

Polysulfide Anions as Visible Light Photoredox Catalysts for Aryl **Cross-Couplings**

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biaryl cross-coupling, borylation, and hydrogenation in a redox catalytic regime involving $S_4^{\bullet-}/S_4^{2-}$ and $S_3^{\bullet-}/S_3^{2-}$ redox couples.

INTRODUCTION

Visible light photoredox catalysis has advanced in the state-ofthe-art chemical synthesis, enabling us to harness lower energy visible light to productively drive various types of useful molecular transformations.¹ Homogeneous photocatalysts such as ruthenium-/iridium-based polypyridyl complexes or organic dyes could be excited under irradiation with visible light, inducing a single-electron transfer (SET) to or from organic substrates to provide reactive open-shell radical intermediates. The employment of heterogeneous semiconductors as a redox active chromophore for the synthesis of complex molecules has recently offered another contemporary trend in photoredox catalysis.² Nonetheless, the further development of new photocatalysts based on inexpensive and abundant elements that can perform productive bond formation processes in a highly efficient fashion is of prominent interest.

Sulfur is known to form various catenated homoatomic polysulfide dianions S_x^{2-} (typically, x = 2-8) and a persistent radical anion $S_3^{\bullet-}$, which is known as a blue chromophore in ultramarine blues.³ In seeking the development of alkali metals-sulfur batteries, the chemical reactivity and redox characters of polysulfide anions have been elucidated in detail.⁴ Polysulfide anions undergo complicated redox, dissociative, and disproportionation processes in the solution states to afford an equilibrium mixture of multiple polysulfide anions, and their steady states depend majorly on the solvents. The in situ spectroelectrochemical studies on the reduction of octasulfur (S₈) identified the ground state redox couples of $S_3^{\bullet -}/S_3^{2-}$ and $S_4^{\bullet -}/S_4^{2-}$, and their electrochemical potentials are estimated at around -1.35 and -0.85 V, respectively,

versus saturated calomel electrode (SCE) in dimethylformamide $(DMF)^5$ (Scheme 1 A). However, the employment of these homoatomic sulfide anions in redox catalysis that engages organic electrophores in the radical-mediated reactions remains an unmet challenge. Nonetheless, their electrochemical potentials dictate that they are incapable of inducing the single-electron reduction of unactivated organic electrophores of highly negative reduction potentials such as aryl halides $[E_{red} < 1.9 \text{ V} \text{ (vs SCE)}]$.⁶ On the contrary, these species show a characteristic absorbance in the ultravioletvisible (UV-vis) spectroscopy and some of them are observed in the visible spectral regions. For example, a degassed solution of cheap and readily available potassium polysulfide (K_2S_{xt} US \$0.12 per gram) in dimethyl sulfoxide (DMSO) shows a blue color and its steady-state UV-vis absorption spectrum indicates the presence of persistent $S_3^{\bullet-}$ (λ_{max} at 618 nm with a wide bandwidth ranging from 450 to 800 nm), S_4^{2-} (λ_{max} at 436 and 333 nm), and S_3^{2-} (λ_{max} at 273 nm) (Scheme 1 B). We posited that on the basis of the redox potentials and visible photon absorptions of S_4^{2-} and S_3^{*-} in their ground state, oxidizible S_4^{2-} could potentially serve as a photoexcited reductant whereas reducible $S_3^{\bullet-}$ could function as a photoexcited oxidant.^{1f} Therefore, we anticipated that these

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Scheme 1. Redox Potential and UV-vis Absorption of Polysulfide Anions

A. Redox potential of S_4^{+}/S_4^{2-} and S_3^{+}/S_3^{2-} redox couples

$$S_4^- + e^- \underbrace{(E_{red} = -0.85 \lor vs SCE)}_{S_4^- + e^-} S_4^{2^-}$$

 $S_3^- + e^- \underbrace{(E_{red} = -1.35 \lor vs SCE)}_{S_3^2 - S_3^{2^-}} S_3^{2^-}$

B. UV-vis spectrum of potassium polysulfide (K_2S_x) in DMSO



polysulfide anions could be engaged seamlessly in SET-driven radical-mediated processes in a redox catalytic manifold under visible light irradiation. Herein, we report the use of polysulfide anions S_4^{2-} and $S_3^{\bullet-}$ as photoredox catalysts for aryl cross-coupling reactions. The reaction design leverages the photo-excitation of S_4^{2-} to induce the single-electron reduction of aryl halides, having reduction potentials ($E_{\rm red}$) as low as -2.4 V (vs SCE). The resulting aryl radicals are engaged in (hetero)biaryl cross-coupling, borylation, and hydrogenation in a redox catalytic regime where the redox interplay between $S_4^{\bullet-}/S_4^{2-}$ and $S_3^{\bullet-}/S_3^{2-}$ redox couples enables the redox-neutral catalytic turnover.

RESULTS

Development of Biaryl Cross-Coupling. At the outset of the project, we explored if the DMSO solution of K_2S_x containing $S_3^{\bullet-}$, S_4^{2-} , and S_3^{2-} could engage aryl halides in radical coupling reactions under visible light irradiation. We selected to investigate a heterobiaryl coupling of 4'bromoacetophenone (1, $E_{\rm red} = -1.89$ V vs SCE)⁶ as an electrophore with N-methylpyrrole (2) as a radical acceptor. The current state-of-the-art strategies for such a radical-based biaryl cross-coupling' leverage highly reducing photoexcited radical anions of polyaromatic hydrocarbons as a photoexcited reductant. In particular, consecutive photoelectron-transfer processes shown by König⁸ and electrophotocatalytic strategies detailed by Lambert and Lin^{9a} and Wickens^{9b} have successfully generated excited radical anions. The leveraging of readily accessible α -aminoalkyl radicals for hydrogen-atom transfer agents was recently proven useful to promote aryl cross-coupling by Leonori, ^{10a} while the inherent net-oxidative nature of the process necessitates the use of a stoichiometric amount of the oxidant. More recently, the same group revealed that α aminoalkyl radicals serve as an initiator and a chain carrier for the aryl cross-coupling between aryl halides (especially aryl iodides) and pyrroles.^{10b} Our optimization of the reaction conditions revealed that the irradiation with blue light (λ_{max} =

Scheme 2. Photoredox Biaryl Cross-Coupling Catalyzed by K_2S_x

A. Optimized reaction conditions^a



^{*a*}Reaction conditions: **1** (0.5 mmol), **2** (20 equiv), K_2S_x (12.5 mol % per S) with H₂O (2 equiv), K_2CO_3 (1.5 equiv), DMSO (2.5 mL), 440 nm light (Kessil lamp), < 30 °C. Isolated yield of **3** was recorded.

Table 1. Screening of Precatalysts for Polysulfide Anions^a



^aReaction conditions: 1 (0.5 mmol), 2 (20 equiv), precatalysts, K_2CO_3 (1.5 equiv), DMSO (2.5 mL), 440 nm light (Kessil lamp), < 30 °C. ^{b1}H NMR yields based on the internal standard. ^cIsolated yield.

440 nm) to the mixture of 1 and 2 in the presence of K_2S_r (12.5 mol % per S atom), potassium carbonate (K₂CO₃, 1.5 equiv), and water (H₂O, 2 equiv) in DMSO enabled an efficient coupling to afford heterobiaryl 3 in 86% yield within 1.5 h (Scheme 2A). The control experiments indicated that K_2S_{xy} the irradiation with blue light, and a buffer (K_2CO_3) are all essential for the process, and the reaction is hampered under an air atmosphere (see the Supporting Information (SI), Table S1). The following synergistic catalytic cycle involving $S_4^{2-}/S_4^{\bullet-}$ and $S_3^{2-}/S_3^{\bullet-}$ redox couples is proposed (Scheme 2B). Photoexcitation by 440 nm light endows S_4^{2-} with highly reducing potential in its excited state,¹¹ allowing for singleelectron reduction of 1 to form arene radical anion I along with generation of $S_4^{\bullet-}$. The single-electron reduction of reducible $S_4^{\bullet-}$ by concomitant ground-state oxidizable S_3^{2-} allows for the regeneration of ground-state S_4^{2-} . Meanwhile, the resulting arene radical anion I undergoes mesolysis of the carbonScheme 3. Bottom-up Generation of Polysulfide Anions from Mono-Sulfides Li₂S and *i*-Pr₃SiSH

A. UV-vis spectra for charge-transfer complex of 1 with Li₂S.



halogen bond to afford aryl radical II,¹² which adds onto 2 to form the radical intermediate III. Single-electron oxidation of III by photoexcited $S_3^{\bullet -}$ followed by deprotonation liberates 3 and ground-state $S_3^{2-.13}$ We measured the quantum yield for the formation of 3 under the optimized reaction conditions, which was determined as $\Phi = 0.07$. This implicates that a possibility of the radical chain process^{10b} is less likely and supports the proposed photoredox catalytic cycle.

Evaluation of Precatalysts. We next screened the precatalysts of the polysulfide anions in the cross-coupling between 1 and 2 (Table 1). The top-down generation of polysulfide anions through the reductive fragmentation of octasulfur $(S_8)^{14}$ in the presence of sodium *tert*-butoxide (NaOt-Bu) in DMSO was amenable for the productive cross-coupling (entry 2). We also found that the bottom-up generation of polysulfide anions from monosulfide species is suitable for the catalysis. For example, the use of dilithium sulfide (Li₂S, 10 mol %) as a precatalyst led to a full conversion of 1 within 2 h to afford 3 in 88% yield (entry 3). Similarly, neutral triisopropylsilylthiol (*i*-Pr₃SiSH), which has commonly

been utilized as a hydrogen-atom transfer catalyst,¹⁵ could also perform as a promising precatalyst (entry 4).

These monosulfides (Li₂S and *i*-Pr₃SiSH) neither showed absorption at the visible region nor facilitated the crosscoupling reaction under the dark conditions due to an insufficient oxidation potential of monosulfide ions (E_{ox} of $S^{2-} = -0.76$ V vs SCE)¹⁶ (see the SI, Figures S13-S15). On the contrary, a charge-transfer absorption band was observed from the mixture of 1 and Li2S (Scheme 3A) and the irradiation with blue light (440 nm) to a mixture of 1 and Li₂S (in 1:1 molar ratio) in DMSO formed acetophenone (4), biaryl 5, and diaryl sulfide 6, all of which could be derived from the corresponding aryl radical (Scheme 3B and Figure S4). We propose that Li₂S triggers the cross-coupling process through the formation of electron-donor-acceptor (EDA) complex V with 1, which induces single-electron transfer upon the irradiation with visible light to produce a radical ion pair (Scheme 3C).¹⁷ The resulting radical anion of 1 undergoes cleavage of the C-Br bond to form the aryl radical, whereas a simultaneously formed monosulfide anion radical $(S^{\bullet -})$ undergoes dimerization to form a disulfide dianion (S_2^{2-}) and its subsequent disproportionation generates the higher order photoredox active polysulfides,¹⁸ which promote the photocatalytic turnover further. Interestingly, *i*-Pr₃SiSH might initiate the bottom-up formation of polysulfide anions in a different manner. We observed that the treatment of *i*-Pr₃SiSH with K₂CO₃ in DMSO immediately stains the solution blue, and the UV-vis absorption spectroscopy unambiguously indicated the generation of polysulfide anions $(S_3^{\bullet-}, S_4^{2-}, S_4^{2-})$ and S_3^{2-}) (Figure S16). The nuclear magnetic resonance (NMR) spectroscopy showed the formation of disulfide (i- $Pr_3SiS)_2$ VI in the solution (Figure S5). Therefore, we postulated that DMSO functions as an oxidant¹⁹ to promote the desilylative oligomerization of *i*-Pr₃SiSH to the higher order polysulfides via disulfide VI (Scheme 3D). The capability of disulfide VI as the catalyst was ascertained as it performed the productive cross-coupling (Table S2).

Substrate Scope on Biaryl Cross-Coupling. We found that this photoredox protocol with polysulfide anions is capable of engaging a wide range of aryl halides for the (hetero)biaryl coupling (Scheme 4A). We first studied the reactivity of 4'-chlolroacetophenone (7), having a reductively inert C-Cl bond.²⁰ We observed a diminished efficiency in the reaction with K_2S_x (12.5 mol % per S atom), resulting in an incomplete conversion of 7 (60%) even after irradiation for 22 h (Table S3). We found that the use of Li₂S and *i*-Pr₃SiSH results in the completion of the process within 4 h to give coupling product 3 in 80% and 75% yields, respectively. These outcomes suggested that a bottom-up preparation of the polysulfide anions from mono sulfides would provide more productive reactivity especially for reductively recalcitrant aryl halides. The method allows for the installation of various polar- π electron-withdrawing groups susceptible to reductive reaction conditions, such as ketone (7-10), aldehyde (11-13), nitrile (14), and ester (15). The protocol could successfully engage five-membered ring heteroaryl halides based on furan (16), thiophene (17, 18), and thiazole (19). The chemistry was also extended to functionalize sixmembered ring heteroaryl halides such as pyridine (20, 21), quinoline (22), and pyrazine (23). We also found that nonactivated aryl halides having a highly negative reduction potential $(E_{red} > -2.4 \text{ V vs SCE})^{21}$ are suitable substrates (24– 27). In these cases, the employment of Li_2S or *i*-Pr₃SiSH (10)

Scheme 4. Reaction Scope on (Hetero)biaryl Cross-Coupling

A. Scope of aryl halides^a cat. S₄²⁻/S₃⁻/S₃²⁻ K₂CO₃ Ar-X DMSO (0.2 M) (X = Br, Cl)Kessil lamp (440 nm) <30 °C CI Br онс онс сно NC MeO₂C 13 8 10 11 12 14 g 15 80%^b (75%)^c 93% 76% 71% 68%^c 79% 67% 81% $66\%^b$ Br Ac B Ν OH NC м́ Br 16 17 18 19 20 21 22 23 59%^b 80% 47%^b 78%^b 75% 81%^b 50%^c 90% MeC CO₂Me Br Br ö Me MeO. MeO Ph 0 Me 27 28 24 25 26 $62\%^{b,d}$ 20%^{b,d} $60\%^{t}$ 58% 64% 29 30 $E_{red} = -2.19 V$ $E_{red} = -2.43 \text{ V}$ $E_{\rm red} = -2.72 \text{ V}$ 82%^b vs SCE 83% vs SCE vs SCE Me Polyhalogenated aromatics NMe онс OHC MeO₂C B MeO₂C 92% 31 80%^b 32 83%^b 33 B NC NC Br 71%^b 34 70%^t 35 B. Scope of arene coupling partners^a



"Reaction conditions: substrates (0.5 mmol), coupling partners (20 equiv), K_2S_x (12.5 mol % per S) with H_2O (2 equiv), K_2CO_3 (1.5 equiv), DMSO (2.5 mL), 440 nm light (Kessil lamp), < 30 °C. Isolated yields of the products were recorded. ^bPrecatalyst: Li₂S (10 mol %). ^cPrecatalyst: *i*-Pr₃SiSH (10 mol %). ^d390 nm light. ^eTen equiv of 3-methylindole was used. ^fThe reaction was run with 5 equiv of 4-methoxyphenol and 5 equiv of NaO*t*-Bu.

mol %) as a precatalyst was optimal. However, the reaction of reductively more inert 4-bromoanisole (28) ($E_{red} = -2.72$ V vs

SCE²² was sluggish. This protocol was found to be capable in the functionalization of nicergoline (29) and indomethacin

Scheme 5. Borylation and Hydrodehalogenation

A. Borylation^a



^{*a*}Reaction conditions: substrates (0.5 mmol), B₂ pin₂ (2 equiv), K₂S_{*x*} (12.5 mol % per S), TMG (1.5 equiv), CH₃CN (5 mL), 440 nm light (Kessil lamp), < 30 °C. Isolated yields of the products were recorded. ^{*b*}Precatalyst: *i*-Pr₃SiSH (10 mol %). ^{*c*}substrates (0.5 mmol), *i*-Pr₂NEt (2 equiv), *i*-Pr₃SiSH (10 mol %), H₂O (20 equiv), CH₃CN (5 mL), 440 nm light (Kessil lamp), < 30 °C.

methyl ester (30) without damaging the other functional groups in these substrates. Finally, we explored if polyhalogenated aromatic substrates could be engaged in chemoselective cross-coupling processes. We were pleased to observe that 2-bromo-4-chlorobenzaldehyde (31) was selectively functionalized on the C-Br bond. Similarly, the coupling reaction of 4-bromo-2-fluoro-1,1'-biphenyl (32) occurs selectively at the C-Br bond. A more reactive C-I bond²³ could be functionalized selectively in the reactions of methyl 3bromo-5-iodobenzoate (33) and 1-bromo-4-iodobenzene (34). Moreover, 3,5-dibromobenzonitrile (35) was found to undergo single functionalization at one of the C-Br bonds. The scope with respect to the trapping (hetero)arenes was next evaluated (Scheme 4B). The use of N-H pyrrole (36) and indole (37) was found to be optimal, while the coupling with thiophene (38) resulted in moderate efficiency. The protocol enables the Minisci type-coupling with pyrazine (39).

Scheme 6. Scale-up in Flow^a

A. Aryl cross-coupling in flow



^{*a*}O'D' = outer diameter; I'D' = inner diameter; $T_{\rm R}$ = residence time; V = volume of the micro-tubing reactor; BPR = back pressure regulator.

Electron-rich benzenes (40, 41) were also found to be compatible as a coupling partner.

Development of Borylation and Hydrodehalogenation. The synthetic utility of this polysulfide anions-based photoredox catalysis was further extended to the dehaloborylation reaction by employing bis(pinacolato)diboron (B₂ pin_2) as the radical trapping reagents (Scheme 5A and Table \$4).^{24,25} The optimization for the borylation of 4'bromoacetophenone (1) with K_2S_r as a precatalyst led to the identification of tetramethylguanidine (TMG) and acetonitrile (CH₃CN) as the optimal base and solvent, respectively, delivering pinacol arylboronate 42 in 84% yield within 1.5 h. This protocol was found to be applicable to the borylation of various functionalized haloarenes (43-47). We also found that the protocol is amenable to hydrodebromination of 1 using diisopropylethylamine (*i*-Pr₂NEt) as a hydrogen donor, providing acetophenone (4) in 96% yield within 2 h (Scheme 5B and Table S5).^{22,26,27} The identified reaction conditions were capable of reductive radical cyclization of 48 to dihydrobenzofuran 49 and hydrodeiodination of secondary alkyl iodide **50** $(E_{\rm red} = -2.35 \text{ V})^{10a}$ to **51**.

Scale-up in Flow. These batch photoredox processes catalyzed by the polysulfide anions stimulated us to explore the scalability of the heterobiaryl cross-coupling and borylation in flow (Scheme 6). The cross-coupling between 1 and 2 was efficiently promoted in a homogeneous system using *i*-Pr₃SiSH as a precatalyst and tetramethylguanidine (TMG) as a base in an operationally simple microtubing continuous-flow reactor²⁸ (Figure S8). The desired product 3 was delivered at a 1.75 g/ hour production rate (78%) with 30 min as the residence time (Scheme 6A). The debromo-borylation of 1 could also be performed in the same flow reactor to afford 42 at a 6.9 g/hour production rate (83%) with 20 min as the residence time (Scheme 6B).

The key enabling advance in the present method takes advantage of unique reactivity of the polysulfide anions as photoredox catalysts, which are capable of engaging a wide range of aryl halides in productive radical cross-coupling chemistry. We anticipate that the broad scope with wide functional group compatibility, operational simplicity, and scalability in flow would bring useful and practical applications of this catalysis strategy in various fields.

ASSOCIATED CONTENT

G Supporting Information

The Supporting Information is available free of charge on the ACS Publications Web site. Procedures and characterization data (PDF) The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c11968.

Discussions of characterization methods used, reaction optimization, experimental protocols, determination of the quantum yield, light on/off experiment, stoichiometric reactions, characterization of products, proposed mechanisms, scale up in flow, and preliminary mechanistic investigation, figures of emission spectra, batch reaction setup, reaction profiles, proposed mechanisms, NMR spectra, proposed catalytic cycles, absorption spectra, UV–vis spectra, steady state fluorescence spectrum, lifetime decay profiles, and reaction pathways, and tables of reaction optimizations (PDF)

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Notes

The authors declare no competing financial interest.

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