

Scheme 2. Iodine-copper exchange with 2-iodo-3-methyl-2-cyclohexenone.

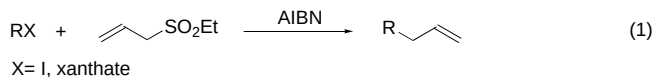
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Tin-Free Radical-Mediated C–C-Bond Formations with Alkyl Allyl Sulfones as Radical Precursors**

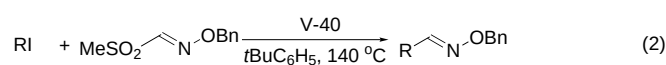
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The synthetic importance of tin-free radical reactions has been well recognized in recent years.^[1] Among several

approaches, an organosulfone-mediated approach is very effective for allylation,^[2] vinylation,^[3] and azidation^[4] [Eq. (1), AIBN = 2,2'-azobisisobutyronitrile]. However, the reported methods did not work well with primary alkyl iodides and xanthates owing to inefficient iodine-atom transfer and xanthate-group transfer, respectively. Recently, we also reported a tin-free acylation approach

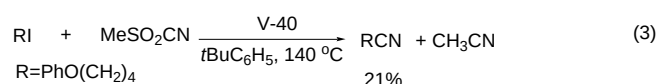


using methanesulfonyl oxime ether, in which primary alkyl iodides and xanthates caused the same problem as a result of the small energy difference between the methyl radical and the primary alkyl radical [Eq. (2), V-40 = 1,1'-azobis(cyclohexane-1-carbonitrile)].^[5]



As our extensive efforts to generate primary alkyl radicals from primary alkyl iodides and xanthates were unsuccessful, we have been interested in developing a new reliable method to generate primary alkyl radicals by using new types of radical precursors that do not require an atom- or a group-transfer step. In this regard, we have studied the possibility of using an alkyl allyl sulfone as a radical precursor. Alkyl allyl sulfones have been widely used as radical acceptors to transfer an allyl group to a radical precursor.^[6] Although alkyl allyl sulfones have been used once as the radical precursor in an allylation reaction,^[2a] primary alkyl allyl sulfones have not been examined. To the best of our knowledge, S-alkoxycarbonyl dithiocarbonates are the only generators of primary alkyl radicals from alcohols, but they cannot be applied to C–C-bond formations owing to the rapid formation of the corresponding xanthates.^[7] We have found that alkyl allyl sulfones are highly efficient and reliable radical precursors for the generation of primary alkyl radicals under tin-free conditions and can be successfully applied to various C–C-bond-formation reactions.

Initially, we focused on radical cyanation,^[8] and began our study with a primary alkyl iodide and methanesulfonyl cyanide.^[9] The reaction of 4-phenoxybutyl iodide with methanesulfonyl cyanide (2 equiv) and V-40 (0.2 equiv) in *tert*-butylbenzene at 140 °C for 5 h afforded 4-phenoxybutyl cyanide in only 21 % yield together with recovered 4-phenoxybutyl iodide (77 %). Notably, methanesulfonyl cyanide was completely consumed, with acetonitrile as the major product [Eq. (3)].^[10] However, allyl sulfone **1** was an effective

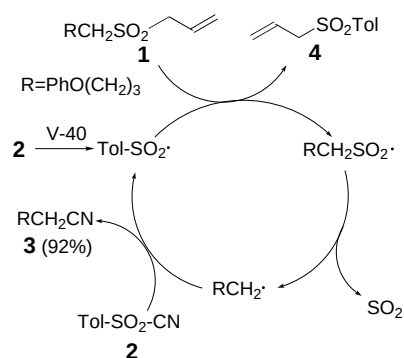


precursor for radical cyanation, and our approach is outlined in Scheme 1. We envisaged that the addition of a *p*-toluenesulfonyl radical to **1** would produce an alkyl sulfonyl radical as well as *p*-tolyl allyl sulfone **4**. Although the alkyl sulfonyl

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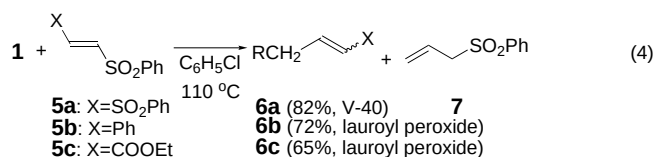
Scheme 1. Pathway for radical cyanation of an alkyl allyl sulfone with *p*-toluenesulfonyl cyanide.

radical could add to **1** and **4**, the former is a degenerate process, and the latter produces the *p*-toluenesulfonyl radical. Thus, neither of the reactions interferes with the desired process. As the addition of an alkyl radical to **4** and **1** is relatively slow,^[11] the alkyl radical that is generated from the thermal decomposition of the alkyl sulfonyl radical, should preferentially add to *p*-toluenesulfonyl cyanide (**2**) and regenerate the *p*-toluenesulfonyl radical for propagation of the radical chain reaction.

The reaction of **1** with **2** (1.5 equiv) in the presence of V-40 initiator (0.2 equiv) in chlorobenzene at 110 °C for 6 h proceeded cleanly, yielding **3** in 92 % yield. The high-yielding formation of **3** indicates that the primary alkyl radical is generated cleanly under tin-free conditions (Scheme 1).^[12] As shown in Table 1, the present method worked well with primary and secondary alkyl allyl sulfones as well as with a benzyl allyl sulfone.

The present approach can be further applied to radical vinylation and allylation reactions. For radical vinylation,^[3,13] we found that (*E*)- and (*Z*)-1,2-bis(phenylsulfonyl)ethylene (**5a**)^[14] are excellent radical acceptors and transfer a vinyl

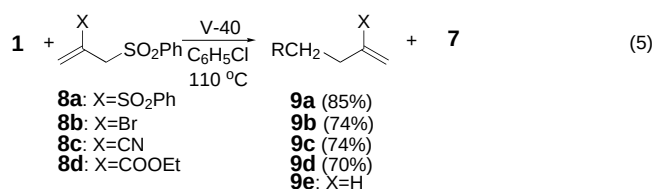
sulfonyl group to an alkyl group.^[15] Treatment of **1** with **5a** and V-40 in chlorobenzene at 110 °C for 6 h afforded **6a** in 82 % yield [Eq. (4)].^[16] Apparently, the addition of the primary



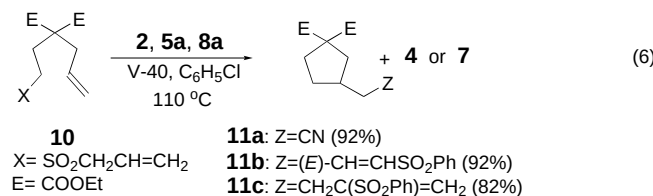
alkyl radical derived from **1** to activated vinyl sulfone **5a** is much faster than the addition to unactivated allyl sulfone **1** and **7**. When (*Z*)-1,2-bis(phenylsulfonyl)ethylene was used, various alkyl allyl sulfones worked equally well, yielding (*E*)-1-phenylsulfonyl alkenes as a major product in high yields (Table 1).^[16]

However, the reaction of **1** with **5b** under similar conditions gave the desired alkene **6b** in 21 % yield. A similar result was also obtained with **5c** (18 %). However, when the reactions were performed with lauroyl peroxide as initiator, much better results were obtained with **5b** and **5c**, showing the generality of vinylation.

We next studied radical allylations with several C2-substituted allyl sulfones [Eq. (5)]. Radical reaction of **1** with **8a**



(1.5 equiv) in chlorobenzene in the presence of V-40 (0.2 equiv) for 6 h afforded 2-phenylsulfonyl alkene **9a** in 85 % yield. Additional experimental results with **8a** clearly demonstrate the synthetic efficiency of the present method (Table 1). When allylations were carried out with other functionalized allyl sulfones (**8b**, **8c**, **8d**) under similar conditions, the corresponding allylated products were isolated in high yields without the formation of **9e**, which results from the concomitant addition of the alkyl radical to **1** and **7**. Finally, when tandem radical reactions involving cyclization and C–C-bond formation were briefly studied with **10**, the reactions worked well, yielding **11** in high yield [Eq. (6)].



In conclusion, we have found that primary alkyl allyl sulfones are one of the most useful and reliable sources of primary alkyl radicals under tin-free conditions and developed highly efficient tin-free radical-mediated cyanation, vinylation, and allylation reactions, which we believe have great synthetic potential.

Table 1. Tin-free cyanation, vinylation and allylation.^[a]

Alkyl allyl sulfone (1) RY (Y = SO ₂ CH ₂ CH=CH ₂)	3 ^[c]	Yields [%] ^[b] 6a ^[d]	9a ^[e]
EtO ₂ C(CH ₂) ₃ Y	93	88	84
	96	94	85
	98	96	90
	98	98	88
		98	98
	98	97	97

[a] The reaction was carried out with V-40 in chlorobenzene at 110 °C for 4–6 h. [b] Yield of isolated products. [c] **3**: Obtained from the reaction of **1** and **2**. [d] (*E*)-**6a**: Obtained from the reaction of **1** and (*Z*)-**5a**. [e] **9a**: Obtained from the reaction of **1** and **8a**.

Experimental Section

Typical procedure: A solution of **1** (0.2 mmol), **2** (0.3 mmol), and V-40 (0.04 mmol) in chlorobenzene (1 mL) was degassed with nitrogen for 10 min and the solution was then heated at 110 °C under nitrogen for 6 h. The solvent was evaporated under reduced pressure and the residue was separated by silica-gel column chromatography (Et₂OAc/*n*-hexane 1:3) to give **3**. ¹H NMR (CDCl₃, 400 MHz): δ = 1.85–1.95 (m, 4H), 2.43 (t, *J* = 7.0 Hz, 2H), 4.00 (t, *J* = 5.6 Hz, 2H), 6.86–6.96 (m, 3H), 7.24–7.29 (m, 2H); ¹³C NMR (CDCl₃, 100 MHz): δ = 17.0, 22.5, 28.2, 66.5, 114.4, 119.5, 120.9, 129.5, 158.6; IR (polymer): $\tilde{\nu}$ = 2927, 1601, 1586, 1497, 1474, 1247, 757, 693 cm⁻¹; HRMS [*M*⁺] calcd for C₁₁H₁₃NO: 175.0997, found: 175.0997.

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