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Use of ArSO$_2$SR$_f$ Reagents: An Efficient Tool for the Introduction of SR$_f$ Moieties

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Abstract: In recent years, a renewal of interest was showed to the quest of new synthetic solutions to directly introduce emergent fluorinated groups (SR$_f$) onto molecules. In this context, a new generation of reagents (ArSO$_2$SR$_f$) as efficient sources of SCF$_3$, SCF$_2$H and more generally SR$_f$ groups, was designed. Hence, potent solutions was developed for the synthesis of SR$_f$-containing molecules, compounds of interest for drug and agrochemical research. This review highlights the recent advances made in the synthesis and the use of this new class of reagents, considerably extending the portfolio of tools for the direct introduction of SR$_f$ moieties.

Introduction

Omnipresent in the daily life, fluorinated molecules$^1$ are compounds of interest in both industry and pharmaceutical fields.$^2$ The importance of these compounds might be directly linked to the remarkable properties of the fluorine atom and the fluorinated groups.$^3$ Although key advances have been made over the years, limitations remain. To answer to the synthetic challenges of this very active research field, innovative methodologies and original reagents were developed. In particular, a strong interest was showed by the scientific community towards the SR$_f$ groups. Indeed, merging the properties of sulfur-containing molecules with fluorinated groups, the corresponding functionalized fluorinated groups (SR$_f$) are of high interest. Therefore, several methods appeared to access these highly valuable SR$_f$-containing compounds. Indirect approaches from sulfur-containing molecules and complementary ones based on the direct introduction of the SR$_f$ groups were designed.$^4,5,6$ Regarding the last strategy, a large panel of SR$_f$ moieties are now available ($R_f = \text{CF}_3$, \text{CF}_2\text{H}, \text{CF}_2\text{C}_n\text{F}_{2n+1}$, \text{CF}_2\text{SO}_2\text{Ph}, \text{CF}_2\text{CO}_2\text{R}, \text{CF}_2\text{PO(OEt)}_2$) thanks to the efforts of key players of the field like Billard,$^7$ Goossen,$^8$ Shibata,$^9$ Shen$^{10}$ and our group,$^{11}$ among others. The design of original nucleophilic and electrophilic sources was successfully achieved, opening new avenues and unprecedented transformations.$^5$ However, these reagents suffer from multi-steps synthesis, difficulty to scale up their preparation and the generation of side products, which considerably hampered wide applications due to atom-economy concerns. Aiming at tackling new synthetic challenges, the generation of reagents, namely ArSO$_2$SR$_f$, was designed acting as efficient electrophilic or radical sources of SR$_f$ moieties. In that review, the major recent advances made in that research field will be presented for the direct introduction of the SR$_f$ moiety.

1. Recent advances in the trifluoromethylthiolation reactions using ArSO$_2$SCF$_3$ reagents

Although impressive progress has been made for the synthesis of SCF$_3$-containing molecules, several strategies based on radical pathways opened new avenues in this very active research field and the development of alternative transformations is still highly desirable. In this section, the recent reports dealing with the trifluoromethylthiolation of compounds using different ArSO$_2$SCF$_3$ reagents will be depicted.
Synthesis and applications of ArSO₂SCF₃ reagents

Synthesis of the PhSO₂SCF₃ reagent I. A strong interest was demonstrated towards trifluoromethylthiolated sulfonate derivatives thanks to their antimicrobial activity. These derivatives were generally prepared by reacting the trifluoromethanesulfenyl chloride with zinc aryl sulfinate dihydrates. With this approach, a large variety of RSO₂SCF₃ derivatives was prepared (R = Alkyl and Aryl, 10 examples, Scheme 1) including the PhSO₂SCF₃ reagent I.

A common synthetic route to prepare trifluoromethylthiolated sulfonates relied on the reaction of sodium sulfinate derivatives with an electrophilic trifluoromethylthiolating reagent as independently reported in 2015 by the groups of Shen and Jereb (Scheme 2). Indeed, in 2015, Shen and co-workers reported the synthesis of an electrophilic trifluoromethylthiolating reagent and its use in several reactions with various nucleophiles. Among them, when sodium sulfinites reacted with the trifluorosulfenate 2 as the electrophilic SCF₃ source under acidic conditions, a panel of compounds (I, II, III and 4) were synthesized in high yields (10 examples, up to 97% yield, Scheme 2, Conditions A). The transformation was tolerant with aromatic sodium sulfinites bearing halogens, electron donating and electron withdrawing groups (II, III, 4a-c) as well as heteroaromatic derivatives (4d and 4e). In the study from the Jereb’s group, only two examples were depicted using the Billard-Langlois reagent 3 in the presence of pTSA H₂O (Scheme 2, Conditions B). In particular, the reagent I was obtained in high yields with these approaches (95% (Conditions A) and 84% (Conditions B), respectively).

In 2017, the group of Qiu depicted a sequential tandem reaction to access trifluoromethylthiolated sulfonates (Scheme 3). Starting from simple anilines, a panel of trifluoromethylthiolated sulfonates was prepared in moderate to high yields in the presence of DABCO(SO₂)₂ as the sulphur dioxide source and the Billard-Langlois reagent 3. The use of a Lewis acid such as bismuth(III) chloride was beneficial in this transformation, presumably activating the PhNHSCF₃ reagent. The reaction worked well with both anilines substituted with electron-donating and electron withdrawing groups (III, 6a-f), the latter being the most...
efficient. Note that heteroaromatic trifluoromethylthiolated sulfonates were not synthesized under these reaction conditions, which constituted the main limitation of this method. Alternatively, starting from a sodium sulfinate salt in the presence of the PhNHSCF$_3$ reagent and $p$TsOH, the synthesis of an heteroaryl trifluoromethylthiolated sulfonate at room temperature was depicted.

![Scheme 3. Synthesis of reagent I and analogs from anilines, SO$_2$ source and PhNHSCF$_3$ reagent.](image)

The same year, the group of Qiu and Sheng developed a complementary approach from sulfonyl chloride via a tandem process (Scheme 4). Indeed, sulfonyl chloride derivatives were first converted into the corresponding sulfinates followed by a reaction with the Billard-Langlois reagent 3 to access the trifluoromethylthiolated sulfonates in the presence of $p$TsOH at room temperature. With this approach, various functional groups on aromatic derivatives were tolerated (CN, CO$_2$Et, NO$_2$, CF$_3$, I, Br, Cl) as well as naphthalene and heterocyclic compounds. Worth mentioning that the reagent I was furnished in a 90% yield.

![Scheme 4. A complementary approach to synthesize the reagent I.](image)

**Applications of the PhSO$_2$SCF$_3$ reagent I as a SCF$_3$ source.** In 2017, the group of Xu reported the difunctionalization of alkenes 11 (Scheme 5). By combining visible light photoredox catalysis with gold catalysis, the concomitant formation of a C(sp$^3$)-SCF$_3$ and C(sp$^3$)-SO$_2$Ph bonds was possible in a regioselective manner. The trifluoromethylthiosulfonylation reaction was successfully applied to styrene derivatives in moderate to good yields as well as to internal alkenes (cyclic and acyclic ones) with high diastereoselectivities.
In the course of their investigations for the Ag-catalysed ring-opening difluoromethylthiolation of cycloalkanols, Hu, Shen and co-workers showed that the transformation might also be extended to the trifluoromethylthiolation reaction.\textsuperscript{18} Hence, under silver catalysis, the trifluoromethylthiolation of cyclobutanols\textsuperscript{13}, cyclopentanols\textsuperscript{15}, cyclohexanols\textsuperscript{17} and the cycloheptanol\textsuperscript{19} was possible in 56-90\% yields, offering an access to the corresponding distally trifluoromethylthiolated ketones (Scheme 6).

The reagent I was successfully used for the trifluoromethylthiosulfonfylolation of alkynes via an atom-transfer addition reaction. Indeed, merging visible light photoredox catalysis and gold catalysis, the difunctionalization of alkynes\textsuperscript{21} led to the trifluoromethylthiolated alkenes containing a sulfone residue (Scheme 7).\textsuperscript{19} (Hetero)aromatic terminal alkynes and one aliphatic alkyne were functionalized, offering an access to the corresponding alkenes as single $E$-isomers. The reaction was not restricted to terminal alkynes, since several internal alkynes were used, yielding the corresponding tetrasubstituted alkenes, the $E$ isomer being the major one. The synthetic utility of the approach was further demonstrated by the functionalization of bioactive compounds and drugs. When enynes\textsuperscript{23} were used, a radical cascade cyclization occurred allowing the concomitant formation of C(sp$^3$)-SO$_2$Ph and C(sp$^3$)-SCF$_3$ bonds (Scheme 7). Thio-functionalized dihydropyrans, other heterocycles and carbocycles were obtained in good to high yields (up to 86\%).\textsuperscript{19}
Note that, in 2018, I was employed in the trifluoromethylthiolation of aldehydes 25 in the presence of PIDA and NaN₃ at room temperature (Scheme 8). Six (hetero)aromatic aldehydes 25 were functionalized in high yields (70-96%).

Recently, Studer and co-workers employed the reagent I in the amide directed remote trifluoromethylthiolation reaction via a 1,5-HAT process. Not only secondary and tertiary C(sp³)-H bonds but also activated primary ones were efficiently and selectively trifluoromethylthiolated in good yields (Scheme 9).
Synthesis and applications of 4-Me-C₆H₄SO₂SCF₃ II and 4-MeO-C₆H₄SO₂SCF₃ III reagents. As above-mentioned, the different approaches depicted were also applied to the synthesis of various ArSO₂SCF₃. Among them, access to the S-trifluoromethyl 4-methylbenzenesulfonothioate II (4-Me-C₆H₄SO₂SCF₃ or TolSO₂SCF₃, Scheme 1, 2 and 4) and the S-trifluoromethyl 4-methoxybenzenesulfonothioate III (Scheme 2, 3 and 4) were reported and these reagents were also successfully applied as SCF₃ sources. Indeed, Zhao and co-workers reported a complementary approach for the synthesis of trifluoromethylthiolated oxindole derivatives 30 (Scheme 10). Using II as a coupling partner and in the presence of K₂S₂O₈, they depicted a silver mediated aryltrifluoromethylthiolation reaction of activated alkenes via a radical process. Under mild conditions, the aryltrifluoromethylthiolation of a panel of activated alkenes 29 was possible with yields up to 80%. The transformation turned out to be tolerant to various functional groups such as nitro and halogens for instance. The authors proposed the following mechanism: in situ generation of AgSCF₃ after reaction of the reagent II and AgF, which then would be oxidized by K₂S₂O₈, leading to the corresponding Ag(II)SCF₃ and its decomposition into Ag(I) and the SCF₃ radical. After addition of the SCF₃ radical on the activated alkene, an alkyl radical I1 would be formed, which would cyclise to provide the aryl radical II1. This later would undergo a final oxidation (intermediate III1) followed by a deprotonation step, which would furnish the trifluoromethylthiolated-substituted oxindole.
In addition, in 2018, the same group developed a methodology allowing the synthesis of aryl trifluoromethylthioether compounds 32 (Scheme 11) via a visible light photocatalytic approach. The S-trifluoromethyl 4-methoxybenzenesulfonothioate III was employed as the SCF$_3$ reagent, which brought a clear added value to the process compared to the previously use of the toxic and volatile (SCF$_3$)$_2$. The trifluoromethylthiolation reaction was conducted either on aryl diazonium tetrafluoroborate 31 or arylamines 33. In the last case, the reaction conditions were slightly modified allowing the in situ generation of the diazonium salt. In both cases, the transformation turned out to be functional group tolerant (COMe, CN, Br, NO$_2$ for instance). The following mechanism was proposed by the authors: in situ generation of the diazonium salt followed by the formation of an aryl radical after a single radical transfer (SET) with the excited photocatalyst (EYH$_2^*$. The later would react with the reagent III to afford the expected product and the sulfone radical. The photocatalyst would be regenerated after a SET with the sulfone radical. Note that very recently, a similar trifluoromethylthiolation reaction on diazonium salts was reported using the reagent II in the presence of a ruthenium-based photocatalyst.
2. Direct introduction of SR₇ groups with PhSO₂R₇ reagents (R₇ = CF₂H, CH₂F and C₂F₅)

In contrast with the recent synthetic advances made using the ArSO₂SCF₃ reagents, the introduction of other SR₇ groups remains limited. However, demonstrating interesting features, any advances made for the incorporation of the SCF₂H group and other SCF₂R₇ ones will be undeniably impact this very active research field. In that context, the main breakthroughs that have been recently developed for the direct introduction of SCF₂H and SCF₂FG groups onto molecules using ArSO₂SR₇ as SR₇ source will be summarized in the following section.

Synthesis and applications of the PhSO₂SCF₂H reagent IV

Synthesis of the PhSO₂SCF₂H reagent IV.

In 2016, the group of Lu and Shen investigated the radical difluoromethylthiolation of several classes of compounds. In that purpose, the design of a new reagent was realized (Scheme 12). The S-(difluoromethyl)benzenesulfonothioate (IV, PhSO₂SCF₂H) was prepared in 79% yield in a 20 mmol scale, according to a one pot two-step sequence from benzyl difluoromethyl sulfide.
The *in situ* generation of HCF$_2$SCI was followed by a nucleophilic substitution with PhSO$_3$Na. Worth mentioning that the synthesis of IV was easily scaled up to a 20 grams’ scale (120 mmol) and IV was obtained in a good 72% yield, showcasing the robustness of the synthesis.

![Scheme 12. Synthesis of the PhSO)$_2$SCF$_2$H reagent IV.](image)

**Applications of the PhSO$_2$SCF$_2$H reagent IV.** This reagent was successfully applied as a SCF$_2$H source, extending further the portfolio of tools for the direct introduction of this emergent moiety on various scaffolds.

First, the efficiency of the reagent was demonstrated in the silver-catalysed difluoromethylthiolating reaction of aryl and alkyl boronic acids 35 (Scheme 13). Indeed, electron-rich and electron-poor arenes, as well as primary alkyl boronic acids were efficiently converted into the corresponding products 36. The reaction was carried out under mild reaction conditions, offering a large functional group tolerance such as halides, ester, ketone, nitro. Nevertheless, limitations remained as heteroaryl boronic acids like the pyridine-3-boronic acid was not suitable substrates. Moreover, secondary and tertiary alkyl boronic acids were less efficient compared to primary ones.

![Scheme 13. Difluoromethylthiolation reaction of aryl and alkyl boronic acids with IV via Ag-catalysis.](image)

To further demonstrate the versatility of the difluoromethylthiolating reagent, other transformations were evaluated. The direct introduction of the SCF$_2$H moiety thanks to a silver-catalysed decarboxylative difluoromethylthiolation reaction in the presence of IV was depicted (Scheme 14). The transformation proceeded smoothly allowing the functionalization of tertiary, secondary ad even primary carboxylic acid derivatives 37, the latter being less efficient. Finally, the Ag-catalysed difunctionalization of alkene derivatives 39 with the concomitant formation of a C(sp$^3$)-SCF$_2$H bond and C(sp$^3$)-SO$_2$Ph bond formation was reported (Scheme 15, conditions A). Note that the reaction also proceeded smoothly under silver-free conditions (Scheme 15, conditions B). A large variety of aliphatic alkenes bearing various functional groups (eg. sulfone, ester, aldehyde, halide) and even heterocyclic moiety were functionalized. Nevertheless, styrenes or $\alpha,\beta$-unsaturated esters remain reluctant substrates.
Then, two years later, during their investigations towards the difunctionalization of alkynes merging photoredox catalysis and gold catalysis, Xu and co-workers showed that the difluoromethylthiosulfonylation reaction was also possible. A panel of terminal alkynes and an internal one 41 were functionalized in moderate to good yields leading to the corresponding SCF$_2$H-substituted alkenes 42 as E-isomers (Scheme 16).
In 2018, Wang and co-workers reported the direct introduction of the \( \text{SCF}_2\text{H} \) group onto aldehydes via a metal-free approach. Indeed, the combination of TBHP and the reagent IV allowed the functionalization of a large panel of (hetero)aromatic aldehydes (Scheme 17). The methodology was also applied to the difluoromethylthiolation of aliphatic and \( \alpha,\beta \)-unsaturated aldehydes, offering a straightforward access to difluoromethylthioethers. The reaction was turned out to be functional groups tolerant (cyano, ester, halides). The same year, Wang, Hu, Shen and co-workers, depicted a complementary approach using \( \text{NaN}_3 \) and PIFA to promote the difluoromethylthiolation of aldehydes via a free radical process (Scheme 18). The reaction occurred under mild conditions (room temperature). In addition, the transformation was not restricted to the \( \text{SCF}_2\text{H} \) part (see other sections). Various aromatic and aliphatic aldehydes, except benzylic aldehydes, were suitable substrates including benzothiophene and thiophene (21 examples, up to 91% yield).

In both studies, a similar mechanism was proposed: 1) generation of an acyl radical with the abstraction of the \( \text{H} \) of the aldehyde either directly with TBHP or with in the \textit{in situ} generated azide radical; then, 2) reaction with IV to afford the expected compound.
In course of the study towards the reactivity of the reagent IV, the group of Li demonstrated its successful use as a radical SCF$_2$H source in visible light promoted innate difluoromethylthiolation of aromatic derivatives 47 (Scheme 19). Indeed, they anticipated that under light irradiation, the homolytic cleavage of the S-S bond would be possible and favoured by the release of the stabilized phenyl sulfanyl radical. Using this metal-free approach, various heteroaromatic derivatives such as indoles, pyroles, azaindoles, pyrazoles, isoxazole, chromones and even thiophene were functionalized under mild conditions in the presence or not of a catalytic amount of tetrabutylammonim iodide (TBAI). The innate introduction of the radical SCF$_2$H group on electron rich arenes was also realized, and the functionalization generally occurred at the most electron–rich and less sterically hindered positions.

The following mechanism was proposed: first, the generation of the SCF$_2$H radical upon light irradiation would occur, which would result from either the homolysis of PhSO$_2$SCF$_2$H or the photo-induced electron transfer (PET) between the PhSO$_2$SCF$_2$H and iodide. Then, after addition of the SCF$_2$H radical onto arenes, the phenyl sulfonium radical would abstract the hydrogen atom to afford the corresponding difluoromethylthiolated product 48. Note that very recently the same group reported the generation of an aryl radical from potassium 4-biphenyl trifluoroborate through an $S_{N}2$ process. Its combination with PhSO$_2$SCF$_2$H as a radical trapping reagent in the presence of diacetyl under h$_\nu$ irradiation afforded the corresponding difluoromethylthiolated arene. 31
Scheme 19. Radical difluoromethylthiolation reaction with (hetero)aromatics under visible light with the reagent IV.

Very recently the visible light promoted difluoromethylthiolation of aryldiazonium salts was reported (Scheme 20). Using a catalytic amount of Ru(bpy)$_3$(PF$_6$)$_2$ in the presence of sodium ascorbate and the reagent IV, 28 (hetero)aryl diazonium salts were functionalized, offering a complementary synthetic route to the one developed by Li and co-workers.

Scheme 20. Radical difluoromethylthiolation reaction with (hetero)aromatics under visible light with the reagent IV.

The difluoromethylthiolation of alkyl ketone derivatives with PhSO$_2$SCF$_2$H at a remote position was investigated by Hu, Shen and co-workers (Scheme 21). In that purpose, the authors reported an efficient approach using cycloalkanols as precursors of functionalized alkyl ketones. Indeed, in 2018, they reported the silver-catalysed difluoromethylthiolation reaction of a variety of cycloalkanols. The construction of difluoromethylthioethers from cyclobutanol derivatives was carried in water in the presence of a surfactant sodium dodecyl sulfate (SDS), a catalytic amount of AgNO$_3$ and a stoichiometric amount of K$_2$S$_2$O$_8$ (15 examples, 50-95%). Not restricted to cyclobutanol, the transformation was extended to cyclopropanols (9 examples,
40-82%), cyclopentanols (3 examples, 60-70%), cyclohexanols (3 example, 61-70%) and cycloheptanol. The authors suggested that the reaction proceeded via the generation of a cycloalkoxy radical, which might lead to an alkyl radical after opening β-scission. Then, the intermediate could react with the reagent IV to afford the expected compounds.


Synthesis and applications of the PhSO₂SCH₂F reagent V

Synthesis of the PhSO₂SCH₂F reagent V. Present in compounds of interest such as the Fluticasone drug, known for its anti-inflammatory properties, the development of efficient tools for the direct introduction of the SCH₂F residue onto molecules is of high importance. In that context, the group of Lu and Shen designed the first electrophilic S-(fluoromethyl) benzenesulfonothioate (V, PhSO₂SCH₂F). This robust reagent was prepared from the commercially PhSO₂SNa 53 either with CH₂FCl or with CH₂FCl (Scheme 22).

Applications of the PhSO₂SCH₂F reagent V. Although restricted so far to a limited number of examples, this reagent V has already showcased its potential in different reactions.

Lu, Shen and co-workers studied the formation of C(sp²)-SCH₂F and C(sp³)-SCH₂F bonds (Scheme 23). Indeed, under copper catalysis, electron-rich and electron-poor aromatic boronic acids 54 were monofluoromethylated in an efficient manner. Note that in case of some products bearing electron-rich groups, an oxidation into the corresponding sulfoxide was necessary to prevent any defluorination of the products as a side reaction. In addition, the reagent V turned to be suitable for the difunctionalization of unactivated alkenes 56 leading to the corresponding monofluoromethylthioethers 57 with good yields, up to 95%. The
transformation was tolerant to various functional groups (halides, ketone, sulfonyl, ...) as well as heteroaryl groups (indolyl, furyl, thienyl moieties).

An additional example to showcase the efficiency of such reagent to introduce the SCH$_2$F group was recently reported by the groups of Wang, as well as Wang, Hu and Shen (Scheme 24). Indeed, Wang and co-workers studied the synthesis of monofluoromethylthioesters from aldehydes via a metal free approach. Using V as a SCH$_2$F radical source, a panel of (hetero)aromatic aldehydes as well as aliphatic and $\alpha,\beta$-unsaturated aldehydes were functionalized (29 examples, 42-91%). The presence of 2,2'-azobisisovaleronitrile (AMBN) as radical initiator was necessary for the generation of the acyl radical from the aldehyde. In the case of Shen and co-workers, the authors demonstrated the versatility and generality of the PhSO$_2$SR$_i$ reagents to introduce a variety of emergent SR$_i$ group on aldehydes including the SCH$_2$F moiety (6 examples, 44-87%).
Synthesis and applications of the PhSO₂SCF₂CF₃ reagent VI

The reagent VI was prepared in three steps from the commercially available benzyl mercaptan 60. The unique example of application of the PhSO₂SCF₂CF₃ reagent VI was recently depicted in the global study from Wang, Hu, Shen and co-workers about the functionalization of aldehydes with PhSO₂SR reagents. Encouraged by their success, the authors reported the synthesis of three unprecedented pentafluoroethylthioesters 62 (Scheme 25).
Conclusion

Extending the portfolio of tools enabling the synthesis of original fluorinated compounds is of high importance. Consequently, over the last years, many efforts focused on the development of original approaches for the introduction of emergent fluorinated motifs. This review showcased and discussed the use of several ArSO₂SR reagents as SR sources, a new emerging trend to access to SR₃-containing molecules. A myriad of transformations allowing the incorporation of the valuable SCF₃, SCF₂H, SCH₂F and SC₂F₅ groups were designed. We strongly believe that this review will bring considerable insights to the organofluorine chemistry community and would stimulate further developments for these promising emergent fluorinated groups.

Conflicts of interest

There are no conflicts to declare.

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Notes and references


For references cited therein, please refer to the original literature cited in the main text.


