Photocatalytic Oxidative C–H Thiolation: Synthesis of Benzothiazoles and Sulfenylated Indoles

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Abstract
We report studies on the photocatalytic formation of C–S bonds to form benzothiazoles via an intramolecular cyclization and sulfenylated indoles via an intermolecular reaction. Cyclic voltammetry (CV) and density functional theory studies suggest that benzothiazole formation proceeds via a mechanism that involves an electrophilic sulfur radical, while the indole sulfenylation likely proceeds via a nucleophilic sulfur radical adding into a radical cationic indole. These conditions were successfully extended to several thiobenzamides and indole substrates.

Key words photocatalysis, benzothiazole, indole, thiolation, C–H functionalization

The formation of carbon–sulfur bonds is an important reaction in synthetic chemistry, as this motif is found in numerous natural products, pharmaceuticals, polymers, and semiconductors.1–8 The most common methods to achieve (C–S) bond formation have utilized transition-metal thiol cross-couplings;9–11 however, these methods typically involve harsh reaction conditions, high temperatures, and require pre-functionalization of the substrate. It would be more desirable to directly functionalize the C–H bond without any intermediate transformation. Direct C–H thiolation has been previously achieved through electrophilic aromatic substitution (SAr) utilizing activated sulfenyl sources such as sulfinyl halides or N-thiosuccinimides.12–16 These reactions are limited primarily to electron-rich aromatics and heterocycles such as substituted indoles. We have recently reported methodologies that function via a Lewis base/Brønsted acid dual catalytic system that allow for the C–H sulfenylation of diverse arenes.17,18 One drawback to this approach is that the formation of activated sulfenyl sources is often cumbersome; thus, methods that could activate readily available thiols in situ would represent a welcome advancement.

Over the past decade, radical chemistry, specifically photoredox catalysis and electrochemistry, has risen as a powerful and popular tool for C–H functionalization.19–22

Scheme 1
Previous photocatalytic and electrochemical methodologies for C–H thiolation of thioamides to benzothiazoles and various electron-rich heterocycles.
The radical species acts as a highly reactive intermediate, which enables synthetic transformations which normally cannot be assessed under reaction conditions involving polar pathways.\textsuperscript{23} In the past five years, there have been multiple accounts of C–H thiolation employing the use of photoredox catalysis (Scheme 1). In 2015, Lei showed that benzothiazoles can be synthesized using a Ru(bpy)\textsubscript{3}(PF\textsubscript{6})\textsubscript{2}/Co dual catalytic system.\textsuperscript{24} Similarly, Xu reported a similar benzothiazole transformation from thioamides using a TEMPO-catalyzed electrochemical C–H thiolation.\textsuperscript{25} Alternatively, Barman and Fan both independently reported the use of Rose bengal and thiophenol for the sulfenylation of 3-substituted indoles and imidazopyridines, respectively.\textsuperscript{26,27} Recently, König and Rehbine showed that electron-rich arenes (such as trimethoxybenzenes) could react with diaryl and dialkyl sulfides with an iridium photocatalyst and a persulfate salt to provide arylthiols.\textsuperscript{28} Herein, we report an oxidative photocatalytic thiolation to synthesize benzothiazoles through an intramolecular synthesis from thioamides, as well as the intermolecular sulfenylation of substituted indoles (Scheme 2).

Notably, mechanistic studies via cyclic voltammetry and density functional theory calculations suggest that even though both reactions use similar conditions, they proceed with markedly different roles for the sulfur, with an electrophilic sulfur radical in the benzothiazole formation, and a nucleophilic sulfur radical in the indole sulfenylation.

While studying whether the Lewis basic thioamide in 1a could act as a directing group for ortho-chlorination via S\textsubscript{2}Ar using Hu’s photocatalytic chlorination conditions\textsuperscript{29} we observed a significant amount of benzothiazole 1b (Table 1, entry 1). Interestingly, removal of the sodium chloride provided a small increase in conversion of 1a into 1b, suggesting this chemistry occurred via a substrate oxidative mechanism rather than sulfur activation through the halogen source (Table 1, entry 2). Removal of both the Ru(bpy)\textsubscript{3}Cl\textsubscript{2} (Table 1, entry 3) and sodium persulfate (Table 1, entry 4) resulted in a significant decrease in conversion; however, there is a still a small benzothiazole background reaction in the presence of persulfate. We then continued our optimization with an evaluation of other common photocatalysts. Because we utilized a 390 nm LED blue light source, we hypothesized that Ru(phen)\textsubscript{3}Cl\textsubscript{2} (1\textsubscript{max} = 422 nm) would be in a higher absorbance range relative to Ru(bpy)\textsubscript{3}Cl\textsubscript{2} (1\textsubscript{max} = 452 nm). However, conversion of the benzothiazole was low at only 15% (Table 1, entry 5). Switching from a transition metal to an organic photocatalyst 4CzIPN also provided no improvement in conversion (Table 1, entry 6), possibly due to the reaction being performed in a biphasic solvent system. Surprisingly, [Ir(dF(CF\textsubscript{3})ppy)\textsubscript{2}(dtbpy)]PF\textsubscript{6}, which has a higher oxidizing potential in its excited state ([Ir(III)/Ir(II) = 1.21 V vs SCE] relative to Ru(bpy)\textsubscript{3}Cl\textsubscript{2} [Ru(II)/Ru(I) = 0.77 V vs SCE]) and would be expected to oxidize 1a more effectively, proved markedly worse than the ruthenium catalyst (Table 1, entry 7). Mechanistically, this implies that while the photocatalyst has a significant effect on the overall conversion of the reaction, its excited state does not directly oxidize the thioamide but rather likely activates the persulfate as a better oxidizing agent (see proposed mechanism). Testing solvent conditions, we observed a decrease in conversion when switching to a more organic composition of MeCN/H\textsubscript{2}O (9:1), suggesting aqueous media is necessary to help solubilize the persulfate salt (Table 1, entry 8). To see if circumventing persulfate activation was a possibility, we added excess amount of sodium persulfate (Table 1, entries 9 and 10); however, we only obtained the acetanilide side product, which is a common degradation product for thioamides under oxidative conditions. Finally, we observed that benzothiazole conversion could be improved markedly (up to 79%) by the addition of two equivalents of pyridine as a base.

We decided to evaluate our conditions from Table 1, entry 11 across a variety of substituted thioamide derivatives (Scheme 3). To confirm our initial hypothesis, we tested the substrates in the absence and presence of pyridine and obtained isolated yields of the benzothiazoles. Varying the electronics at the aryl ring R\textsubscript{1} (2a–5a) provided minor decreases in yield relative to the unsubstituted 1a (isolating between 49–63% yield for 2b–5b). Notably, we observed no effect when adding pyridine for naphthyl-based substrate 6a (54% with no pyridine, 55% with pyridine for 6b) and substrate 7a (32% without, 34% with pyridine for 7b). This lack of pyridine effect held for other substrates that possessed these aryl groups 9a (43% no pyridine, 38% with pyridine for 9b), and 15a (31% no pyridine, 30% with pyridine for 15b). Replacing the thioamide tert-butyl group 1a with a phenyl group in 8a resulted in a marked decrease in yield (79% to 32% of 8b); however other phenyl-containing thioamides resulted in decent yields (i.e., 10a resulted in 68% yield 10b). Finally, when we replaced the thioamide substitution with aliphatic groups other than tert-butyl (11a–14a), we isolated the corresponding benzothiazoles in good
yield (55–73% 11b–14b). Surprisingly, when the thioamide substitution was a methyl (16a) we only isolated the corresponding amide. Alternatively, when we evaluated trifluoromethyl containing 17a, we observed no reaction of any kind, perhaps due to the thioamide being significantly more electron poor and possessing a higher redox potential, or a lower innate nucleophilicity.

Table 1  Optimization of Intramolecular Benzothiazole Synthesis of 2,2-Dimethyl-N-phenylpropanethioamide

<table>
<thead>
<tr>
<th>Entry</th>
<th>Catalyst</th>
<th>Oxidant (equiv)</th>
<th>Additive (equiv)</th>
<th>Solvent</th>
<th>Conversion (%) b</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Ru(bpy)_3Cl_2</td>
<td>Na_2S_2O_8 (2)</td>
<td>NaCl (3)</td>
<td>MeCN/H_2O (1:1)</td>
<td>52</td>
</tr>
<tr>
<td>2</td>
<td>Ru(bpy)_3Cl_2</td>
<td>Na_2S_2O_8 (2)</td>
<td>none</td>
<td>MeCN/H_2O (1:1)</td>
<td>57</td>
</tr>
<tr>
<td>3</td>
<td>Ru(bpy)_3Cl_2</td>
<td>none</td>
<td>none</td>
<td>MeCN/H_2O (1:1)</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>none</td>
<td>Na_2S_2O_8 (2)</td>
<td>none</td>
<td>MeCN/H_2O (1:1)</td>
<td>5</td>
</tr>
<tr>
<td>5</td>
<td>Ru(phen)_3Cl_2</td>
<td>Na_2S_2O_8 (2)</td>
<td>none</td>
<td>MeCN/H_2O (1:1)</td>
<td>15</td>
</tr>
<tr>
<td>6</td>
<td>4CzIPN</td>
<td>Na_2S_2O_8 (2)</td>
<td>none</td>
<td>MeCN/H_2O (1:1)</td>
<td>30</td>
</tr>
<tr>
<td>7</td>
<td>[Ir(df(CF_3)ppy)_2(dtbbpy)]PF_6</td>
<td>Na_2S_2O_8 (2)</td>
<td>none</td>
<td>MeCN/H_2O (1:1)</td>
<td>20</td>
</tr>
<tr>
<td>8</td>
<td>Ru(bpy)_3Cl_2</td>
<td>Na_2S_2O_8 (2)</td>
<td>none</td>
<td>MeCN/H_2O (9:1)</td>
<td>34</td>
</tr>
<tr>
<td>9</td>
<td>Ru(bpy)_3Cl_2</td>
<td>Na_2S_2O_8 (5)</td>
<td>none</td>
<td>MeCN/H_2O (1:1)</td>
<td>0 (1c obtained)</td>
</tr>
<tr>
<td>10</td>
<td>Ru(bpy)_3Cl_2</td>
<td>Na_2S_2O_8 (10)</td>
<td>none</td>
<td>MeCN/H_2O (1:1)</td>
<td>0 (1c obtained)</td>
</tr>
<tr>
<td>11</td>
<td>Ru(bpy)_3Cl_2</td>
<td>Na_2S_2O_8 (2)</td>
<td>pyridine (2)</td>
<td>MeCN/H_2O (1:1)</td>
<td>79</td>
</tr>
</tbody>
</table>

*Reactions were performed on a 0.130 mmol scale (approximately 25 mg) with 5 mol% photocatalyst loading in 1 mL of solvent mixture of MeCN/H_2O.

*Conversions were measured by NMR integrated spectra; the results are reported as an average of two trials. See Supporting Information for the details.
To explain the subsequent transformation, we propose the following mechanism (Scheme 4), which is supported by several key experiments. Due to the increase in yield upon addition of pyridine, we believe there is a Lewis base effect wherein the pyridine coordinates to the N–H thioamide bond, providing extra stability to the radical cation that forms upon initial oxidation. This hypothesis is substantiated through cyclic voltammetry experiments on substrate \( A \).

In pure acetonitrile without the presence of additive base, we observed two half-wave oxidation potentials at 1.5 V and 1.9 V vs SCE, meaning both values are out of the range of the Ru(bpy)\(_3\)Cl\(_2\) reduction potential in its excited state \([\text{Ru(II)/Ru(I)} = 0.77 \text{ V vs SCE}]\). Upon titration of pyridine, we noticed a distinct shift in the two oxidation potentials to 1.2 V and 1.5 V vs SCE. Interestingly, the first oxidation potential of \( A \) with pyridine is now within the range of the ground-state reduction potential of Ru(bpy)\(_3\)Cl\(_2\) \([\text{Ru(III)/Ru(II)} = 1.29 \text{ V vs SCE}]\). This suggests that, in its excited state, the photocatalyst reduces persulfate to the \( \text{SO}_4^{2-} \) anion and the \( \text{SO}_4^{•-} \) anion radical, followed by the resultant Ru\(^{2+}\) complex oxidizing the thioamide substrate to radical cation \( B \). Additionally, we utilized density functional theory (DFT) calculations to predict the electron-density maps for several thioamide intermediates and consequently predict the most favorable sites for oxidation. In the first map, we see a large concentration of electron density at the sulfur relative to the rest of the molecule \( A \), implying it is the most favorable site for initial oxidation; this pathway is also supported by recently reported work from Nicewicz on allylic thioamides.\(^{32,33}\)

At this point, \( B \) will likely undergo radical cyclization to \( C \). This is supported by CV scan-rate experiment; as we sweep from 0.1 V/s to 5 V/s, the second half-wave oxidation peak begins to diminish and completely disappears at the highest scan rate. One explanation for this observation is there is a new intermediate reaction between the first and second oxidation (i.e., thioamide cyclization) and that faster voltage sweeps can kinetically outpace the reaction, thereby hindering subsequent oxidation. Additionally, the predicted electron map of \( B \) suggests that the sulfur is now

### Scheme 4

Cyclic voltammetry (CV) measurements of substrate \( 1a \) with variation in potential scan rate (a) and with pyridine additive (b). Both processes show two oxidation peaks in an irreversible process. Increasing the scan rate shows disappearance of the second oxidation peak, implying a chemical reaction step between the first and second oxidation towards product formation. Titration of pyridine shows a lowering of both oxidation potentials. CV experiments were run vs Ag wire reference electrode, a glassy carbon working electrode, and a platinum counter electrode, followed by standard conversions to saturated calomel electrode (SCE). (c) A proposed mechanism for intramolecular benzothialation is shown, with electron-density maps derived from density functional theory (DFT) structure optimizations. The key experiments suggest an initial oxidation at the sulfur to form the thiol radical cation, which then undergoes intramolecular cyclization. Coordination of the pyridine additive to the substrate lowers the first half-wave oxidation due to coordination with the N–H thioamide bond, providing a more favorable single-electron-transfer process.
more electron deficient compared to the aryl ring; thus, it is likely that the cyclization will occur via the arene acting as a nucleophile and the sulfur acting as an electrophilic radical. Due to the comparable yields of electron-deficient thioamides without the presence of pyridine (i.e., 10a), we believe that the additive is beneficial towards initial oxidation but not necessary as persulfate can also promote formation of the thyl radical cation; it is the subsequent radical cyclization which drives the reaction favorably towards the benzothiazole product. Upon cyclization, the electron-density map shows a more electron-rich intermediate which should be easily oxidized by the persulfate radical anion to give Wheland arenium ion D, which will rapidly undergo aromatization to the final product E.

We also explored whether our methodology for intramolecular C–H thiolation could be applied to other arenes for intermolecular functionalization, specifically the sulfenylation of electron-rich heterocycles such as indoles. Our initial experiment utilized our optimized conditions for benzothiazole synthesis without pyridine, using 18a melatonin as the substrate and 4-methyl thiophenol as the sulfonylating reagent, and obtained 29% yield of 18b (Table 2, entry 1). Upon addition of pyridine (Table 2, entries 2 and 3), we observed a similar trend as the yields increased to 40%. Just like the previous reaction, removal of the photocatalyst diminishes the yield significantly to 8% (Table 2, entry 4), however, there is still a background reaction from just persulfate exclusively. Interestingly, reintroduction of the photocatalyst but cutting the persulfate equivalent in half reduced the overall yield to 5% (Table 2, entry 5). As expected, complete removal of persulfate provides no reaction (Table 2, entry 6).

Similar trends also hold for changing the ratio of solvents, as we see almost no variation going from 1:1 to 9:1 MeCN/H2O, and a lowering of 15% yield switching to completely acetonitrile (Table 2, entries 7 and 8). After evalu-
ing a number of photocatalysts (Table 2, entries 9–11), we observed that \( \text{[Ir(dF(CF}_3)ppy]_2(dtbpy)} \text{PF}_6 \) gave a much higher yield at 52% compared with \( \text{Ru(bpy)}_3\text{Cl}_2 \). Finally, variation of the base to potassium hydroxide (Table 2, entries 12–14) provided an increase of yield to 68%. Interestingly, we noticed a trace amount of a disulfenylated side product 18c when the iridium photocatalyst is used, suggesting that at some point the thiophenol reagent (or product sulfide) is oxidized and reacts with a second equivalent of thiophenol.

With our optimized conditions, we evaluated a number of substituted indoles and report the isolated yields of both the mono- and disulfenylated product, with a majority of substrates providing exclusively the monosulfenylated product in 9–36% yield (Scheme 5, 19–24). Similar to melatonin, N-methyl 3-methylindole (20a) also gave a mixture of monosulfenylated 20b and disulfenylated 20c (28% and 14%, respectively) Additionally, we tested a number of benzenethiol reagents (25a–27a) and varied the electronics off the aryl ring; this gave attenuated yields ranging from 16–31% yield (25b–27b). While these yields are moderate compared to other conditions (both via traditional \( SEAr \), and photocatalysis), we find it notable that this sulfenylation worked on biologically relevant scaffolds such as melatonin and tryptophan. We also found this transformation mechanistically interesting as the conditions were nearly identical to those of the benzothiazole synthesis and performed a series of mechanistic studies.

We first determined the experimental redox potentials of melatonin and 4-methylbenzenethiol sulfenylating reagent. We observed that the melatonin 18a has a first half-wave oxidation potential of 1.10V vs SCE while the latter has a higher half-wave oxidation potential of 1.49 V vs SCE (see Supporting Information). Consequently, in its excited triplet state, the \( \text{[Ir(dF(CF}_3)ppy]_2(dtbpy)} \text{PF}_6 \) photocatalyst would be out of the potential range for oxidation of 4-methylbenzenethiol, and initial oxidation likely occurs at melatonin to form the cation radical F. Stern–Volmer quenching studies between the iridium photocatalyst, melatonin, and 4-methylbenzenethiol supports this hypothesis as melatonin (\( K_{sv} = 0.2 \text{ M}^{-1}\text{L} \)) is quenched at a much higher rate than the thiophenol (\( K_{sv} = 0.1 \text{ M}^{-1}\text{L} \)) (see Scheme 6 and Supporting Information). Additionally, we ran the photocatalytic reaction in the absence of indole, observing a significant amount of the disulfide byproduct, which is known to undergo homolytic cleavage under UV light to form the thyl radical. To test whether the disulfide was an intermediate, we evaluated the reaction using phenyl disulfide as the sulfur source, observing comparable yields to that of thiophenol. Stern–Volmer quenching of the photocatalyst with 4-methylthiophenyl disulfide provided a slight increase (\( K_{sv} = 0.6 \text{ M}^{-1}\text{L} \)) relative to the thiophenol but still significantly less than melatonin. To confirm our findings, we ran a sulfenylation cross experiment using both 4-methylbenzenethiol and phenyl sulfide, observing the methylated indole as the main product via mass spectrometry (see Supporting Information). Based on these experiments, two plausible simultaneous mechanisms can occur. Once indole cation radical F is formed, deprotonated thiophenol can nucleophically attack F to form radical intermediate G, which will be oxidized by persulfate and aromatize to form the sulfenylated product 18b (Scheme 6, pathway 1). Alternatively, under photocatalytic conditions, thiophenol can be converted into disulfide which can homolytically dissociate to form the thyl radical. The radical can undergo radical coupling with F to form Wheland intermediate H, followed by aromatization to form product 18b. Both pathways can occur simultaneously; however, we believe that the nucleophilic pathway is predominant as shown by Stern–Volmer quenching studies and sulfenylation cross experiment.

In conclusion we have developed an operationally simple and economical method to synthesize benzothiazoles via photocatalytic C–H thiolation and have extended these conditions to indole sulfenylation.
nistic studies that suggest that the sulfur displays divergent activities (nucleophilic or electrophilic radical) in the two reactions.

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**Supporting Information**

Supporting information for this article is available online at https://doi.org/10.1055/s-0039-1690107.

**References and Notes**

in a 10 mL scintillation vial, thioabenzamide (1.0 equiv), azoles D.

2-(tert-Butyl)benzyl[d]thiazole (1b)

55% with no pyridine, 54% with pyridine, yellow-green solid. 1H NMR (500 MHz, CDCl3): δ = 8.82 (dt, J = 8.0, 0.8 Hz, 1 H), 7.93 (d, J = 8.1 Hz, 1 H), 7.87 (d, J = 8.7 Hz, 1 H), 7.75 (d, J = 8.7 Hz, 1 H), 7.65 (dd, J = 8.2, 6.9, 1.3 Hz, 1 H), 7.55 (dd, J = 8.2, 6.9, 1.3 Hz, 1 H), 1.59 (s, 9 H). 13C NMR (101 Hz, CDCl3): δ = 180.7, 149.2, 131.8, 131.2, 128.6, 127.9, 126.6, 125.7, 125.0, 119.0, 38.4, 31.0. MS-APCI: m/z calcd for C13H12NS [M+H]+: 242.4; found: 242.1.

N-[2-(5-Methoxy-2-[p-tolythio]-1H-indol-3-yl)ethyl]acetamide (1bb)

68%, tan solid. 1H NMR (400 MHz, CDCl3): δ = 8.12 (s, 1 H), 7.22 (d, J = 8.8 Hz, 1 H), 7.02-7.06 (m, 3 H), 6.98 (d, J = 8.3 Hz, 2 H), 6.91 (dd, J = 8.8, 2.4 Hz, 1 H), 5.50 (s, 1 H), 3.86 (s, 3 H), 3.52 (q, J = 6.4 Hz, 2 H), 3.05 (t, J = 6.5 Hz, 2 H), 2.27 (s, 3 H), 1.78 (s, 3 H). 13C NMR (126 Hz, CDCl3): δ = 170.28, 154.36, 136.17, 133.22, 132.07, 130.04, 128.13, 127.00, 119.64, 70.6. MS-APCI: m/z calcd for C23H19N2O2S [M+H]+: 355.5; found: 355.5.


(36) General Procedure for Synthesis of Substituted Benzothiazoles

In a 10 mL scintillation vial, thioabenzamide (1.0 equiv), Ru(bpy)3Cl2 (5 mol%), Na2S2O4 (2.0 equiv), and pyridine (2.0 equiv) were added to a solution of 1:1 MeCN/H2O (25 mg/1 mL). The reaction was stirred under blue LEDs for 12 h at room temperature. The resulting solution was quenched with H2O, extracted with EtOAc, dried over Na2SO4, and concentrated in vacuo. The product was purified via FCC and prep TLC with gradient from hexanes to 8:2 Hex/EtOAc.

General Procedure for the Synthesis of Sulfenylated Indoles

In a 10 mL scintillation vial, substituted indole (1.0 equiv), [Ir(dF(CF3)ppy)2(dtbbpy)]PF6, photocatalyst (1 mol%), Na2S2O4 (2.0 equiv), KOH (2.0 equiv), and thiophenol (1.2 equiv) were added to a solution of 1:1 MeCN/H2O (25 mg/1 mL). The reaction was left to stir under blue LEDs for 12 h at room temperature. The resulting solution was quenched with H2O and extracted with EtOAc, dried over Na2SO4, and concentrated in vacuo. The product was purified via FCC and prep TLC (8:2 Hex/EtOAc). Melatonin substrates were purified via FCC with (2:8 Hex/EtOAc).