ChemComm



COMMUNICATION

View Article Online
View Journal | View Issue



Cite this: *Chem. Commun.*, 2022, 58, 10241

Received 9th June 2022, Accepted 26th July 2022

DOI: 10.1039/d2cc03262g

rsc.li/chemcomm

Utilizing a needle as a source of iron in synergistic dual photoredox catalytic generation of alkoxy radicals†

transformations (Scheme 1A).

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A visible-light mediated alkoxy radical generation is described, which allows for a structurally divergent oxidative $C(sp^3)-H$ bond functionalization. This protocol employs a photoredox/iron dual catalysis allowing for an unprecedented chemoselective single-step transformation of alcohol derivatives giving access to two valuable sets of products, tetrahydrofurans and γ -bromoketones, under one set of conditions. Addition of iron, by slow corrosion of a needle, provides superior reaction efficiency as compared to various iron(III) complexes.

Oxygen-centered radicals are open-shell, highly reactive species that can participate in a variety of valuable reactions. Despite their rich synthetic possibilities, there is a limited number of reliable strategies for their generation and utilization in the construction of molecular complexity. Typically, facile generation of alkoxy radicals often requires stoichiometric quantities of very strong oxidants.²⁻⁷ Consequently, the most common approach was the employment of alkoxy radical precursors. However, the preparation and/or utilization of historically common alkoxy radical precursors often involves explosive reagents or strong oxidants, 8,9 tedious synthesis 10 or unstable nature of precursor itself,11 necessity of organotin initiator,10 UV irradiation or elevated temperatures 12-14 which naturally diminished their synthetic potential. More recently, photocatalysis 15-17 has emerged as a powerful platform for alkoxy radical generation under mild conditions. Recent work from Knowles group (Scheme 1B)18 has shown that hydroxyl group can be directly oxidized via reductive quenching of a photocatalyst, thereby generating an oxygencentered radical. In efforts toward alkoxy radicals, research groups of Zhu (Scheme 1B)19,20 and Almqvist21 have developed protocols for efficient alkoxy radical initiated ring-opening of tertiary cycloalkanols via β-scission of cyclic C-C bonds. The

radicals and photoredox catalysis, we have investigated the possibility to access these moieties utilizing *N*-alkoxythiazolethione precursors. Given their capacity as an efficient alkoxy radical source.²⁴ we hypothesized that redox-active thiazolethione (Thi)

research group of Chen,22 as well as our own laboratory,23 have

recently reported an approach relying on oxidative quenching of

the photocatalyst, utilizing hypervalent iodine(III) alkoxy radical

precursors in 1,5-hydrogen atom transfer (1,5-HAT) mediated

As a part of our ongoing engagement in oxygen-centered

source,²⁴ we hypothesized that redox-active thiazolethione (Thi) functionality could be strategically employed in oxidative quenching of the photocatalyst, subsequently activating δ -C(sp³)-H position in new C-O bond-forming events. Herein, we present a new method that allows for a structurally divergent strategy providing the opportunity to selectively transform linear alcohols to cyclic

A. Our previous work: alcohol etherification utilizing hypervalent iodine (III) alkoxy radical precursors

ethers and linear γ -bromoketones by employing a synergistic dual

C. This work: alcohol etherification and ring-opening of cyclic ethers utilizing alkoxy radicals



Scheme 1 (A) Our previously reported etherification method utilizing hypervalent iodine(III) alkoxy radical precursors, (B) previously reported methods towards halogenated ketones utilizing alkoxy radicals, (C) visible-light mediated etherification and bromoketone synthesis utilizing alkoxy radicals and 1,5-HAT.

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† Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d2cc03262g

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catalytic system using a photocatalyst in combination with a needle acting as a source of Fe (Scheme 1C).

Initial efforts were focused on establishing an efficient 1,5-HAT system. For this, N-alkoxythiazolethione 1, derived from 4phenylbuthanol, was chosen as our model substrate to allow for an efficient δ -C(sp³)-H abstraction together with subsequent oxidation and cyclization giving the corresponding tetrahydrofuran derivative. Bromotrichloromethane (CBrCl₃) was chosen as external oxidant since it is known to oxidatively quench several common photoexcited catalysts.²⁵ Extensive screening of photocatalysts, solvents, additives, and stoichiometry of the reagents was conducted (Table 1 and ESI†).

Surprisingly, initial experiments yielded a mixture of tetrahydrofuran 2 and bromoketone 3. This rather intriguing result opens for the possibility of a chemodivergent method providing access to two distinctively different structural features by simple tweaking of the conditions. Monitoring the reaction over time revealed that tetrahydrofuran 2 serves as an intermediate for bromoketone 3 formation (see ESI†). Consequently, we set out to optimize the conditions for both 2 and 3. The optimal loading of CBrCl₃ towards tetrahydrofuran 2 was found to be 1.5 equiv. (62%, entry 1). If the loading was lowered, even after prolonged reaction time, the reaction did not proceed with full conversion. With 10 equiv. of CBrCl₃ we were able to obtain satisfactory results for bromoketone 3 (64%, entry 2). Increasing the temperature to 55 °C increased the yield slightly, but more importantly, did so with a significantly shortened reaction time from 15 h to 6 h (entry 4). To further facilitate the ring-opening of 2, we reasoned that an external source of bromide would increase the yield of desired ketone 3. Various external bromide sources were evaluated (entries 5-7) and we were pleased to find that the addition of tetrabutylammonium bromide (TBAB) led to almost a quantitative 95% yield of 3

Table 1 Summary of main optimization experiments

				Yield ^e (%)	
Entry	$Ru(bpy)_3 (mol\%)$	CBrCl ₃ (equiv.)	Additive	2	3
1^a	5	1.5	_	62	43
2^a	5	10	_	23	64
3^a	5	20	_	_	69
4^{bc}	5	10	_	_	66
5^{bc}	5	10	LiBr	49	15
6^{bc}	5	10	NBS	28	9
7^{bc}	5	10	TBAB	_	95
8^{bc}	5	1.5	TBAB	54	24
9^b	5	10	TBAB	72	19
10^{bd}	5	10	TBAB	_	_
11^b	0	10	TBAB	_	_

^a Reaction time 15 h, reaction temperature 25 °C. ^b Reaction time 6 h, reaction temperature 55 °C. c Reaction performed with a metallic needle inserted into the system. d Reaction performed in the dark. Yields were determined by ¹H NMR using ethylene carbonate as an internal standard.

(entry 7). Throughout the initial optimization, discrepancies in the results were observed for several conditions. After meticulous investigation of all variables, we observed that having a metallic needle, used as a part of the experimental setup to keep the system under a N2 atmosphere, present during the entire reaction time, reproducible yield of 3 was obtained (entry 7). Employing identical conditions with the needle removed provided 2 in 72% yield (entry 9). The notable difference in reaction outcome can be explained by that earth abundant transition metals present in the needle entered the reaction solution and mediated the conversion of 2 to 3. Observation of slow corrosion of the metallic needle, likely due to formation of HBr during the progression of the reaction, is consistent with this assumption. In our previous efforts, we have observed that ring-opening of cyclic ethers can be facilitated under similar conditions, in the presence of trace amount of transition metals,²⁶ which further supports the hypothesis of metal catalysis. Therefore, the metal composition of the needle was evaluated by XRF, and major metals present in the needle were tested in our system (see ESI†). Although the evaluated metal complexes (Fe[III], Ni[II], Cr[III]) mediated the conversion of 2 to 3, none could match the efficiency of introducing the needle into the system. ICP-MS analysis of the crude reaction mixture clearly showed that the main metals present in the needle indeed did corrode into the reaction mixture (see ESI†). Lastly, no products were detected in control experiments where the photocatalyst or blue light irradiation were excluded from the system supporting their necessity in the given transformation (entry 11 and 12).

With optimized conditions in hand, we explored the scope of N-alkoxythiazolethione derivatives. Initially, we dedicated our efforts toward primary N-alkoxythiazolethione derivatives (Scheme 2). Aromatic substituents with various electronic properties are well tolerated for both transformations. Both electron-donating and electron-withdrawing aryl substituents were evaluated, and we were pleased to obtain both tetrahydrofurans 2a-2j and corresponding bromoketones 3a-3j derived from primary alcohols in good to excellent yields. Worth noting is that a p-nitro functionality, which is very often found to be problematic in photoredox catalysis, was compatible with our conditions and gave rise to tetrahydrofuran 2c in 54% yield and the corresponding bromoketone 3c in high 78% yield. Aromatics bearing p-halo substituents gave rise to tetrahydrofurans 2f, 2h and 2j in high yields (94%, 84% and 79%) and 2g with p-chloro substituted phenyl ring in slightly lower 55% yield. In terms of their subsequent conversion to the corresponding bromoketones 3f-3i, these were obtained in good to high yields (65-86%). Additionally, the electron rich heteroaromatic thiophene moiety was also compatible with this transformation and delivered tetrahydrofuran 2j and the corresponding ketone 3j in good 58% and 50% yields, respectively. These examples illustrate that the process has a broad scope in terms of the electronic nature of the arene moiety.

Next, we turned to explore the scope of secondary alkoxy radicals where the 4-phenylbuthanol fragment of the molecule was retained while the α-substituent of the alcohol was

3i 50%

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blue LED blue LED 2a - i 3a - ja 1a - j C - O bond formation starting precursor remote halogenation C - O bond formation starting precursor remote halogenation 3a 90% (83%)b 2f 94% 2a 62% (61%)b 3f 65% **2b** 95% 3b 85% 2g 55% 1g **3g** 68% 1b 1h 2h 84% 3h 84% 2c 54% 3c 78% 3d 50% 2i 79% **3**i 86% 2d 59%

Scheme 2 Scope of primary N-alkoxythiazolethione derivatives. Reaction conditions: **1** (0.1 mmol), Ru(bpy)₃(PF₆)₂ (5 mol%), CBrCl₃ (10 equiv.), TBAB (1.1 equiv.) in CH₃CN (0.05 M), blue LED irradiation for 6 h at 55 °C, ^a needle inserted into the vial, ^b reaction conducted on 1 mmol scale. Yields refer to isolated products.

3e 68%

alternated (Scheme 3). We were pleased to find that all investigated secondary N-alkoxythiazolethiones led to desired tetrahydrofurans 2k-2r as well as corresponding bromoketones 3k-3r in good to high yields. Both methyl and sterically demanding isobutyl substituted substrates provided tetrahydrofurans 2k and 2l in 68% and 70% yields, respectively. Their corresponding bromoketones 3k and 3l were obtained with yields of 64% and 65%, respectively. For longer butyl substituents, two possible pathways for 1,5-HAT are available. However, a benzylic C(sp³)-H bond is more activated towards 1,5-HAT, resulting in more thermodynamically favoured carbon-centered radical. Indeed, tetrahydrofuran 2m was isolated as a major regioisomer in 58%. The formation of the other possible regioisomer was observed in negligible quantity. Bromoketone 3m was obtained in slightly lower 48% yield. Considerably bulkier phenethyl N-alkoxythiazolethione yielded desired tetrahydrofuran 2n in 75% and bromoketone 3n in 65% yields. Next, we moved our attention towards α-aryl-substituted N-alkoxythiazolethiones (10–1r). The transformation to tetrahydrofurans 20, 2p and 2r proceeded smoothly in 69%, 59% and 63% yields, respectively. The reaction of α -phenyl substituted N-alkoxythiazolethione alcohol and its p-chloro derivative, proceeded towards bromoketone 3p and 3r in 70% and 62% yields, respectively, with reaction time shortened to 3 h. Longer reaction time converted bromoketones 3p and 3r into their corresponding dehalogenated derivatives. This result can be rationalized by the ability of Ru(II) to initiate visible-light mediated dehalogenation in the presence of HCCl₃ as H-atom donor. 15,27 Electron-rich o-methoxy substituted derivative provided trichloromethyl functionalized tetrahydrobenzoxepine 30 in 54% yield (see ESI†).

To demonstrate the synthetic utility of γ -bromoketones, we converted 4-bromo-1-phenylbutane-1-one 3 to various structural

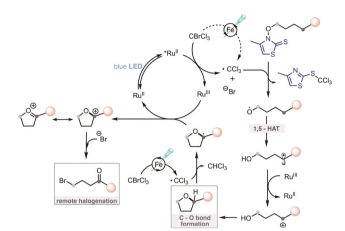
Scheme 3 Scope of secondary N-alkoxythiazolethione derivatives. Reaction conditions: ${\bf 1}$ (0.1 mmol), Ru(bpy) $_3$ (PF $_6$) $_2$ (5 mol%), CBrCl $_3$ (10 equiv.), TBAB (1.1 equiv.) in CH $_3$ CN (0.05 M), blue LED irradiation for 6 h at 55 °C, a needle inserted into the vial, b reaction time 3 h, c regioisomeric ratio. Yields refer to isolated products.

entities of general significance (Scheme 4). Asymmetric reduction of 3, followed by ring closure generated (*S*)-2-phenyltetrahydrofuran 4 in 87% yield with 86% ee. Functionalized 2-methyl-2-phenyltetrahydrofuran 5 was obtained in 85% yield whereas

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Scheme 4 Demonstration of synthetic applications of γ-bromoketones

ulating the presence of an earth-abundant metal catalyst in combination with a common photocatalyst, we have shown that it is possible to selectively access cyclic ethers and γ -bromoketone derivatives under visible-light irradiation conditions. The protocol is operationally simple and allows for synthetic diversity for simple alcohols under one set of conditions, giving rapid access to a broad collection of structurally diverse compounds.



Proposed mechanism based on generated data

cyclopropane 6 and dihydrooxazine 7 were generated in nearly quantitative yields.

A plausible mechanism for the developed process is depicted in Scheme 5. The photoexcited catalyst Ru^{II^*} ($E_{1/2}^{II^*/III} = -0.81 \text{ V vs.}$ SCE in CH₃CN)¹⁵ is oxidatively quenched by CBrCl₃ to afford Ru^{III} along with •CCl₃ and Br⁻. Addition of •CCl₃ radical to the thione sulfur, followed by the homolytic cleavage of the O-N bond liberates a desired oxygen-centered radical. Subsequently, the alkoxy radical undergoes 1,5-HAT generating a stabilized benzylic carbon-centered radical. This radical intermediate ($E_{ox} = ca. 0.35 \text{ V}$ vs. SCE in $CH_3CN)^{28}$ can be further oxidized by Ru^{III} ($E_{1/2}^{III/II} = 1.29 \text{ V}$ vs. SCE in CH₃CN)¹⁵ into its corresponding carbocation. This oxidation event returns the photocatalyst to its ground state. The new C-O bond is formed via intramolecular cyclization yielding the tetrahydrofuran derivative. Further transformation into the corresponding bromoketone proceeds via hydrogen abstraction mediated by another molecule of °CCl3 radical affording cyclic radical intermediate ($E_{ox} = ca. 0.16 \text{ V} \text{ vs. SCE in CH}_3\text{CN}$)²⁸ which can be oxidized by RuIII (or alternatively CBrCl3) into the corresponding cyclic carbocation. This process is further facilitated by an iron catalyzed generation of both °CCl3 and carbocation under blue-light irradiation.²⁶ Lastly, the ring-opening event of carbocation by bromide ion provides the final bromoketone.

In summary, this study set out to develop a new method for alkoxy radicals generation under mild, photon-driven conditions,

Conflicts of interest

There are no conflicts to declare.

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