+\text{(SAr})]_2[\text{NET}_4]\).


A DFT study indicates that binding of thiolate to \([\text{Cu}^+\text{(SAr})]\) to produce \([\text{Cu}^+\text{(SAr})]^2\) is unfavorable by \(\sim 20\) kcal mol\(^{-1}\) (see the ESI†).

Less than 5% decomposition of complex 1 was observed after irradiation (Hg lamp) in CH\(_2\)CN at 0 °C for five hours.

The lower limit of this range is based on the onset of emission, and the upper limit of this range is calculated from the midpoint of the maxima of the emission and excitation spectra (inset of Fig. 7).


Time-dependent DFT calculations were performed using the BP86 functional within the Tamm–Dancoff approximation employing the def2-TZVP basis set. The 50 lowest-lying singlet excited states were calculated, based on the DFT-optimized geometry of complex 1.


The triplet state of Ph–I is 3.5–3.6 eV (see ref. 30), whereas the triplet-state energy of 1* is 1.8 eV based on an emission maximum of 675 nm. These data exclude the possibility of Dexter energy transfer.


We have investigated the possibility that C–S bond formation is occurring through a nucleophilic aromatic substitution reaction \((\text{S}_\text{Ar})\) of the copper(--)thiolate with the aryl diazonium salt. For the cross-coupling of complex 1 with an aryl diazonium salt that bears a pendant olefin that is suitably positioned for intramolecular addition by an aryl radical, we observe none of the “direct” coupling product, indicating that substitution \(\text{via an } \text{S}_\text{Ar}\) pathway is not favored under these conditions (see the ESI†).


When the aryl diazonium salt is mixed with FeCp\(_3\) in CD\(_3\)CN at \(-20\) °C for 30 minutes, anisole is produced in \(\sim 85\%\) yield.

For details, see the ESI†.


We have conducted control experiments that establish that these aryl iodides are stable to irradiation under the indicated conditions.

When the cross-coupling illustrated in entry 2 of Table 1 is conducted at different concentrations \((0.010\ M\ \text{versus} \ 0.0050\ M)\), the yield of the coupling product does not change, which is also consistent with a single copper complex serving as the electron donor and the source of SAr.

When the cross-coupling illustrated in eqn (10) is conducted with NaSAr in place of \([\text{Cu}^+\text{(SAr})]_2\text{Na}\), a different ratio \((1:18)\) of uncyclized : cyclized products is observed, consistent with the suggestion that different thiolates (copper versus sodium) are coupling with the organic radical in the two processes.


As determined by spin quantification, the copper[II] species is present at a concentration of 0.2 mM, which corresponds to \(\sim 5\%\) of the starting copper(II) complex.


Intermolecular trapping of \([\text{Cu}^+\text{(SAr})]_2\) by free thiolate requires cage escape by the aryl radical generated \(\text{via}\) electron transfer to Ar–I. While our radical-clock studies are most consistent with in-cage recombination between a copper(II)–thiolate and an aryl radical at 0 °C, bimolecular reactions that follow Eyring behavior can, when there is no coulombic term and the viscosity change is not dramatic, be expected to exhibit enhanced cage escape as the temperature is lowered in fluid solution, according to the Eigen equation (see: J. Olmsted and T. J. Meyer, J. Phys. Chem., 1987, 91, 1649). In general, the
temperature dependence of the partitioning between in-cage versus out-of-cage radical pathways is challenging to predict (see the ESI† for additional information).

47 According to a DFT calculation, binding of iodide anion to CuII(SAr)2 is approximately thermoneutral (see the ESI†).

48 We have examined whether evidence for [CuII(SAr)3]− might be obtained, if electrochemical oxidation of [CuI(SAr)2]Na (1) is conducted in the presence of NaSAr. However, the cyclic voltammogram of complex 1 is unchanged in the presence of 10 equivalents of NaSAr, and its oxidation remains irreversible. This result is consistent with our expectations. As evident in Fig. 6, the onset for the oxidation feature of NaSAr is cathodically shifted relative to that for complex 1.