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Photoredox-catalyzed sulfonylation of diaryliodonium salts with DABSO and silyl enolates involving the insertion of SO_2^+

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A versatile photoredox-catalyzed three-component sulfonylation of diaryliodonium salts with DABSO and silyl enolates involving the insertion of SO₂ was developed. Moreover, by employing β -alkyl substituted silyl enolates as substrates, the sulfonylation would give α -alkyl substituted β -keto sulfones, which are difficult to accessed by previous method involving the insertion of $SO₂$.

b-Keto sulfones are privileged frameworks in many pharmaceuticals and display remarkable biological activities and pharmacological properties such as anti-bacterial, anti-fungal, anti-hepatitis, and non-nucleoside inhibition (Fig. 1).¹ Additionally, b-keto sulfones are utilized as versatile intermediate synthons in diverse synthetic transformations due to the simultaneous existence of multiple functional groups including carbonyl, sulfonyl and active methylene moieties.² Therefore, there is high demand for developing efficient synthesis methods to construct β -keto sulfones.

Conventional approaches for the synthesis of β -keto sulfones include oxidation of 2-oxo-sulfides with strong oxidants³ and the sulfonylation of α -halo-ketones with preinstalled sulfonylcontaining segments,⁴ which have several drawbacks, such as the employment of poorly accessible and smelly organosulfur compounds as starting materials, limited substrate applicability, and harsh conditions. In recent years, sulfonylation involving the insertion of sulfur dioxide via a radical process has emerged as a powerful method to access sulfone derivatives,⁵ in which various SO_2 surrogates, like DABSO, metabisulfites, rongalite reagents and SOgen, are employed instead of SO₂ gas.⁶ Outstanding contributions to synthesize diverse β -keto sulfones from silyl enolates via sulfur dioxide insertion have been made by Wu's groups using aryldiazonium tetrafluoroborates,⁷ aryl/ alkyl halides,⁸ and thianthrenium salts⁹ as active radical precursors in some cases. These transformations proceed with good tolerance of functional groups, easily enabling the incorporation of various sulfonyl skeletons into ketones.

On the other hand, diaryliodonium salts can participate in various transformations as a type of versatile, easily available, non-toxic, environmentally, and air stable solid arylating reagent.¹⁰ So far, diaryliodonium salts as active radical precursors in multi-component sulfonylation reactions involving the insertion of sulfur dioxide has been developed initially. Jiang and coworkers reported a straightforward protocol for the synthesis of diverse functionalized diarylannulated sulfones through SO_2/I exchange of iodonium(III) salts. In this reaction, the aryl radical generated from diaryliodonium salt was captured by $\text{Na}_2\text{S}_2\text{O}_5$ to form a SO_2 radical anion, which served as the key intermediate to realize the exchange strategy (Scheme 1a).¹¹ In 2018, Manolikakes¹² and Zhang¹³ demonstrated that visible-light-induced reduction of diaryliodonium salts would generate aryl radicals, which could undergo cascade cyclization in combination with $SO₂$ for the synthesis of sulfonylated coumarins (Scheme 1b) and 3-arylsulfonylquinoline derivatives, respectively. Very recently, Piguel and coworkers established the first visible-light photoredox catalyzed C-H sulfonylation of imidazopyridines with diaryliodonium salts and DABSO for the straightforward synthesis of novel C-3 PAPER
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Fig. 1 Examples for β -keto sulfones and related sulfones with biological activities.

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$$
\begin{array}{c}\n\text{OSiMe}_{3} \\
\downarrow \\
\text{PABSO} \\
\downarrow \\
\text{Ph}_{2} \text{LOTf}\n\end{array}\n\quad\n\begin{array}{c}\n4 \text{CzIPN} \\
\text{blue LED} \\
\text{dioxane, rt} \\
\downarrow \\
\text{dioxane, rt}\n\end{array}\n\quad\n\begin{array}{c}\n0 & Q \\
\downarrow 0 \\
\downarrow 0 \\
\downarrow 0\n\end{array}
$$

Scheme 1 Diaryliodonium salts as active radical precursors in the insertion of sulfur dioxide.

sulfonylated imidazoheterocycles (Scheme 1c).¹⁴ Herein, we would like to report a novel 4CzIPN-catalyzed three-component sulfonylation of diaryliodonium salts with DABSO and silyl enolates via a radical process involving SO_2 insertion. This photoredox catalysis with the assistance of visible light represents a green and sustainable approach for the synthesis of β keto sulfones (Scheme 1d).

Trimethyl((1-phenylvinyl)oxy)silane 1a, diphenyliodonium tri flate 2a and 1,4-diazabicyclo^[2.2.2]octane bis(sulfur dioxide) (DABSO) were selected as starting materials to optimize the sulfonylation conditions. Initially, irradiation of all three reaction partners in 1,4-dioxane with blue LED (25 W) at room temperature for 12 hours under N_2 led to the desired β -keto sulfone 3a in 47% yield (Table 1, entry 1). Other SO_2 surrogates such as $K_2S_2O_5$, $Na₂S₂O₅$, $Na₂S₂O₄$ displayed lower activity (Table 1, entries 2–4). A variety of solvents were examined. 1,4-Dioxane proved to be superior to tetrahydrofuran (THF), acetonitrile, 1,2-dichloroethane (DCE), N,N-dimethylformamide (DMF) and toluene (Table 1, entries 1 and 5–10). When SO_2 -MeCN solution was used as SO_2 source instead of DABSO, the reaction mixture in MeCN gave 3a in a slightly increased yield (Table 1, entries 6 and 7). Next, a range of experiments were carried out under various light sources such as blue LED (10 W, 25 W, 30 W), white LED (24 W), compact fluorescent lamp (CFL, 18 W), Kessil light (390 nm) and UV (600 W). As a result, 25 W of blue LED has the best performance (Table S1 in ESI†). Despite extending the time to 24 h, only 55% yield of 3a can be obtained (Table S1,† entry 8). The addition of photoredox catalysts was beneficial for the transformation. For example, the yield of 3a sharply rose to more than 80% when 2.0 mol% of either $[\text{Ir{dFCF}_3ppy}_{2}](bpy)]PF_6$ or 1,2,3,5tetrakis(carbazol-9-yl)-4,6-dicyanobenzene (4CzIPN) was added (Table 1, entries 11 and 14). Considering cost and availability

Table 1 Optimization of three-component sulfonylation

^a Reaction conditions: 1a (2.0 equiv.), 2a (0.1 mmol, 1.0 equiv.), $[SO_2]$ (2.0 equiv.), PC (2 mol%), and $n-C_{12}H_{26}$ (10 μ L) in solvent (1.0 mL) at room temperature for 12 h, irradiation with 25 W blue LED, under N₂. GC yields. ^c Under air atmosphere. ^d No light. Ir-1 = [Ir ${dFCF_3ppy}_2(bpy)]PF_6$, Ru-1 = [Ru(bpy)₃]Cl₂·6H₂O. SO₂ solution: SO₂ solution in MeCN (∼7.9 M).

factors, 4CzIPN was chosen as the optimal photocatalyst. When the reaction was carried out under air atmosphere, 60% yield of 3a was generated (Table 1, entry 15). In addition, the threecomponent sulfonylation cannot proceed without light (Table 1, entry 16). In addition, the introduction of various bases, such as NaO^tBu, NaOH, Na₂CO₃, NaHCO₃ and NEt₃, has caused the reaction yield to decrease to varying degrees (Table S2 in ESI†).

We investigated the scope of this three-component sulfonylation transformation with the optimal conditions in hand. The treatment of diverse silyl enolates 1 with diphenyliodonium triflate 2a and DABSO afforded the corresponding β -keto sulfone derivatives in moderate to good yields (Scheme 2). Silyl enolates possessing methyl and alkoxy groups on the phenyl ring could react with 2a and DABSO smoothly to give the desired products (3b–3h). Halogen atoms, especially chlorine, can be compatible under the optimal conditions (3i–3l), which revealed possible further transformations of the resulting halogenated products with other nucleophiles.¹⁵ Representative electron-withdrawing groups, such as cyano, nitro, and tri fluoromethyl, were also tolerated in this three-component sulfonylation transformation under the optimal conditions (3m– 3p). When trimethyl((1-(naphthalen-2-yl)vinyl)oxy)silane was treated with 2a and DABSO, the desired 3q was isolated in 61% yield. In addition, heterocyclic substituted silyl enolate could also be transformed to 3r in 59% yield. It is notable that α substituted β-keto sulfones cannot be obtained from reported radical transformations involving the insertion of SO_2 .⁷⁻⁹ To our

delight, β-alkyl substituted silyl enolates could work well in this three-component sulfonylation to give a-alkyl substituted bketo sulfones 3s and 3t in acceptable yields.

Subsequently the scope of diaryliodonium salts was examined. As shown in Scheme 3, both diaryliodonium triflates and diaryliodonium tetrafluoroborates were suitable substrates for this photocatalytic reaction. Diaryliodonium salts bearing different functional groups, containing electron-donating methyl (4a and 4b), tert-butyl (4c), methoxy (4d) and electronwithdrawing fluorine $(4e)$, chlorine $(4f)$, bromine $(4g)$, trifluoromethyl $(4h$ and $4i)$ and nitro $(4j$ and $4k)$, were readily compatible in this 4CzIPN-catalyzed three-component sulfonylation.

We next investigated the 4CzIPN-catalyzed three-component sulfonylation with unsymmetrical diaryliodonium salts. When unsymmetrical salt possessing a p-methylphenyl and a bulky mesityl group was used, only 4a was obtained in 68% via a selective transfer of the p-methylphenyl group (Scheme 4a). The reaction of 5b afforded methoxy substituted sulfone 4d and cyano substituted sulfone 4l in 15% and 62%, respectively (Scheme 4b). When unsymmetrical diaryliodonium salt 5c reacted with 1a and DABSO, the β -keto sulfone 4f was isolated as the main product in 65% with selective transfer of the electrondeficient p -chlorophenyl group over the electron-rich p methoxyphenyl moiety (Scheme 4c). These results indicate that

Scheme 3 The scope of diaryliodonium salts. a Diaryliodonium tetrafluoroborates was used.

Scheme 4 Reactions with unsymmetrical diaryliodonium salts: (a) the sulfonylation with p-tolyl-mesityl iodonium salt 5a; (b) the sulfonylation with (p-cyanophenyl)-(p-methoxyphenyl) iodonium salt 5b; (c) the sulfonylation with (p-chlorophenyl)-(p-methoxyphenyl) iodonium salt 5c.

the electron-deficient aryl moiety is more easily reduced by photocatalyst to generate an aryl radical.

The scope of enolates in addition to silyl enolates was checked. 1-Phenylvinyl acetate 6 can afford the desired product in moderate yield in the presence of either Ir-1 or 4CzIPN (Scheme 5a), while 1,3-diketone derived enol ester 7 was not compatible (Scheme 5b). Under the standard conditions, the combination of acetophenone and sodium tert-butoxide was adopted, which can generate enolate in situ to replace the enol silane. As a result, the generation of 3a was not observed (Scheme 5c).

Scheme 5 The scope of other enolates: (a) the sulfonylation with 1 phenylvinyl acetate 6; (b) the sulfonylation with 1,3-diketone derived enol ester 7; (c) the sulfonylation with acetophenone in the presence of sodium tert-butoxide.

To gain more insight into the mechanism, the following radical inhibition experiments were performed. The standard sulfonylation transformation was completely restrained in the presence of 3.0 equiv. of 2,2,6,6-tetramethylpiperidine 1-oxyl (TEMPO) (Scheme 6a). At the same time, the yield of 3a decreased to 47% when 3.0 equiv. of butylated hydroxytoluene (BHT) was added in the reaction system (Scheme 6b). These results implied that a radical process might be involved in the mechanism.

Based on the above observations and previous work, plausible mechanism of the photoredox-catalyzed sulfonylation involving the insertion of SO_2 is proposed. For the pathway in the presence of photocatalyst (Scheme 7, path a):⁹ The photoexcited 4CzIPN* reduces diaryliodonium salt 2 through single electron transfer (SET) process to give an aryl radical and the oxidized photocatalyst A. Aryl radical is trapped by SO_2 to generate the sulfonyl radical IM1. Then, the addition of sulfonyl radical IM1 into the double bond of silyl enolate 1 affords radical intermediate IM2, which can be oxidized to cation intermediate IM3 by photocatalyst A via another SET along with the regeneration of the photocatalyst 4CzIPN. Finally, cation intermediate IM3 undergoes desilylation with nucleophilic anion species to give the desired β -keto sulfone 3. For the pathway in the absence of photocatalyst (Scheme 7, path b):¹² the interaction of iodonium salt with DABSO $(DABCO \cdot (SO_2)_2)$ would produce DABCO radical cation, dioxide, and diaryl iodine radical. Fragmentation of diaryl iodine radical furnishes an aryl

Scheme 6 Parallel control experiments: (a) radical inhibition experiment with TEMPO; (b) radical inhibition experiment with BHT.

Scheme 7 Plausible mechanistic pathway

radical, which is trapped by $SO₂$ to generate the sulfonyl radical IM1. Then, the addition of sulfonyl radical IM1 into the double bond of silyl enolate 1 affords radical intermediate IM2, which can be oxidized to cation intermediate IM3 by DABCO radical cation. Finally, cation intermediate IM3 undergoes desilylation to give 3.

Conclusions

In summary, a versatile strategy for the synthesis of diverse β keto sulfones via photoredox-catalyzed sulfonylation of diaryliodonium salts with DABSO and silyl enolates involving the insertion of $SO₂$ has been established. In this reaction, bluelight-induced reduction of diaryliodonium salts afford the aryl radical, which would be trapped by $SO₂$ to form sulfonyl radical as the key intermediate. This novel photoredox catalysis with the assistance of visible light represents a green and sustainable approach for the synthesis of β -keto sulfones and features wide substrates scope, good reactivity, and broad functional group tolerance. In addition, by employing b-alkyl substituted silyl enolates, this three-component sulfonylation would give α -alkyl substituted β -keto sulfones, which cannot be accessed by previous method involving the insertion of $SO₂$.

Conflicts of interest

There are no conflicts to declare.

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