

# Green Chemistry

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#### **Green Chemistry**

Kai Yang,<sup>a</sup> Miaolin Ke,<sup>b</sup> Yuanguang Lin<sup>b</sup> and Qiuling Song<sup>\*</sup>,<sup>a</sup>

**Ambient Temperature** 

**Sulfonamides Formation from Sodium Sulfinates and** 

Amines or Ammonia under Metal-Free Conditions at

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### COMMUNICATION

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 $RSO_{2}Na + NHR^{1}R^{2} \xrightarrow{l_{2} (1 \text{ equiv})}_{EtOH \text{ or water, air}} R \xrightarrow{O}_{O} \xrightarrow{R^{1}}_{R^{2}} R^{1}$   $R, R^{1}, R^{2} = H \text{ or aliphatic} \text{ rt} \qquad rt$   $R, R^{1}, R^{2} = H \text{ or aliphatic} \text{ software or EtOH as solvent} R \xrightarrow{O}_{O} \xrightarrow{R^{1}}_{R^{2}} R^{2}$ 

very broad substrate tolerability

A novel, practical and highly efficient method for the construction of a variety of sulfonamides mediated by  $I_2$  was demonstrated. The reaction proceeds readily at room temperature using a variety of sodium sulfinates and amines or ammonia in water in a metal-, base-, ligand-, or additive-free protocol. Primary, secondary and tertiary sulfonamides were obtained in good to excellent yields with a broad range of functional group tolerability.

Sulfonamides are a common structural motif in biologically active compounds and pharmaceutical interesting molecules, owing to their well-known anticonvulsant, antibacterial, anticancer, antitumor, anti-inflammatory and HIV protease inhibitory activities (Figure 1).<sup>[1-7]</sup> In addition, sulfonamides are a good type amino protection group due to their easily removal property.<sup>[8,9]</sup> Consequently, many endeavors have been made towards the construction of sulfonamides. Traditional method for the formation of sulfonamides stems from sulfonyl chloride and amino compounds,<sup>[10,11]</sup> later on transition metal catalyzed alternative method was developed between primary sulfomides and aryl halides<sup>[12-17]</sup> or arylboronic acids.<sup>[18]</sup> Very recently, Jiang and coworkers developed an efficient method to build up sulfonamides between sodium sulfinates and amines via a copper-catalyzed aerobic oxidative conditions.<sup>[19]</sup> Kim et. al. reported a mild copper catalyzed Chan-Lam type coupling by using sulfonyl azide and boronic acids at room temperature.<sup>[20]</sup> Although great progress has been achieved on the synthesis of sulfonamides, there are still many drawbacks existing in the current methods, such as hazardous starting materials (mutagenic sulfonyl chlorides or precautioushandling organic azides), harsh reaction conditions, long reaction

time, poor functional group tolerability and transition metal usage which might cause pollution in the final product and so on. Therefore development of a general, practical and efficient method for the construction of sulfonamides under mild conditions is still a challenge and highly desirable.



Figure 1. Drugs with sulfonamide structure motif

Herein, we report a transition-metal-free sulfonamides formation between sodium sulfinates and amines mediated by I2 at room temperature. There are several obvious advances of the present method: 1) Cheap and readily available I<sub>2</sub> emerges as an efficient catalyst and oxidant as well, instead of a transition-metal catalyst, which is usually quite expensive and is required to be completely removed from products, particularly during the process of drug synthesis; 2) The reaction shows broad substrate scopes and generality: various sodium sulfinates, including aromatic, heteroaromatic or aliphatic ones and numerous amines, such as aromatic, heteroaromatic or aliphatic, natural occurred amino ester, or even aqueous ammomia, are well tolerable under the standard conditions and the desired sulfonamide compounds were obtained at room temperature in open air smoothly within 3 h. Therefore we have reasons to believe that pharmaceutical molecules and bioactive compounds could be manipulated by this strategy; 3) The I<sub>2</sub>/EtOH or water system is low cost, easily handled and mild with high efficiency and safety.

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In our previous research, we found that  $I_2$  could functionalize as a surrogate for CuI in the process of acylation of benzothiazoles with acetophenones.<sup>[21]</sup> This reaction inspired us, since sulfonamide could be formed under a copper-catalyzed aerobic oxidative conditions between sodium sulfinates and amines, we propose that  $I_2$  should replace copper salt, yet just play the same role to the reaction between sodium sulfinates and amines, thus making the reaction take place under transition-metal free conditions.



#### Scheme 1. Overview of the synthetic approach for sulfonamides.

In order to verify the hypothesis, aniline (1a) and sodium benzenesulfinate (2a) were chosen as the substrates in the model reaction (Table 1). To our delight, the desired product 3aa was formed in 60% yield (Table 1, entry 4) with 1 equivalent of  $I_2$  in CH<sub>3</sub>CN just at room temperature in an open flask! This reaction condition was much milder than the copper-catalyzed one. Further oxidant screening revealed that I<sub>2</sub> was the superior oxidant to NBS, NCS, NIS, K<sub>2</sub>S<sub>2</sub>O<sub>8</sub>, DDQ and DTBP (Table 1, entries 1-4, 6-8), also better than catalytic amount of I2 with one equivalent of TBHP (70% in water) (Table 1, entry 5). Solvent effect was also investigated (Table 1, entries 8-14), and both water and EtOH showed optimal influence on the reaction and afforded desired products in 73% and 75% yields respectively (Table 1, entries 9 and 14), indicating this reaction was environmentally benign. Gratifying, shortening the reaction time from 18 hours to 3 hours did not reduce the efficiency of the reaction, and 76% of desired product was obtained correspondingly (Table 1, entry 15). Yet the decreasing on the amount of I<sub>2</sub> to 0.2 equivalents did cause the decreasing on the yield of desired product from 76% to 23% (Table 1, entries 16 and 17). Notably, catalytic amount of I<sub>2</sub> (20 mol%) with TBHP (70% in water) did not bring satisfactory yield, only 19% of desired product was obtained after 18 hours reaction. Based on the above results, EtOH was the optimal solvent for this reaction and water could be an alternative one for this transformation. It is well known, both EtOH and water are environmentally benign solvents and there are many advantages with EtOH and water as solvent over other organic ones. The property makes the whole process quite green and sustainable. Together with mild reaction conditions, the process might be used in late-stage manipulation of complex molecules.

#### Table 1. Condition Screening.

	NH <sub>2</sub> +	`ONa solver	ant nt, rt	O S O H
	1a 2	a		3aa
Entry	Additive (equiv.)	Solvent (mL)	Time (h)	Yield (%) <sup>c</sup>
1	NBS (1)	CH₃CN	18	21
2	NCS (1)	CH₃CN	18	37
3	NIS (1)	CH₃CN	18	12
4	I <sub>2</sub> (1)	CH₃CN	18	60
5	I <sub>2</sub> (0.1) + TBHP (1)	CH₃CN	18	8
6	K <sub>2</sub> S <sub>2</sub> O <sub>8</sub> (1)	CH₃CN	18	trace
7	DDQ (1)	CH₃CN	18	trace
8	DTBP (1)	CH₃CN	18	0
9	I <sub>2</sub> (1)	DCM	18	69
10	l <sub>2</sub> (1)	H₂O	18	73 <sup>d</sup>
11	I <sub>2</sub> (1)	THF	18	49
12	<b>I</b> <sub>2</sub> (1)	dioxane	18	27
13	I <sub>2</sub> (1)	DCE	18	68
14	I <sub>2</sub> (1)	toluene	18	59
15	I <sub>2</sub> (1)	EtOH	18	75 <sup>d</sup>
16 <sup>6</sup>	l <sub>2</sub> (1)	EtOH	3	76 <sup>d</sup>
17 <sup>b</sup>	l <sub>2</sub> (0.2)	EtOH	3	23
18	I <sub>2</sub> (0.2) + TBHP (2)	EtOH	18	19

Reaction Condition: *a.* aniline (**1a**) (0.5 mmol), sodium benzenesulfinate (**2a**) (1 mmol), oxidant (0.5 mmol), solvent (2 mL), air, rt. *b.*  $l_2$  and sodium benzenesulfinate (**2a**) were mixed up and stired at rt for 20 min, then EtOH (2 mL) and aniline (**1a**) were added, the mixture was continued to stir at rt for 3 h. *c.* GC yields. *d.* Isolated yields.

With the optimized reaction conditions in hand, the substrate scope of amines were explored firstly (Table 2). As listed in Table 2, both aromatic amines, such as aniline derivatives and aliphatic amines worked well under the mild reaction conditions. The electronic property of the substituents on the aromatic rings of anilines had some effect on yields, with electron-deficient substitutions usually giving lower yields of the sulfonamides in comparison to electron-rich substitutions (Table 2, 3ba-3ja), perhaps due to a result of lower nucleophilic property of the electrondeficient ones. Secondary aniline, such as N-methylaniline also reacted well with sodium benzenesulfinate and gave the corresponding sulfonamide 3ka, albeit with slightly lower yield. All aliphatic amines showed good reactivity on this novel transformation. Notably, both terminal alkyne and alkene functionalities were tolerable in this reaction (Table 2, 3ra and 3sa) and provide the feasibility for further structural manipulation. Interestingly, a bis-sulfonamide 3ta was obtained in 66% yield with a diamine as substrate. A special aromatic secondary amine, benzoimidazoles, which reacted with a variety of sulfonamide, was also applicable in the standard reaction condition to give the desired products in moderate to good yields (Table 2, 3ua-3ud, 3va). Remarkably, the natural occurred amino esters, which are bioactive molecules, were also proven to be compatible with this reaction and generated the corresponding products in reasonable yields (Table 2, 3wa-3xa), which showed our methods potential application in late stage modification of complex molecule synthesis.

## Table 2. I2 mediated synthesis of sulfonamides from amines 1 and sodium sulfinates (2).

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a. Reaction Condition:  $l_2$  (0.5 mmol) and sodium sulfinete 2 (1.0 mmol) were mixed up and stired at rt for 20 min, then EtOH (2 mt) and amine 1 (0.5 mmol) were added, the mixture was continued to stirt at rt for 3 h. b. EteVH (2 equiv.) was added due to the starting materials amino ester exist in H2 salt.

The scope of sodium sulfinates was investigated by reacting with morpholine (1q) under the standard conditions (Table 3). Both electron-rich and electron-deficient substituents on the aromatic ring of sodium benzenesulfinates were tolerant under the novel transformation (Table 3, 3qb-3ql), yet the latter ones usually showed lower efficiencies than the former ones in term of the isolated yields, which might be explained by the electron-inductive effect. Different position of substituents on the aromatic ring of sodium benzenesulfinates do have effect on the efficiency of the reaction, with the ortho position having lower yield compared with the meta and the para position, which might be explained by the steric hindrance of the *ortho* position. In Table 3, these phenomena were clearly shown by compounds **3qj**, **3qk** and **3ql**. 2-Naphthyl sodium sulfinate was also a good reactant for this reaction (Table 3, 3qm). Intriguingly, quinolinyl group was survived in the standard conditions and gave the corresponding product 3qn in 60% yield. Gratifyingly, this reaction was applicable to aliphatic sodium sulfinates as well, and cyclopropyl sodium sulfinate was a good candidate for this novel metal-free reaction with 81% isolated yield (Table 3, 3qo).

Table 3.  $I_2$  mediated synthesis of sulfonamides from morpholine (1q) and sodium sulfinates 2.



Reaction Condition:  $I_2$  (0.5 mmol) and sodium sulfinate 2 (1.0 mmol) were mixed up and stired at rt for 20 min, then EtOH (2 ml) and morpholine (1q) (0.5 mmol) were added, the mixture was continued to stir at rt for 3 h.

Intrigued by the above results, we further applied the same conditions to ammonia in water. Delightfully, primary benzenesulfonamide was obtained in 75% yield when sodium benzenesulfinate was interacted with aqueous ammonia under the standard conditions (Table 4, 4a). Without optimization, numerous primary sulfonamides were found to be formed through various sodium sulfinates with ammonia in water in moderate to good yields under the standard conditions (Table 4), with the similar substrate scope as the examples list in Table 3.

Primary sulfonamide was a type of very useful building blocks which could be used as starting materials in the cross coupling reaction for polysubstituted sulfonamide synthesis as well as key scaffolds in many bioactive molecules and pharmaceutical compounds. Additionally, this reaction showed a very wide range of functional groups tolerability and all primary, secondary and tertiary sulfonamides could be generated under the standard conditions smoothly. Therefore this is a universal method for the formation of sulfonamides in a transition-metal-free, ligand-, base- and additivefree conditions at ambient temperature.

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Reaction Condition: a.  $I_2$  (0.5 mmol) and sodium sulfinate 2 (0.5 mmol) were mixed up and stired at rt for 20 min, then ammonia (1 mL) was added, the mixture was continued to stir at rt for 3 h. b.  $I_2$  (0.5 mmol) and sodium sulfinate 2 (0.5 mmol) were mixed up and stired at rt for 20 min, then ammonia (1 mL) and EtOH (1 mL) were added, the mixture was continued to stir at rt for 3 h.

To our delight, this reaction could be easily scaled up to 10 mmol without dramatically losing the efficiency. As shown below, morpholine (1a) and sodium benzenesulfinate (2a) were interacted with each other under the standard conditions to generate desired product 3ga in 70% yield (1.6 gram) (Scheme 2), which might suggest a potential application in industry.



Scheme 2. Scale up reaction with morpholine (1q) and sodium benzenesulfinate (2a).

Control experiments were performed in order to understand and gain insight into the mechanism (Scheme 3). A radical scavenger BHT (2,6-di-tert-butyl-4-methylphenol) was added to the reaction system, 65% of desired product **3aa** was obtained (Scheme 3, eq. 1), almost no significant influence on the reaction. Furthermore, no radical intermediate was trapped by radical inhibitor TEMPO (2,2,6,6-tetramethylpiperidine-N-oxyl). These two reactions suggested that a radical reaction might not be involved in the reaction. N-iodomorpholine (1q') and 4-MeObenzene sulfinyl iodide (2b') were prepared according to the literature,<sup>[22-23]</sup> and they were used as the substrates with 4-MeObenzene sodium sulfinate and morpholine correspondingly under the standard conditions, remarkably, the desired product was obtained in 85% and 53% respectively. These two reactions suggested that 1q' and 2b' might be the key intermediates for this novel and sustainable reactions.



#### Scheme 3. Control experiment

On the basis of the above results, two tentative reaction mechanisms for this I2-mediated sulfonamide formation were given in Scheme 4: in **path a**, amine might be activated in the presence of iodine to give N-iodoamine 5, which reacted with sodium benzenesulfinate to give the sulfonamide with the release of HI; or in **path b**, sodium benzenesulfinate reacted firstly with  $I_2$  to give benzenesulfonyl iodide 6 which further react with amine to afford the desired product sulfonamide.



Scheme 4. Plausible reaction mechanism for sulfonamide formation

#### Conclusions

In summary, a highly efficient, practical, and chemoselective sulfonamides synthesis have been developed. All types of sulfonamides, such as primary, secondary and tertiary sulfonamides could be generated in good to excellent yields. Wide substrate scope compatibility and mild reaction conditions plus transition-metal-, base-, ligand- and additive-free features recommends that this method might be applied for the synthesis of bioactive compounds and medically important compounds, also could be used in late stage structural manipulation of complex molecules. This method will widely expand the utility of sodium sulfinates in organic synthesis.

#### Notes and references

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Electronic Supplementary Information (ESI) available: <sup>1</sup>H NMR and <sup>13</sup>C NMR analysis for all products . See DOI: 10.1039/c000000x/

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## Sulfonamide Formation from Sodium Sulfinates and Amines or Ammonia under Metal-Free Conditions at Ambient Temperature

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#### Abstract



A practical and highly efficient method for the construction of a variety of sulfonamides mediated by  $I_2$  was demonstrated. The reaction proceeds readily at room temperature using a variety of sodium sulfinates and amines or ammonia in water in a metal-, base-, ligand-, or additive-free protocol. Primary, secondary and tertiary sulfonamides were obtained in good to excellent yields with a broad range of substrates tolerability.