Thioether-functionalized trifluoromethyl-alkynes, 1,3-dienes and allenes: divergent synthesis from reaction of 2-trifluoromethyl-1,3-conjugated enynes with sulfur nucleophiles†

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A divergent synthesis of thioether-functionalized trifluoromethyl-alkynes, 1,3-dienes and allenes via regioselective nucleophilic addition of sulfur nucleophiles to 2-trifluoromethyl-1,3-conjugated enynes was developed. The addition patterns depend on the type of enyne, sulfur nucleophile and reaction conditions used. 1,4-Addition leading to thioether-functionalized trifluoromethyl-allenes was realized when enynes possessing electron-withdrawing aryl groups on the alkyne moiety were used as reaction partners and alkanethiols were used as nucleophiles, whereas solvent-controlled construction of thioether-functionalized 1,3-dienes and allynes was realized, respectively, via a 3,4-addition pattern or 1,2-addition pattern if thiophenols were applied as nucleophiles. The three types of compounds containing both sulfur and fluorine elements are valuable building blocks for synthesis of multifunctional fluorinated vinyl sulfides and thiophene derivatives.

Introduction

Owing to their unique structural properties and reactivity, allene chemistry,1 alkyn chemistry,2 as well as 1,3-diene chemistry,3 have attracted particular attention by chemists in the past few decades. They have proven themselves to be important and valuable building blocks toward complex molecular targets in organic synthesis. As a consequence, substantial efforts have been made over the years by synthetic organic chemists in order to design and develop methods allowing efficient access to a variety of functionalized allenes,4 alkynes5 and 1,3-dienes.6 Meanwhile, the widespread use of fluorinated, especially trifluoromethylated, compounds in pharmaceutical chemistry, agrochemistry7 and materials science8 has attracted considerable interest in development of various synthetic methodologies for synthesis of these trifluoromethylated species.9 In this context, special attention has naturally been brought to trifluoromethyl-allene,10 alkyn11 and 1,3-diene12 building blocks and how to access them.

During the last few years, our research group and others have demonstrated that 2-trifluoromethyl-1,3-conjugated enynes are readily available fluorinated building blocks for the synthesis of fluorinated heterocycles and carbocycles. They could act as four- or two-carbon components in palladium catalyzed intermolecular formal [4 + 2] or [3 + 2] cycloaddition reactions for trifluoromethyl benzenes or exomethylene cyclopentane derivatives synthesis (Scheme 1a).13 They also could serve as novel electrophiles to react with bisnucleophiles such as hydroxylamine, primary amine, aminomalonate and 1,3-dicarbonyl compounds for various fluorinated heterocycles synthesis such as pyrrrole, N-hydroxypyrrrole, isoxazole, cyclic nitroene, pyrroline, pyrrolidine and fluorinated carbocycles synthesis such as cyclopentene derivatives, etc. in transition metal catalysed or simple base mediated cascade reactions (Scheme 1b).14

Sulfur-containing compounds (SCCs) have important applications in pharmaceuticals,15 materials,16 and foods.17 Within this context, we became interested in the regioselective nucleophilic addition of sulfur nucleophiles to 2-trifluoromethyl-1,3-conjugated enynes to construct useful building blocks containing both sulfur and fluorine elements, which are both leading constituents of the pharmaceuticals that comprise our medicinal history.18 We herein report our investigation toward this aim and found that 1,4-addition leading to thioether-functionalized trifluoromethyl-allenes was realized when enynes possessing electron-withdrawing aryl groups on the

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alkyne moiety were used as reaction partner and alkanethiols were used as nucleophiles, whereas solvent-controlled construction of functionalized 1,3-dienes and alkynes were realized, respectively, via 3,4-addition pattern or 1,2-addition pattern if thiophenols were applied as nucleophiles (Scheme 1c).

**Results and discussion**

We initiated this project using enyne 1a and 4-methylthiophenol 2a as model substrates for screening reaction conditions. Under base free conditions, no reaction occurred, the result prompted us to test base as additives. Considering that the replacement of the alkaline metal ion with ammonium cation formed by deprotonation of hydrosulfuryl of thiophenol will weaken the metalfluorine interaction, thus suppress the defluorination process,\textsuperscript{18,19} we choose organic base such as tertiary amine for reaction screening (for detailed reaction conditions screening, please see ESI Table 1). During our initial screens, we found that two main regioselective nucleophilic adducts, that is, functionalized alkyne 3aa via 1,2-addition pattern and functionalized 1,3-diene 4aa via 3,4-addition pattern were formed, which relied heavily on the solvent and base used. In order to control regioselectivity and to have the optimized reaction conditions for the formation of 3aa and 4aa, respectively, extensive optimizations were performed. After many attempts, we were pleased to find that the combination of Et₃N as base and ether as solvent was found to be the optimal reaction conditions for 1,2-nucleophilic addition pattern, and functionalized alkyne 3aa could be obtained in 77% isolated yield. Interestingly, simply use DMSO instead of ether as solvent invert the regioselectivity to 3,4-nucleophilic addition pattern, and functionalized 1,3-diene 4aa could be obtained in 80% isolated yield. We hypothesize that thiophenol has a relatively high acidity and is easily ionized with increasing nucleophilicity in a polar solvent with relatively high dielectric constant such as DMSO compared with Et₂O under basic reaction conditions,\textsuperscript{20} which facilitate both allene intermediate formation and subsequent stereospecific nucleophilic addition to allene and β-heteroatom elimination tandem process, leading to functionalized 1,3-diene formation.

With the optimal reaction conditions established, the enyne substrate scope as well as thiophenol substrate scope were then examined and demonstrated with sixteen examples for these two regioselective nucleophilic addition patterns, respectively, as shown in Scheme 2. There are several points that are noteworthy: (1) in most cases, the reactions are solvent-controlled regioselectivity, functionalized alkynes 3 and 1,3-dienes 4 are two predominate adducts while the amount of the allene 5 is negligible, which accounts for the mass balance; (2) in one case for synthesis of functionalized alkynes 3ha, enyne 1h bearing strong electron-attracting group such as nitro group in para-position of the benzene ring on the alkyne moiety exhibits substrate-controlled regioselectivity and allene 5ha is formed predominately (61% isolated yield); (3) enynes 1e, 1f bearing relatively weakly electron-attracting group such as acyl, nitrile group afford functionalized alkynes 3ea, 3fa in 22%, 33% isolated yield along with the formation of 1,3-butadiene 4ea and 4fa in 60%, 52% isolated yield, respectively; (4) in most cases, both enyne and thiophenol show good functional groups tolerance for 3,4-nucleophilic addition pattern, and good to excellent yields of the 1,3-butadiene products could be obtained with high stereoselectivity, most of addition products were observed as single double bond E-isomers (4aa–4da, 4ia–4ja, 4ab–4ag), and (5) the structure of compound 4ag was further confirmed by means of single-crystal X-ray crystallography (Fig. 1).\textsuperscript{22} (6) owing to its insolubility in ether, 4-nitrobenzenethiol 2g failed to afford corresponding alkyne 3ag in ether, however, it reacts well with enyne 1a in DMSO to afford corresponding 1,3-diene 4ag in 84% isolated yield.
Next, we evaluated the use of alkanethiol instead of thio-phenol as nucleophile. Initially, benzyl mercaptan was selected as a representative alkanethiol reagent to react with enyne under above two optimal reaction conditions. It is interesting to note that regioselective nucleophilic addition pattern varied from 1,2-/3,4-addition pattern to 1,2-/1,4-addition pattern and functionalized alkynes and allene were two predominately formed adducts according to crude 1H NMR analysis, while the amount of the 1,3-diene was negligible [eqn (1)]. The interesting results prompted us to further explore optimal reaction conditions for allene synthesis. Based on the formation of allene and our previously published paper for cyclopentene synthesis, concerning the significant role of electronic effects in allene formation, enyne with an ester substituent was selected as model substrate to react with benzyl mercaptan for optimal reaction conditions screening. After numerous attempts, we were pleased to find that allene could be obtained in 75% isolated yield when reaction was performed in the co-solvent of DMSO and CHCl₃ (v/v = 1 : 1) at room temperature by using 10% TMEDA as catalytic base (TMEDA = N,N,N′,N′-tetramethyl-ethane-1,2-diamine) (see ESI Table 2†).

With these results in hand, the enyne substrate scope as well as alkanethiol scope were then carried out (Scheme 3). Points to note: (1) in general, enynes possessing electron-withdrawing aryl group on the alkyne moiety are general good candidates for allene formation and the desired thioether-functionalized allenes could be obtained in moderate to good yield (5gh–5ht); (2) although 1,4-addition pattern proceeded well with alkanethiols and enyne bearing electron-withdrawing groups, product mixture were generally observed and functionalized alkynes were predominated by-products; (3) a variety of...
substituted benzyl thiols, including hindered substrate underwent 1,4-addition smoothly and corresponding functionalized alkenes could be obtained in moderate to good yield \((5\text{hi}-5\text{hp})\); (4) both primary and secondary alkanethiol \((5\text{hs})\) as well as tertiary alkanethiol \((5\text{hs})\) are suitable nucleophiles. Using alkanethiols as nucleophiles, 3,4-nucleophilic addition pattern leading to functionalized 1,3-diene was suppressed. We hypothesize that this may be due to its relatively low acidity and hardly ionization in DMSO compared with thiophenols, which reduces its ability to nucleophilic attack to allene intermediate to form functionalized 1,3-diene.²¹

To highlight the synthetic utilities of present transformation, several selective transformations of the representative functionalized alkyne \(3\text{aa}, 1,3\)-dien 4aa and alkenes 5eh, 5hh are shown in Scheme 4. The results show that they are versatile organic building blocks in organic synthesis. For example, upon treatment of 3aa with molecule iodine, a I⁻ induced electrophile cyclization and subsequent \(I^+\) induced ring-opening reaction occurred and diiodic compound 6 featured with alkynyl and alkyl iodide was obtained in 50% isolated yield, which was further confirmed by an X-ray crystallography analysis.²² Interestingly, synthetically valuable \(\beta\)-halo alkynyl sulfides 7 or 8 could be delivered in quantitative yield but with poor stereoselectivity by the treatment of 3aa with \(N\)-halosuccinimides (NXS). As we know, vinyl sulfides have wide applications in organic synthesis, material science and pharmaceutics, thus, novel and efficient methods for constructing them are still of important significance to the synthetic community and have received continuous attention.²³ Notably, functionalized 1,3-diene 4aa can also serve as novel electrophiles for the nucleophilic addition. 1,3-Diene 4aa can react readily with benzylamine to afford multifunctional tetrasubstituted alkenes 10 in 86% isolated yield via a consecutive regioselective nucleophilic addition followed by double bond isomerization. Furthermore, 1,3-diene 4aa could undergo Diels–Alder reaction with \(N\)-phenyl-maleimide in chlorobenzene at 150 °C, yielding the corresponding cycloaddition product 9 in 62% isolated yield with moderate diastereoselectivity. It is also important to highlight that halogenated trifluoromethylated thiophenes 11–13 could be delivered in good isolated yield upon treatment of allene 5eh, 5hh with three equivalents of \(N\)-halosuccinimide (NXS) in toluene at room temperature via sequential electrophilic cyclization/oxidation/debenzylation cascade process.²⁴

**Conclusions**

In summary, we have discovered the divergent synthesis of thioether-functionalized trifluoromethyl-alkynes, 1,3-dienes and alkenes from the regioselective nucleophilic addition reactions of 2-trifluoromethyl-1,3-conjugated enynes with sulfur nucleophiles. The addition patterns depend on the type of enynes, sulphur nucleophiles and reaction conditions used. The three types of compounds containing both sulfur and fluorine elements are valuable building blocks for synthesis of multifunctional fluorinated vinyl sulfides and thiophenes derivatives.

**Experimental**

**General procedure for the synthesis of thioether-functionalized trifluoromethyl-alkynes 3**

To the solution of 2-trifluoromethyl 1,3-conjugated enynes 1 (1.0 mmol), thiophenol 2 (1.5 mmol) in toluene (3.0 mL) under nitrogen at room temperature was added Et₃N (2.0 mmol), the reaction was stirred at room temperature for 24 h. After it was completely consumed, which was determined by TLC analysis, the solvent was removed under reduced pressure and the crude reaction mixture was purified by flash column chromatography on silica gel (petroleum ether : ethyl acetate = 1:0-5:1) to give the desired 3.

**General procedure for the synthesis of thioether-functionalized trifluoromethyl-1,3-dienes 4**

To the solution of 2-trifluoromethyl 1,3-conjugated enynes 1 (1.0 mmol), thiophenol 2 (1.1 mmol) in DMSO (5.0 mL) under nitrogen at room temperature was added Et₃N (2.0 mmol), the reaction was stirred at room temperature for 24 h. After it was completely consumed, which was determined by TLC analysis, water (15 mL) was added to reaction mixture and extracted with ethyl acetate (3 x 5 mL). Organic layers were combined, washed with brine (4.0 mL) and dried over MgSO₄, after filtration and evaporation, the residue was purified through flash column chromatography on silica gel (petroleum ether : ethyl acetate = 1 : 0–5 : 1) to give the desired 4.

**General procedure for the synthesis of thioether-functionalized trifluoromethyl-allenes 5**

To the solution of benzyl mercaptan 2 (1.0 mmol) in DMSO/CHCl₃ (v/v = 1 : 1, 5.0 mL) under nitrogen at room temperature...

![Scheme 4 Synthetic transformation of alkyne 3aa, 1,3-diene 4aa and alkenes 5eh, 5hh.](https://example.com/scheme4.png)
was added TMEDA (0.1 mmol), then added 2-trifluoromethyl 1,3-conjugated enynes 1 (1.6 mmol), the reaction was stirred at room temperature for 4 h. After 2 was completely consumed, which was determined by TLC analysis, water (2.0 mL) was added and the reaction mixture was extracted with ethyl acetate (3 : 4.0 mL). The combined organic extracts were washed with brine (4.0 mL) and dried over MgSO4. After filtration and evaporation, the residue was purified by flash column chromatography on silica gel (petroleum ether: ethyl acetate = 100 : 0–50 : 1) to give the desired 5.

**Conflicts of interest**

There are no conflicts to declare.

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**Notes and references**


5. For selected reviews on synthesis of functionalized alkyne, see: (a) E. Bernoud, R. Veillard, C. Alayrac and A. C. Gaumont, *Molecules*, 2012, 17, 14573; (b) K. Lauder, A. Toscani, N. Scalacci and D. Castagnolo, *Chem. Rev.*, 2017, 117, 14091.


22 X-Ray data and ORTEP depiction for compounds 4ag (CCDC 1867549) and 6 (CCDC 1860794) please see ESL†.
