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PII:	S1001-8417(19)30755-7		
DOI:	https://doi.org/10.1016/j.cclet.2019.12.040		
Reference:	CCLET 5398		
To appear in:	Chinese Chemical Letters		
Received Date:	27 November 2019		
Revised Date:	25 December 2019		
Accepted Date:	31 December 2019		

Please cite this article as: Luo D, Min L, Zheng W, Shan L, Wang X, Hu Y, Introduction of N,N'-disulfonylhydrazines as new sulfonylating reagents for highly efficient synthesis of (E)- β -iodovinyl arenesulfones under mild conditions, *Chinese Chemical Letters* (2020), doi: https://doi.org/10.1016/j.cclet.2019.12.040

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Chinese Chemical Letters

journal homepage: www.elsevier.com

Communication

Introduction of *N*,*N*'-disulfonylhydrazines as new sulfonylating reagents for highly efficient synthesis of (E)- β -iodovinyl arenesulfones under mild conditions

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Article history: Received 27 November 2019. Received in revised form 25 December 2019, Accepted 27 December 2019

Available online

Graphical Abstract



N,*N*'-Disulfonylhydrazines were introduced as new sulfonylating reagents to achieve a highly efficient synthesis of (E)- β -iodovinyl arenesulfones under extremely mild conditions.

ARTICLE INFO

ABSTRACT

N,N'-Disulfonylhydrazines have been proven to be the most reactive precursors of the sulfonyl radicals among all types of sulfonyl substituted hydrazines as early as half a century ago. However, the sulfonyl radicals generated from these compounds have not been used in organic synthesis except the simple self-dimerization synthesis of disulfones controlled by the "solvent-cage-effects". In this article, N,N'- disulfonylhydrazines were introduced as new sulfonylating reagents and their combinations with NIS were disclosed as new iodosulfonylating reagents of alkynes. Finally, a highly efficient method for the synthesis of (E)- β -iodovinyl arenesulfones was developed by mixing an alkyne, a N,N'-disulfonyl- hydrazine and NIS in aqueous THF at room temperature for 5 min.

Keywords: Alkynes: Radical reactions; N,N'-Disulfonylhydrazines; Iodosulfonylation; Sulfonyl radicals

Vinyl sulfones are a type of important organic compounds and have been widely applied in organic synthesis and drug discovery [1,2]. (*E*)- β -Iodovinyl sulfones are one of structurally special types of vinyl sulfones (Scheme 1). Their iodine atoms are responsible for many unique transformations, such as: (a) dehydroiodination [3], (b) nucleophilic substitution [4], (c) Suzuki coupling [5], (d) Sonogashira coupling [6]. They have been widely used as versatile synthesis of the synthesis of functionalized sulfones.



Scheme 1. (*E*)- β -Iodovinyl sulfones as versatile synthons

In literature, the iodosulfonylation of alkynes has been recognized as the most practical method for the synthesis of (E)- β -iodovinyl sulfones. The first iodosulfonylation was reported by Truce [3e] in 1971, in which RSO₂I (1a) was used as both a sulfonylating reagent and an iodine source under illumination (Scheme 2a). However, this most straightforward procedure lost its appeal soon because RSO₂I (1a) are highly instable compounds for preparation and storage. In the past decades, many modified procedures [2] have been developed to use RSO₂Na (1b) [3d,5a-c,7], RSO₂H (1c) [6b,8], TosMIC (1d) [9], DABCO·(SO₂)₂ (1e) [10] or RSO₂NHNH₂ (1f) [11] as a sulfonylating reagent in the presence of an iodine source and an oxidant (Scheme 2b). Although each of reagents has its own advantages, they all have their own unavoidable drawbacks. For example, RSO₂Na (1b) is the most often used sulfonylating reagent, but this salt usually suffers from the poor solubility in organic solvents. In recent years, RSO₂NHNH₂ (1f) received much more attention due to its satisfactory stability, solubility and preparation, but its reactions usually suffers from the uses of large amounts of peroxide oxidants under prolonged heating conditions.



Scheme 2. Previous iodosulfonylations and this work.

Herein, we report a novel method for the synthesis of (E)- β -iodovinyl arenesulfones (4), in which N,N'-disulfonylhydrazines [(ArSO₂NH)₂, 3] were introduced as new sulfonylating reagents. As shown in Scheme 2c, the desired product 4 was synthesized efficiently by mixing an alkyne (2), a N,N'-disulfonylhydrazine (3) and NIS (N-iodosuccinimide) together in aqueous THF at room temperature within five minutes.

The mechanism studies [3e,5a-c,6b,7-11] indicated that the iodosulfonylations of alkynes similarly went through a radical addition pathway. As shown in Scheme 3, this radical addition was initiated by a sulfonyl radical (RSO₂•) [3e] that was generated *in situ* from a sulfonylating reagent. Therefore, the methodology study for the synthesis of (E)- β -iodovinyl sulfones actually has been became an issue how to conveniently and efficiently generate the sulfonyl radicals.





Thus, we made a comprehensive investigation for different types of sulfonyl substituted hydrazines. We were surprised to find that, as early as 1970 [12], *N,N'*-disulfonylhydrazines [Ar¹SO₂NHNHSO₂Ar² (**3**)] have been proven to be the most convenient and efficient precursors of the sulfonyl radicals. As shown in Scheme 4, when compound **3** was treated with an oxidant (such as Br₂ [12], NBS [12], KMnO₄ [12], Chloranil [12], NaOCl [13] or HNO₃ [14]), it quickly released N₂ *via* an azo-intermediate (**5**) to generate two sulfonyl radicals (**6**). Finally, two sulfonyl radicals were dimerized into a disulfone R¹SO₂-SO₂R² (**7**, Ar¹ = Ar² or Ar¹ \neq Ar²). For example, the disulfone Ts-Ts (**7a**) was produced in 52% yield by treatment of TsNHNHTs (**3a**) [14b] with HNO₃ at room temperature for 1 h. The disulfone PhSO₂-SO₂Ph (**7b**) was obtained in 41% yield when (PhSO₂NH)₂ (**3b**) was treated with NaOCl at 0 °C for 5 min [13]. Remarkably, unsymmetrical disulfone Ts-SO₂Ph (**7c**) was obtained as single product in 67% yield from the oxidation of unsymmetrical TsNHNHSO₂Ph (**3h**) [14b]. Clearly, these disulfons **7** were formed by a self-dimerization controlled strongly by the "solvent-cage-effects" [12].



Scheme 4. N,N'-disulfonylhydrazines (3) as precursors of sulfonyl radicals.

Our investigation also showed that N,N'-disulfonylhydrazines (**3**) have had several important synthetic applications in recent years [15]. In these works, only the reactions of N-atom in the compounds **3** were emphasized. In fact, N,N'-disulfonyl-hydrazines (**3**) have not been used as the precursors of sulfonyl radicals in organic synthesis except the simple self-dimerization synthesis of disulfones (**7**) in nearly half a century. This phenomenon may be caused by the negative influence of the "solvent-cage-effects" on sulfonyl radicals. However, we carefully found that a mixture including the product Ts-Br (**8**) was obtained when **3a** was treated with NBS in aqueous MeCN (Scheme 5) [12]. We strongly believed that the formation of **8** resulted from the fact that a radical Ts• was captured by a radical Br•. This result also indicated that the "solvent-cage-effects" on sulfonyl radicals may be broken by an external radical scavenger.

Ts-NHNH-Ts $\xrightarrow{NBS, aq. MeCN, r.t.}$ Ts-Ts + Ts-Br + Ts-OH 3a 7a (10%) 8 (53%) 9 (25%) Scheme 5. Sulfonyl radicals captured by bromine radicals.

Thus, we hypothesized that (E)- β -iodovinyl arenesulfones (4) may be synthesized from an alkyne (2, as an external radical scavenger) and a $(ArSO_2NH)_2$ (3) in the presence of NIS (as an oxidant and an iodine source). As shown in Table 1, a group of conditional experiments were tested by using PhC=CH (2a) and TsNHNHTs (3a) as model substrates. When the solution of 2a and 3a in anhydrous MeCN was treated with NIS for 2 h, the desired product 4a was obtained in 9% yield (entry 1). However, the N₂ released quickly to give the product 4a in 82% yield within 10 min when the same reaction proceeded in aqueous MeCN (96 vol%) (entry 2). This result indicated that NIS was not strong enough to convert 3a into the corresponding azo-intermediate 5a. The "real oxidant" may be HOI that was formed *in situ* by hydrolysis of NIS in water. None of the product 4a was obtained when toluene was used as a solvent, possibly due to the poor solubility of the substrates (entry 3). THF was the best solvent to give the product 4a in 92% yield within 5 min (entry 7). The results in entries 8 and 9 indicated that the yield of 4a was obtained when the reaction proceeded in water because both of substrates were not soluble in water (entry 10). The yield of 4a was improved slightly by addition of 0.5 equivalents of TBAI as a phase-transfer-catalyst (entry 11).

Next, the different oxidants and iodine sources were tested as the alternatives of NIS (Table 2). Although ICl and I_2 are the oxidative iodine sources, they showed poor reactivity for this reaction (entries 1 and 2). When the combinations of NaOCl/I₂ (entry 3), NaOCl/NaI [16] (entry 4), NCS/I₂ (entry 5) or NCS/NaI [17] (entry 6) were used, the N₂ released completely within 5 min to give the product **4a** in 18%-66% yields. These unsatisfactory results indicated that NaOCl and NCS were good oxidants for the decomposition of TsNHNHTs (**3a**). However, I₂ and NaI were unsuitable iodine sources, possibly because the formation of iodine radicals (I•) from these iodine sources was a slow process. When the mixture of **2a** and **3a** was treated with the combination of I₂/NIS, **4a** was obtained in 90% yield (entry 7).

At this time, we realized that we may find one of the most efficient methods for the synthesis of (E)- β -iodovinyl arenesulfones (4). To the best of our knowledge, no such method has been reported in literature so far. Thus, the scope of this method was tested under standard conditions (Scheme 6). By fixing TsNHNHTs (3a), different products 4a-4k were obtained in high to excellent yields from aryl ethynes 2a-2k. The product 4l was obtained smoothly in 70% yield from pyridin-2-yl ethyne (2l). Although the product 4m was obtained in 36% yield from cyclohexyl ethyne (2m), only complicated mixtures were obtained from other alkyl ethynes. By fixing phenyl ethyne (2a), all tested disulfonylhydrazines 3a-3g proved to be highly efficient sulfonylating reagents. The desired products 4n-4s were obtained in excellent yields and no comparable electronic effects were observed.



Scheme 6. The scopes of substrates and products.

The *N*,*N*'-disulfonylhydrazines (**3**) proved to be so such strong sulfonylating reagents that they reacted smoothly with internal alkynes (**2n-2q**) to give the tetra-substituted (*E*)- β -iodovinyl sulfones **4t-4z**. Under the influences of the steric and/or electronic effects of the substituents on the internal alkynes, the products **4t** and **4u** were obtained in relatively low yields. Usually, the hydroxyl group could not survive under the conditions of the existing methods. However, the products **4w-4z** bearing the versatile hydroxymethyl groups were synthesized in moderate yields extremely short reaction times and mild conditions were used in our method. Since the products **4w-4z** are new compounds, the structure of **4z** was determined by X-ray single crystal diffraction and its (*E*)-configuration was confirmed. Under the standard conditions, the product **4a** was prepared on 2-gram scales in 90% yield. However, *N*,*N*'-dialkyl sulfonyl-hydrazines have been proved to be unsuitable sulfonylating reagents for our methods.

To further understand the mechanism of this method, more conditional experiments were tested under the standard conditions. As shown in Schemes 7a and 7b, no differences were observed when the reaction proceeded in the dark or under O_2 atmosphere. None of the product **4a** was detected in the presence of one equiv of TEMPO (Scheme 7c). These results indicated that our method may go through a radical addition pathway and the tosyl radical (Ts•) is generated by an oxidative decomposition of **3a**. As was expected, two products **4a** and **4n** were obtained when unsymmetrical *N*,*N*'-disulfonylhydrazine **3h** was used as a substrate (Scheme 7d).





Under the standard conditions, some of the existing sulfonylating reagents were tested as shown in Schemes 8a-8e. The results clearly indicated that all these existing sulfonylating reagents were not suitable for our method.



Scheme 8. The tests for different sulfonylating reagents.

Thus, a possible pathway for our method was proposed as shown in Scheme 9. Initially, NIS was hydrolyzed by H_2O to form the stronger oxidant HOI, by which *N*,*N*'-disulfonyl-hydrazine (3) was oxidized into the corresponding azo-intermediate (5). The azo-intermediate (5) was then spontaneously decomposed into two sulfonyl radicals (6). When the sulfonyl radical (6) was captured by an alkyne (2), a vinyl radical (10) was formed. Finally, the vinyl radical (10) attacked NIS to get an iodine radical to yield the target product (4). In this pathway, NIS served as the precursors for both oxidant HOI and iodine radical (1·). Since no additional peroxide oxidant needed, our method was performed under extremely convenient and safe conditions.



Scheme 9. A proposed pathway for the formation of 4.

In conclusion, a new method for highly efficient synthesis of (E)- β -iodovinyl arenesulfones was developed by simply mixing an alkyne, a *N*,*N*'-disulfonylhydrazine and NIS together in aqueous THF for 5 min at room temperature. Its success is mainly due to two distinctive works: (a) *N*,*N*'-disulfonyl-hydrazines were introduced as highly efficient sulfonylating reagents; (b) the combinations of *N*,*N*'disulfonylhydrazines/NIS were discovered as highly efficient iodosulfonylating reagent of alkynes.

Declaration of interests

 \boxtimes The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgments

This work was supported by National Natural Science Foundation of China (Nos. 21971138 and 21472107).

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Table 1

Effects of the solvents, H₂O and reaction times.^a

_			NIS, <mark>H₂O</mark> I	Ts		
Ph— + 2a		⁺ isnhnhts – 3a	solvent, r.t. Ph 4a 0-92%			
	Entry	Solvent	$H_2O(vol\%)$	Time (min)	4a $(\%)^b$	
	1	MeCN		120	9	
	2	MeCN	4 ^c	10	82	
	3	PhMe	4	120	0	
	4	1,4-dioxane	4	10	45	
	5	DCM	4	10	60	
	6	DME	4	10	88	
	7^d	THF	4	5	92	
	8^e	THF	4	5	59	
	9 ^f	THF	4	5	90	
	10	THF	100	5	7	
	11^{g}	THF	100	5	35	

^a The mixture of **2a** (0.5 mmol) and **3a** (0.5 mmol) in a solvent (3 mL) was treated by NIS (1 mmol) at room temperature for given times. ^b The isolated yields.

 c H₂O (4 vol%) was used in the reference 13.

^d Similarly excellent results were obtained by uses of 3-10 vol% of water.

^e0.25 mmol of **3a** was used.

 $^{\rm f}$ 0.4 mmol of **3a** was used.

^g 0.5 equiv. of TBAI was used.

Table 2

Effects of the oxidants and iodine sources.^a

Ph + 2a	TsNHNHTs 3a	oxidant, [l]-s	., 2 h Ph 4a 10%-90%	
Entry	Oxidant		[I]-source	4a $(\%)^b$
1	ICl		ICl	31
2	I_2		I_2	10
3	NaOCl		I_2	25
4	NaOCl		NaI	60
5	NCS		I_2	18
6	NCS		NaI	66
7 ^c	I_2		NIS	90

^a The mixture of **2a** (0.5 mmol) and **3a** (0.5 mmol) in aq. THF (3 mL) was treated by an oxidant (1 mmol) in the presence of an iodine source (0.5 mmol) at room temperature for 2 h.

^b The isolated yields.

^c The reaction was finished within 5 min.