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REVIEW

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Recent trends in dehydroxylative trifluoromethylation, -methoxylation, -methylthiolation, and -methylselenylation of alcohols

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Owing to the prevalence of hydroxyl groups on molecules, much attention has been paid to the synthesis of functionalized organic compounds by dehydroxylative functionalization of parent alcohols. In this context, dehydroxylative trifluoromethylation. trifluoromethoxylation, trifluoromethylthiolation. trifluoromethylselenylation of readily available alcohols have recently emerged as intriguing protocols for the single-step construction of diverse structures bearing C-CF₃, C-OCF₃, C-SCF₃, and C-SeCF₃ bonds, respectively. This Mini-Review aims to summarize the major progress and advances in this appealing research area with special emphasis on the mechanistic features of the reaction pathways.

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Scheme 1 Selected examples of CF₃/OCF₃/SCF₃-based drugs.

Introduction

The incorporation of fluorinated moieties and particularly the trifluoromethyl (CF_3) , trifluoromethoxy fluoromethylthio (SCF3), and trifluoromethylseleno (SeCF3) groups into organic molecules such as pharmaceuticals and agrochemicals can often substantially improve their physical, chemical, and biological properties because of the electronic properties, unique size, lipophilicity, and metabolic stability of these groups.^{1,2} There are different examples of commercially available human and veterinary drugs that contain a CF₃, OCF₃, or SCF₃ moiety in their structures (Scheme 1).3 However, there are no analogous CF₃Se-containing drugs, which is likely due to the limited synthetic options for their preparation.4 Compounds containing the titled functionalities are versatile synthetic intermediates and can function as suitable building blocks for the preparation of many valuable functional materials.5 Considering the widespread biological activities and synthetic usefulness of CF₃/OCF₃/SCF₃/SeCF₃-containing compounds, it has been of great synthetic interest to develop new, efficient, and practical methods for the introduction of these privileged moieties into organic molecules. 6-9

In recent years, direct dehydroxylative functionalization of alcohols has become one of the hottest research topics in organic chemistry because it is a powerful and general strategy

for the construction of various valuable functionalized organic compounds from inexpensive and abundantly available alcohols without isolation of intermediates.¹⁰ In this regard, dehydroxylative trifluoro-methylation, -methoxylation, -methylthiolation, and -methylselenylation of alcohols have captured the imagination of the organic chemical community and have become promising synthetic methods for constructing C-CF₃, C-OCF₃, C-SCF₃, and C-SeCF₃ bonds, respectively. These synthetic processes are advantageous because the starting materials possess high selectivity and stability, are abundant and inexpensive, with low toxicity, and there is no need for isolation of intermediates. In continuation of our interest in organofluorine chemistry¹¹ and modern organic synthesis, 12-18 in this Mini-Review, we will highlight the most important advances and progress in the arena of dehydroxylative trifluoro-methylation, -methoxylation, -methylthiolation, and -methylselenylation of alcohols (Scheme 2), with a particular emphasis on the mechanistic aspects of the reaction pathways.

Dehydroxylative trifluoromethylation

After their seminal work on the conversion of allylic alcohols to through esterification/ trifluoromethanes a two-step decarboxylative trifluoromethylation procedure,19 Altman and



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[CF₃] [OCF₃]

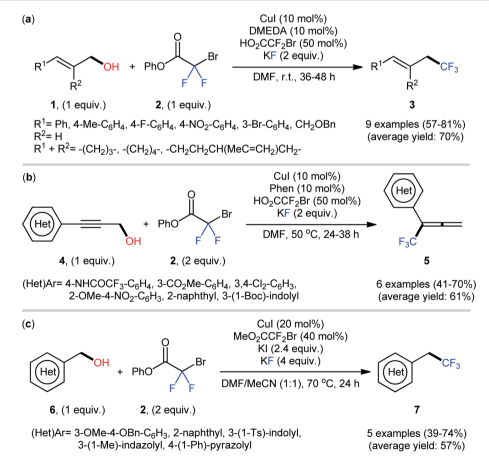
| CF₃ | OCF₃ | OCF₃ | OCF₃ | R¹ | R² | R² | R²

Scheme 2 Direct dehydroxylative trifluoro-methylation, -methoxylation, -methylthiolation, and -methylselenylation of alcohols.

[SCF₃]

SeCF₃

colleagues developed bench-stable phenyl bromodifluoroacetate (PhO₂CCF₂Br) as a new nucleophilic trifluoromethylation reagent, which can directly convert alcohols to trifluoromethanes in a single operation without isolation of intermediates.²⁰ They showed that treatment of various allylic alcohols 1 with over-stoichiometric amounts of phenyl bromodifluoroacetate 2 in the presence of the CuI/DMEDA/ HO₂CCF₂Br combination as a catalytic system in DMF at room temperature afforded the corresponding deoxytrifluoromethylated products 3 in good to high yields (Scheme 3a). The substrate scope was evaluated on nine terminally substituted allylic alcohols, which proved that various linear cinnamyl alcohols and endocyclic alkenols were compatible with this reaction. Moreover, this strategy was successfully applied for the direct deoxytrifluoromethylation of a series of propargylic alcohols 4 (by replacement of the DMEDA ligand with Phen) and (hetero)benzylic alcohols 6 (by employing MeO₂CCF₂Br reagent instead of HO₂CCF₂Br and using overstoichiometric amounts of KI) (Scheme 3b and c). Of note, propargylic alcohols underwent rearrangement and resulted in allenylic structures. Furthermore, the synthetic utility of this procedure was highlighted by gram-scale synthesis of (4,4,4trifluorobut-1-en-1-yl)benzene (75% yield on a 20 mmol scale). Finally, the synthetic potentiality of this strategy was showcased by preparing a fluorinated analogue of tebufenpyrad, an acaricide. Regrettably, the authors did not investigate the applicability of secondary and tertiary alcohols in their methodology. On the basis of several control experiments, a plausible mechanism was proposed by the authors for this trifluoromethylation protocol involving the following key steps transesterification (Scheme 4): (i) between



Scheme 3 Cu-catalyzed deoxytrifluoromethylation of (a) allylic alcohols 1; (b) propargylic alcohols 4; and (c) (hetero)benzylic alcohols 6 with phenyl bromodifluoroacetate 2 developed by Altman.

Scheme 4 The plausible mechanism for the reactions in Scheme 2.

bromodifluoroacetate 2 and the starting alcohol 1 to generate alkyl bromodifluoroacetate intermediate A; (ii) formation of the active $Cu-CF_3$ species B via interaction of the L_nCuI precatalyst with HO_2CCF_2Br ; (iii) direct nucleophilic trifluoromethylation

of intermediate **A** with Cu-CF₃ species **B** to give the desired product **3** and carboxylate-coordinated [Cu] complex **C**; and (iv) CO₂ liberation and halogen exchange to recover the active Cu-CF₃ species **B** and completion of the catalytic cycle.

Scheme 5 (a) Wu-Xiao's synthesis of (2,2,2-trifluoroethyl)arenes 10. (b) The proposed pathways for the formation of (2,2,2-trifluoroethyl)arenes 10.

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Scheme 6 Metal-free direct dehydroxytrifluoromethoxylation of alcohols 11 with trifluoromethyl tosylate 12.

In another report, this research group also developed [1,1′-biphenyl]-4-yl-2-bromo-2,2-difluoroacetate (BBDFA) as an efficient trifluoromethylating reagent for Cu-catalyzed dehydroxylative trifluoromethylation of alcohols.²¹ The reagent was synthetized on a 100 g scale in 93% yield *via* chlorination of commercially available bromodifluoroacetic acid with oxalyl chloride in the presence of a catalytic amount of DMF, followed by esterification of the resulting acid chloride with 4-phenylphenol, and it showed good thermal stability and high ability for trifluoromethylation of examined alcohols. However, the usefulness of this reagent was only demonstrated by deoxytrifluoromethylation of cinnamyl alcohol and 2-naphthalenemethanol, without any substrate scope exploration.

Drawing inspiration from these works, very recently, the Wu group, in collaboration with the Xiao group, described an interesting Cu(0)-catalyzed dehydroxylative trifluoromethylation of a library of (hetero)benzylic alcohols 8 with Chen's reagent (methyl fluorosulfonyldifluoroacetate; 9) in the presence of the Ph₃P/ICH₂CH₂I system, which acted as the activator of the hydroxyl group.²² The reactions were performed in DMF at 80 °C, tolerated a series of synthetically useful functionalities (e.g., -Br, -I, -CO₂Me, -CN, -NO₂), and provided the desired (2,2,2-trifluoroethyl)arenes 10 in modest to good yields (Scheme 5a).

Regarding the influence of electronic effects of substituent groups in the phenyl ring periphery of benzylic alcohols, electron-rich substrates afforded higher yields compared to electron-poor ones. It is noteworthy that although aryl bromides remained intact under these conditions, undesired trifluoromethylation of C-I bonds was observed in the case of aryl iodides. Regrettably, alkyl alcohols did not exhibit reactivity under standard reaction conditions. It was noteworthy that under relatively similar conditions, dehydroxylative difluoromethylation and trifluoromethylthiolation of a diverse range of aliphatic alcohols with TMSCF2H and AgSCF3, respectively, also effectively proceeded to afford the corresponding difluoromethylated and trifluoromethylthiolated compounds in reasonable yields. While the detailed mechanistic picture remains unclear, the authors speculated that phosphonium A, iminium C, or (hetero)benzyl halide D might be key intermediates for this trifluoromethylation reaction (Scheme 5b).

3. Dehydroxylative trifluoromethoxylation

At the outset of 2018, Tang and colleagues discovered that a combination of nucleophilic trifluoromethoxylating reagent 12 (trifluoromethyl tosylate), quaternary ammonium salt

Scheme 7 Mechanism proposed to explain the formation of alkyl trifluoromethyl ethers 13.

Scheme 8 Ph₃P/ICH₂CH₂I-promoted dehydroxylative trifluoromethoxylation of aliphatic alcohols **14** using AgOCF₃ as a nucleophilic trifluoromethoxylating reagent.

(tetramethylammonium bromide; TMABr), and fluoride source (CsF) enabled direct dehydroxylative trifluoromethoxylation of various primary and secondary alcohols **11**, giving the corresponding alkyl trifluoromethyl ethers **13** in moderate to high yields, ranging from 40% to 87% (Scheme 6).²³ The synthetic broad scope of the protocol was established using a library of various alkyl, benzyl, allyl, and propargyl alcohols. Significantly, the authors showcased the potentiality of this process by dehydroxytrifluoromethoxylation of complex bioactive molecules such as cholane-3,12,24-triol (a natural product) and

pleuromutilin (an antibacterial drug). The reaction demonstrated a high degree of chemoselectivity, and in the presence of a secondary or tertiary alcohol, the reaction preferentially took place at the position of a primary alcohol.

Based on mechanistic studies (isotope labelling and ¹⁹F NMR experiments), the author proposed a plausible mechanistic pathway for the above transformation, as depicted in Scheme 7.²⁴ The reaction begins with the release of trifluoromethoxide anion (CF₃O⁻) from reagent **12** under the action of a fluoride salt. Subsequently, CF₃O⁻ undergoes

$$ICH_{2}CH_{2}I \xrightarrow{Ph_{3}P} Ph_{3}P-I I \xrightarrow{Ph_{3}P-1} Ph_{2}Ph_{2}Ph_{3}P-I I \xrightarrow{Ph_{3}P-1} Ph_{3}P-I I \xrightarrow{Ph_{3}P-1}$$

Scheme 9 Mechanistic proposal for the formation of trifluoromethyl ethers 15.

Ar
$$= \frac{OH}{R^2} + CuSCF_3 = \frac{BF_3.Et_2O (2 \text{ equiv.})}{MeCN, r.t., 30 \text{ min}} = \frac{SCF_3}{R^2}$$

16, (1 equiv.) (1.5 equiv.) 17

18 examples (50-99%)

 $\begin{array}{l} \text{Ar= Ph, 4-OMe-C_6H_4, 4-Ph-C_6H_4, 4-F-C_6H_4, 4-CF_3-C_6H_4, 2-F-C_6H_4, } \\ 3,4-(OMe)_2-C_6H_3, 3,4-OCH_2$O-$C_6$H_3, 2,4,6-Me_3-$C_6$H_2, } \\ 3,4,5-(OMe)_3-$C_6$H_2, 2-(5-Me)-thienyl, 1-ferrocenyl \\ R^1= H, Me, Ph, 4-OMe-C_6H_4, 2-furyl \\ R^2= H, Me \\ R^1+ R^2= -(CH_2)_5- \end{array}$

(average yield: 88%)

$$Ar + R^1 = \bigcup_{s}^{h} \bigcup_{s}^{h}$$

Scheme 10 Lewis acid-mediated dehydroxylative trifluoromethylthiolation of benzylic alcohols 16 with CuSCF $_3$, as reported by Rueping.

decomposition to produce carbonic difluoride **A** which, upon esterification with alcohol **11**, generates alkyl fluoroformate intermediate **B**. Finally, nucleophilic substitution reaction of activated species **B** with *in situ-*generated CF_3O^- affords the final product **13**. Of note, the studies indicated that the presence of TMABr in the reaction mixture is crucial for improving the nucleophilicity of OCF_3 anion, while in the absence of any quaternary ammonium salt, inferior results in terms of product yield were observed.

In an attempt to further demonstrate the strength of this novel and interesting alkyl trifluoromethyl ether synthesis, Lin and Xiao along with their co-workers documented an elegant Ph₃P/ICH₂CH₂I-promoted trifluoromethoxylation of aliphatic alcohols using AgOCF₃ as a nucleophilic trifluoromethoxylating reagent, which allowed very rapid access to the corresponding dehydroxytrifluoromethoxylated products.²⁵ Through exploration and optimization of this dehydroxylative functionalization, the authors identified that the reaction rate is highly dependent on the nature of solvent. Among several solvents tested (*e.g.*,

DMSO, DMF, NMP, and toluene), DMF was found to be the most effective. Furthermore, the outcome of this transformation was also dramatically dependent on the reaction temperature. The best results were obtained by performing the process at 80 °C. A higher or lower temperature resulted in lower yields. With these optimized reaction conditions, 25 trifluoromethyl ethers 15 were synthesized in 16-76% yields from the corresponding aryl/ benzyl/allyl/propargyl alcohols 14 (Scheme 8). Notably, a diverse range of functional groups such as fluoro, chloro, bromo, iodo, cyano, nitro, ester, and ether functionalities were demonstrated to be well-tolerated by this protocol. However, the major drawback of this synthetic protocol was its very low efficiency for functionalization of alkyl alcohols. Intriguingly, the authors nicely solved this problem by replacing Ph₃P with Ph₂PCH= CH₂ and performing the process at 100 °C. However, the only reported case of a secondary alkyl alcohol led to a mediocre vield.

Mechanistically (Scheme 9), the reaction may be initiated by the reaction of Ph₃P with ICH₂CH₂I to give diiodophosphonium salt **A**, which upon coordination with the reaction solvent DMF, furnishes complex **B**. Subsequently, substitution of an alcohol **14** with a DMF molecule in complex **B** yields complex **C**, and after nucleophilic attack by trifluoromethoxy anion, generated from AgOCF₃ by precipitating AgI, affords the observed alkyl trifluoromethyl ether **15**. In another possibility, a sequential P–O bond formation and C–O bond cleavage process converts complex **B** into a triphenylphosphine oxide (Ph₃P=O) and the Vilsmeier–Haack-type intermediate **D**. Later, nucleophilic substitution of alcohol **14** with intermediate **D** leads to the formation of intermediate **E**, which after nucleophilic trifluoromethoxylation with CF₃O⁻, provides the final product **15**.

4. Dehydroxylative trifluoromethylthiolation

The first report of the direct dehydroxylative trifluoromethylthiolation of alcohols was published by Rueping

Scheme 11 Qing's synthesis of trifluoromethylthioethers 19

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Scheme 12 Proposed reaction mechanism for the synthesis of trifluoromethylthioethers 19 starting from alcohols 18 with AgSCF₃.

and co-workers in 2014,26 who disclosed that the treatment of benzylic alcohols 16 with CuSCF3 as a stable and readily available SCF₃ source and BF₃·Et₂O as a Lewis acid additive under an atmosphere resulted in the formation fluoromethylthioethers 17 in moderate to quantitative yields within 30 min (Scheme 10). All three types of alcohols (primary, secondary, and tertiary) were applicable to this reaction, and in all cases, the corresponding trifluoromethylthioethers were selectively obtained. However, due to the strong acidic conditions, poor functional group tolerance occurs with this method. It should be mentioned that the presence of BF₃·Et₂O was essential for the success of this C-S bond formation reaction.

No product was detected with the lack of this additive, even at elevated temperatures.

Other Brønsted or Lewis acids such as MsOH, TsOH, TFA, Sc(OTf)3, Bi(OTf)3, and In(OTf)3 were also tested and proved to be completely ineffective. The identical reaction conditions were also applied for trifluoromethylthiolation of allylic alcohols to give the corresponding allylic trifluoromethyl thioethers in good to excellent yields (9 examples, 73-96% yield) and high regioselectivities, in which regardless of the substitution pattern, conjugated aryl/olefin products were predominantly formed in the case of aryl-substituted allyl alcohols. Mechanistic investigations indicated that the reaction may occur via an S_N1-type process, as evidenced by the formation of racemic products from enantiopure alcohols.

Immediately after, Qing and collaborators developed a similar dehydroxytrifluoromethylthiolation of alkyl alcohols 18 with AgSCF₃, employing a large excess of the mild reagent n-Bu₄NI as activator and toluene as solvent (Scheme 11).²⁷ The reaction was shown to be quite general, and a diverse range of primary aliphatic, benzylic, allylic, and propargylic alcohols participated in the trifluoromethylthiolation. Moreover, secondary alcohols also accomplished production of the corresponding products albeit the addition of a very large amount of another activator, KI (8 equiv.), and elevated reaction temperature (120 °C) were needed to prevent the competitive elimination reaction. However, tertiary alcohols were not suitable substrates for this transformation.

Scheme 13 Qing's synthesis of α -aryl- β -(trifluoromethylthio)acrylates 21

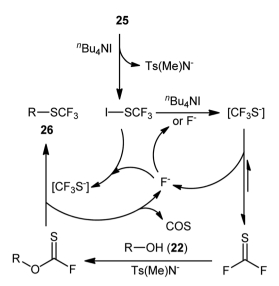
Scheme 14 Synthesis of 3-(((trifluoromethyl)thio)methylene)indolin-2-ones 23 via "Bu₄NI-mediated dehydroxytrifluoromethylthiolation of 3-(hydroxymethylene)indolin-2-one derivatives 22 with AgSCF₃.

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Interestingly, several biologically active alcohols such as idebenone **18a** (an anti-Alzheimer's drug), galantamine **18b** (an anti-Alzheimer's and anti-dementia drug), and epiandrosterone **18c** (a steroid hormone) also responded to the reaction. Notably, the authors observed that changing the ratio of $AgSCF_3/n-Bu_4NI$ from 1:3 to 1:1 led to the selective formation of alkyl fluorides instead of the expected alkyl trifluoromethylthioethers. A plausible mechanism based on previous studies is outlined in Scheme **12**.

In their subsequent studies, this research group extended the scope of their methodology to enols. Thus, a library of (Z)-ethyl 2-(aryl)-3-hydroxyacrylates **20** were reacted with AgSCF₃, n-Bu₄NI, and KI in toluene at 120 °C leading to the respective α -aryl- β -(trifluoromethylthio)acrylates **21** in moderate to excellent yields and satisfactory stereoselectivities in favor of the (E)-products (Scheme 13). Under the same conditions, they also executed the direct dehydroxytrifluoromethylthiolation of a small series of 3-(hydroxymethylene)indolin-2-one derivatives **22**, offering a decent yield of the desired 3-(((trifluoromethyl) thio)methylene)indolin-2-ones **23** (Scheme 14). Interestingly, two electron-deficient phenols were also tested and gave products in satisfactory yields. To the best of our knowledge, this is the first and only reported example of dehydroxylative trifluoromethylthiolation of C(sp²)-OH bonds.

At the outset of 2016, Billard's research group devised an elegant metal-free method for dehydroxylative



Scheme 16 Proposed pathway of alkyl trifluoromethylthioethers 25 formation.

trifluoromethylthiolation of alkyl alcohols **24** using the second-generation trifluoromethanesulfenamide reagent **25** as an SCF₃ source and *n*-Bu₄NI as an activator in refluxing acetone to afford the corresponding alkyl trifluoromethylthioethers **26** in an efficient manner (Scheme 15a).²⁹ The experiments

(a) OH
$$R^2$$
 + T_S SCF $_3$ R^2 R^2 24, (1 equiv.) 25, (2 equiv.) 26 12 examples (27-96%) (average yield: 70%) R^1 = Ph, 4-OMe-C $_6$ H $_4$, 4-Ph-C $_6$ H $_4$, 2-Br-C $_6$ H $_4$, 2-Br-5-F-C $_6$ H $_3$, 3-Me-4-NO $_2$ -C $_6$ H $_3$, phenylacetylenyl, -CH $_2$ CH=CH $_2$, -CH $_2$ CH(Me)CH $_2$ CH=CMe $_2$, -CH=C(Me)CH $_2$ CH=CMe $_2$, R^2 = H, Me R^1 + R^2 = -CH $_2$ C(Me) $_2$ CH $_2$ C(Me)=CH- (b) R^1 + R^3 + AgSCF $_3$ R^3 R^4 R^3 + AgSCF $_3$ R^3 R^3 R^4 R^3 + AgSCF $_3$ R^4 R^3 + AgSCF $_3$ R^4 R^3 + AgSCF $_3$ R^4 R^3 R^4 R^5 R^5 R^5 R^5 R^5 R^6 R

Scheme 15 (a) Metal-free dehydroxylative trifluoromethylthiolation of alkyl alcohols 24 with trifluoromethanesulfenamide 25; (b) solvent-free microwave-assisted synthesis of alkyl trifluoromethylthioethers 28 from alkyl alcohols 27 and AgSCF₃.

(b)
$$R^{+}$$
 OTf R^{2} $NEt(^{i}Pr)_{2}$ (2 equiv.) R^{2} R^{2}

 $\begin{array}{l} R^{1} = Ph, \ 4-Me-C_{6}H_{4}, \ 4-Ph-C_{6}H_{4}, \ 4-NO_{2}-C_{6}H_{4}, \ 4-CO_{2}Me-C_{6}H_{4}, \ 4-Cl-C_{6}H_{4}, \ 4-Br-C_{6}H_{4}, \ 4-LCl-C_{6}H_{4}, \ 4-Br-C_{6}H_{4}, \ 4-LCl-C_{6}H_{4}, \ 4-Br-C_{6}H_{4}, \ 3-Br-C_{6}H_{4}, \ 3-NO_{2}-C_{6}H_{4}, \ 2,4,6-Cl_{3}-C_{6}H_{2}, \ 2-Br-5-F-C_{6}H_{3}, \ C_{6}F_{5}, \ C_{6}Br_{5}, \ 1-naphthyl, \ 2-paphthyl, \ 2-pyridyl, \ -CH_{2}Bn, \ -(CH_{2})_{2}Bn, \ -(CH_{2})_{3}Bn, \ -CH_{2}-1-naphthyl, \ -(CH_{2})_{3}-4-OMe-C_{6}H_{4}, \ ^{n}C_{8}H_{17}, \ ^{n}C_{9}H_{19}, \ ^{n}C_{11}H_{23}, \ ^{n}C_{13}H_{27}, \ NPhth, \ phenylacetylenyl \ R^{2} = H, \ Me, \ ^{n}Pr, \ Ph \end{array}$

Scheme 17 Hünig's base-mediated dehydroxylative trifluoromethylthiolation of alkyl/benzyl/propargyl alcohols 30 with benzothiazolium reagent 29.

demonstrated that the outcome of this reaction was not highly sensitive to the electronic nature of substrates, and therefore benzylic alcohols with either neutral, electron-donating, or electron-withdrawing substituents gave the trifluoromethylthiolated products in relatively similar yields. However, the steric effect was very strong (92% yield for benzyl alcohol to 46% for α -methylbenzyl alcohol). Unfortunately, the applicability of tertiary alcohols as starting materials was not explored in this study.

A presumptive mechanism for this dehydroxylative trifluoromethylthiolation reaction is represented in Scheme 16. Subsequently, a straightforward and greener approach for the synthesis of alkyl trifluoromethylthioethers **28** by the reaction between alkyl alcohols **27** and 1-*n*-butyl-3-methylimidazolium trifluoromethylthiolate ([bmim][SCF₃]) generated *in situ* from [bmim][I] and AgSCF₃ was reported by Pégot, Magnier, and coworkers.³⁰ The reactions were implemented under microwave irradiation and solvent-free conditions, tolerated primary, secondary, as well as tertiary alcohols, and rapidly provided the desired products in poor to excellent yields (Scheme 15b). Recycling tests indicated that the ionic liquid can be reused in

$$\begin{array}{c} \text{OTf} \\ \text{SCF}_3 \end{array} \xrightarrow{\text{NEt}(^i\text{Pr})_2} \\ \text{RCH}_2\text{OH} (\textbf{30}) \end{array} \longrightarrow \begin{array}{c} \text{NEt}(^i\text{Pr})_2 \\ \text{RCH}_2\text{SCF}_3 \end{array} + \begin{array}{c} \text{NET}(^i\text{Pr})_2 \\ \text{RCH}_2\text{SCF}_3 \\ \text{RCH}_2\text{SCF}_3 \end{array} + \begin{array}{c} \text{NCH}_2\text{SCF}_3 \\ \text{RCH}_2\text{SCF}_3 \end{array} + \begin{array}{c} \text{NCH}_2\text{SCF}_$$

Scheme 18 Mechanism proposed to explain the formation of trifluoromethylthioethers 31.

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Ph₃P/ICH₂CH₂I/n-Bu₄NI-mediated direct conversion of alcohols 32 into the corresponding trifluoromethyl thioethers 33 using Scheme 19 AqSCF₃.

several consecutive trials without significant loss of its activity (from 96% in the first run to 96% in the fifth run).

In 2019, Hopkinson and co-workers designed and synthesized a new bench-stable 2-trifluoromethylthio-substituted benzothiazolium salt (BT-SCF3; 29) through visible-lightinduced, Ir-catalyzed trifluoromethylthiolation of inexpensive 2-mercaptobenzothiazole disulfide (MBTS) with the Langlois reagent (NaSO₂CF₃) and subsequent reaction of the generated 2-SCF₃-substituted benzothiazole with MeOTf in DCM at room temperature (Scheme 17a).31 The activity of this purely organic trifluoromethylthiolating reagent has been evaluated in the dehydroxylative trifluoromethylthiolation of a broad set of alkyl/ benzyl/propargyl alcohols 30 in the presence of Hünig's base (NEt(ⁱPr)₂) in MeCN. Moderate to almost quantitative yields of the target trifluoromethyl thioethers 31 were obtained within 1-2 h at room temperature (Scheme 17b). The reaction exhibited satisfactory tolerance for an array of catalytically reactive functional groups (e.g., F, Cl, Br, I, CF₃, CO₂Me, SMe, NO₂), and thus, promised further elaboration of the end products. It is worthwhile to note that the authors nicely adapted this approach to the direct construction of SeCF3-substituted compounds from

alcohols by developing a similar trifluoromethylselenylsubstituted benzothiazolium salt (BT-SeCF₃).

As for the mechanism, the authors speculated that the reaction most likely proceeds through the formation of key electrophilic 2-alkoxybenzothiazolium species A via nucleophilic attack of the alcohol 30 in the presence of $NEt(^{i}Pr)_{2}$ at the C2-position of the BT-SCF₃ reagent 29 and subsequent nucleophilic substitution reaction with in situ generated -SCF₃ anion (Scheme 18). Guided by the same principle, a similar dehydroxylative functionalization strategy was applied by the same research group towards the synthesis of various perfluoroalkyl thioethers32 and thioesters.33

In a recent report, Wu, Xiao, and colleagues accomplished the direct conversion of alcohols 32 into the corresponding trifluoromethyl thioethers 33 using AgSCF₃ as a source of F₃CS group and the Ph₃P/ICH₂CH₂I/n-Bu₄NI combination as an activation system in a 2:1 mixture of DMF and MeCN.22 The reaction was compatible with a variety of functionalized benzylic alcohols, as well as heterobenzylic alcohols such as hydroxymethyl pyridines, quinolines, thiazoles, benzothiols, even simple allylic and propargylic alcohols (Scheme 19). As

OH
$$R^2$$
 + SeCF₃ NEt(i Pr)₂ (2 equiv.) R^2 R^2 R^2 R^2 R^2 R^2 R^2 R^2 R^3 R^4 (1equiv.) 35, (1.25 equiv.) R^4 R^4

Scheme 20 Hopkinson's synthesis of trifluoromethylselenoalkanes 36.

for alkyl alcohols, the corresponding products were obtained in poor yields. In this case, when Ph₂PCH=CH₂ was used instead of Ph₃P, the product yields were significantly increased. The plausible mechanism for this transformation is analogous to the one depicted in Scheme 5. It should be mentioned that this mechanism is tentative and lacks experimental evidence.

5. Dehydroxylative trifluoromethylselenylation

Compared to trifluoromethylation, trifluoromethoxylation, and trifluoromethylthiolation. the direct fluoromethylselenylation of alcohols is relatively less explored, although there has been much recent attention on the development of novel and efficient methodologies for the synthesis of SeCF₃-substituted compounds.³⁴ In 2019, in the same study describing the deoxytrifluoromethylthiolation of alcohols with the benzothiazolium salt BT-SCF3, Hopkinson's research team reported the successful NEt(1Pr)2-promoted preparation of trifluoromethylselenoalkanes 36 from the respective alcohols 34 employing 2-trifluoromethylseleno-substituted benzothiazolium salt (BT-SeCF₃; 35) (Scheme 20).31 Notably, BT-SeCF₃ was synthesized via the same general strategy used to prepare BT-SCF₃.

In 2020, using [Me₄N][SeCF₃] salt as a stable, non-volatile, and readily accessible source of nucleophilic SeCF₃, Zhang and colleagues engineered the direct dehydroxylative trifluoromethylselenylation of alcohols 37 for the synthesis of valuable alkyl trifluoromethyl selenoethers 38 under catalyst-free conditions.³⁵ By employing 3-phenylpropyl alcohol as the model reactant, several additives such as CaF₂, CaCl₂, CaBr₂,

Ca(OTf)₂, Ca(C₂O₄), CaSO₄, HCl, and LiI were carefully screened. Among them, excellent results were obtained for this transformation with CaCl₂, whereas MeCN was found to be the most effective solvent among the other common organic solvents tested (DMA, DMF, NMP, DMSO, MeCN, DCM, toluene, and 1,4-dioxane).³⁶ Evaluation of the substrate scope clearly demonstrated that the reaction was tolerant to a variety of primary and secondary aliphatic alcohols (Scheme 21). However, tertiary alcohols provided complicated mixtures. In order to elucidate the mechanism of the reaction, the authors performed several control experiments, such as GC-MS analyses, ¹⁹F NMR studies, and others.³⁷

From these results, the authors proposed two possible pathways for this transformation. The first pathway (Scheme 22, path A) starts with the formation of carbonoselenoic difluoride intermediate A through the decomposition of SeCF3 with CaCl₂ as a fluoride scavenger, which after reaction with another two equivalents of SeCF₃ in the presence of Ca²⁺ cations provides bis(trifluoromethyl)-carbonotriselenoate B (this intermediate was detected by ¹⁹F NMR and GC-MS analyses). Subsequently, the nucleophilic substitution of key intermediate B by alcohol 37 leads to a carbonoselenoate C. Finally, the nucleophilic attack of intermediate C with SeCF3 anion provides the target trifluoromethylselenylated product 38. The key steps of the second possible route (Scheme 22, path B) are the generation of O-alkyl carbonofluoridoselenoate D via straightforward reaction of intermediate A with alcohol 37 and its nucleophilic substitution by SeCF₃ to form the observed product 38. According to the authors, pathway B is not likely the in the Ca-mediated dehydroxy fluoromethylselenylation,38 especially when using a large excess of [Me₄N][SeCF₃].

Scheme 21 Selected examples of CaCl₂-promoted dehydroxylative trifluoromethylselenylation of alcohols 37 with [Me₄N][SeCF₃].

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Scheme 22 Mechanistic proposal for the reaction in Scheme 21

Conclusion

Because of their unique physicochemical and biological properties, there has been broad interest in fluorinated compounds in various research fields such as pharmaceuticals, medical imaging, agrochemicals, and materials science. Among all fluorine-containing functionalities, the trifluoromethyl (CF₃), trifluoromethoxyl (OCF₃), trifluoromethylthiol (SCF₃), and trifluoromethylselenyl (SeCF3) groups are becoming increasingly prominent in new drugs due to their strong electron-withdrawing nature and high lipophilic properties. Therefore, the high demand for biologically active compounds has stimulated significant efforts to develop efficient methods for the installation of these groups into organic molecules. As shown in this Mini-Review, recently direct dehydroxylative trifluoromethylation, tritrifluoromethylthiolation, fluoromethoxylation, fluoromethylselenylation of alcohols have emerged as efficient new methods for the construction of CF₃/OCF₃/SCF₃/SeCF₃-functionalized compounds from inexpensive and readily available starting materials without the need for time-consuming prefunctionalization steps.

As illustrated, various aliphatic and benzylic alcohols were applicable in these reactions. However, aromatic alcohols were mainly unsuitable substrates. Therefore, many more studies are needed to develop efficient procedures that allow trifluoromethylation, -methoxylation, -methylthiolation, and -methylselenylation reactions of aromatic alcohols. Moreover, there are insufficient reported examples for some reactions such as trifluoromethylselenylations, and thus, additional study is necessary to determine the scope and limitations of these reactions.

Conflicts of interest

There are no conflicts to declare.

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