Aromatic and heterocyclic perfluoroalkyl sulfides. Methods of preparation

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Abstract
This review covers all of the common methods for the syntheses of aromatic and heterocyclic perfluoroalkyl sulfides, a class of compounds which is finding increasing application as starting materials for the preparation of agrochemicals, pharmaceutical products and, more generally, fine chemicals. A systematic approach is taken depending on the mode of incorporation of the SR\textsubscript{F} groups and also on the type of reagents used.

Review
1. Introduction
Perfluoroalkyl sulfides of aromatic and heterocyclic compounds have been an important aspect in the general development of organofluorine chemistry over the last twenty years.

Alkyl aryl sulfides containing partly fluorinated aliphatic moieties have been widely used for a number of years. Their methods of preparation, for example, by the reaction of thiols with fluoro-olefins or with chloropolyfluoroalkanes are well known and have been widely used. In contrast, sulfides with fully fluorinated aliphatic chains have been limited to trifluoromethylated compounds. This was due to the unique preparation (at that time) of such compounds by means of two consecutive reaction steps: the chlorination of the side chain followed by replacement of the chlorine atoms by fluorine. This procedure enabled only the preparation of CF\textsubscript{3}S-derivatives because it is not possible to synthesize perchloroalkylated aromatic sulfides larger than CCl\textsubscript{3}S. This is currently still the case. Iodoperfluoroalkanes as perfluoroalkylating agents have only emerged rather recently.

New synthetic procedures to access this class of compounds have appeared which make use of novel intermediates. Thus, single-electron oxidation or reduction enables the generation of perfluoroalkyl radicals. Two-electron reduction of perfluoroalkyl iodides generates perfluoroalkyl carbanions, which may be stabilized by organophosphorus and organosilicon ligands and even by dimethylformamide.

One of the driving forces for the synthesis of perfluoroalkyl sulphides is the high lipophilic properties of perfluoroalkythio
groups (the greatest Hansch constant $\pi = 1.44$, belongs to SCF$_3$ group [1]), which increases the ability of such molecules to cross lipid membranes and creates opportunities for the modification of known and new drugs. Thus this group is a useful substituent in agrochemicals and pharmaceuticals [2-4]. Examples of bioactive compounds containing SCF$_3$, SOCF$_3$ and SO$_2$CF$_3$ groups are shown in Figure 1 and Figure 2.

The synthesis of a large number of potential hypotensive agents containing SR$_F$ and SO$_2$RF groups of the 1,4-dihydropyridine class and also of Losartan (Dup 753) analogues which are used clinically for the treatment of cardiovascular diseases have also been developed [5,6] (Figure 3).
Other patented compounds containing perfluoroalkyl thio substituents are illustrated in Figure 4 and Figure 5 along with their pharmacological functions [7-11]. These examples represent only a small number of the vast array of organic compounds with SR_F, SOR_F or SO_2R_F groups which display pharmacological activity and interest in such analogues continues to grow.

Previous reviews in this area are either dated [19] or focus on specialist aspects such as perfluoroalkyl radicals [20-22], fluorinated carbanions [23], organometallic compounds [24,25], per-
fluoroalkyl sulfenyl halides [26], perfluoroalkyl silicon reagents [27-32], the trifluoromethylthio anion [29] or electrophilic perfluoroalkylating agents [33]. Others are devoted to particular methods such as trifluoromethylation initiated by sodium dithionite [34] or the electrochemical introduction of fluoroalkyl groups in organic molecules [35]. Moreover, many of the reviews on the subject are very general [28,30,32,36].

The present work reviews synthetic methods employed to prepare aromatic and heterocyclic perfluoroalkyl sulfides and is systematized depending on the mode of constructing the SRF groups and also on the nature of the starting materials.

1. The halogenation of SAIk-derivatives with subsequent replacement of the halogen atoms by fluorine.
2. The introduction of SRF-moieties into aromatic compounds by both electrophilic and nucleophilic reagents.
3. Various modes of perfluoroalkylation of organosulfur compounds including cationic, anionic, radical and ion-radical variants.

2. Substitution of halogen atoms by fluorine in aryl-α-polyhalogenoalkyl sulfides

Substitution of the halogen atoms in SAIkHlg groups (mainly chlorine) using antimony trifluoride [37], is the oldest method of perfluoroalkylsulfide preparation and is still commercially significant.

The reaction is carried out by heating a mixture of aryl trichloromethyl sulfide with an excess of SbF₃ in the absence of a solvent. For industrial processes, dry hydrogen fluoride is used as the fluorinating agent (Scheme 1).

The presence of halogen atoms and electron-withdrawing groups such as NO₂, CF₃ or COCl in the aromatic ring of trichlorothioanisole does not influence the fluorination and the reaction is not hindered by bulky ortho-substituents e.g., phthalic acid imide [53] or N-substituted anilines [54]. Other reactive substituents, for example 3-SCl₃ or 4-COCI are also fluorinated and form 1,3-bis(SCF₃) benzene [38-40] and 4-SCF₃-benzoic acid fluoride, respectively [55].

The use of hydrogen fluoride has some advantages. Due to its low boiling point (+19.4 °C) and good solubility in water, excess HF is easily removed from the reaction mixture. Unlike HF, reactions with SbF₃ can be carried out in glass. The SbF₃ must be freshly sublimed and used in a corrosion-proof vessel. Attempts to use less aggressive fluoride ion sources, e.g., KF/18-Crown-6 in CH₃CN or KF/Bu₄N⁺Cl⁻ under phase-transfer conditions, have been unsuccessful [56].

The method does not give access to longer perfluoroalkyl sulfides because the required aryl perchloroalkyl sulfide precursors are not easily accessible [57,58]. However, pentafluoroethyl ethers of various thiophenols (or phenols) can be obtained by the more sequential process as shown in Scheme 2 [59].

Use of mixed (Cl/F) polyhalogenofluoro alkanes as partial fluorinated alkylation agents generates the corresponding sulfides which are appropriate precursors for subsequent conversion to perfluoroalkyl thioethers. For example, α,α-difluoro polyhalogenoalkyl sulfides and α,α-dichlorotrifluoroethyl sulfide can be obtained by reaction of thiophenols with dihalogenodifluoro methane [60-62], per(halogenofluoro)ethanes [60,63,64] and 2,2,2-trifluorotrichloroethane.

The Cl- and Br-substituents can then be replaced by fluorine without use of HF or SbF₃ [61]. As shown in Scheme 3 [65], bromine to fluorine exchange is possible by the use of other heavy metal fluorides, and even by silver tetrafluoroborate under mild conditions.
The halex-method allows the selective preparation of α,α-di-fluoroalkyl aryl sulfides (and also ethers, sulfoxides and sulfones) as intermediate products in the synthesis of herbicides [66,67]. Interestingly, the reaction of anhydrous hydrogen fluoride with aryl α,α,β-trichloroisobutyl sulfide at 20 °C leads only to substitution of the α-chlorine atoms, whilst at a higher temperature and pressure a more complete fluorination with rearrangement is observed [67] (Scheme 4).

Hydrogen fluoride/fluoride complexes such as H$_2$F$_3$ stabilized on a polymer [68] show even greater selectivity. For example, only one chlorine atom of the α,α-dichloromethylene group of benzyl alkyl sulfide is substituted by the reagent (Scheme 5).

Thus, halogen atoms replacement by fluorine is an effective and cheap method for preparing aromatic and heterocyclic perfluoroalkyl sulfides. Application of the appropriate conditions allows control and a degree of selectivity thus making this method an important industrial process.

3. Introduction of the aryl SR$_F$ moiety
3.1. Electrophilic introduction of SR$_F$ groups

Perfluoroalkyl sulfenyl chlorides react with electron rich aromatic and heterocyclic compounds, to give SR$_F$ derivatives. Thus, phenol, o-hydroquinone and their derivatives react with CF$_3$SCl to yield p-hydroxyaryl trifluoromethyl sulfides (Scheme 6).

The best yields are achieved when electron-donating substituents are present on the ring. In the case of m-cresol and m-chlorophenol a small degree of o-substitution was observed. Phenol is a poor substrate in the reaction (Scheme 6) however, when FeCl$_3$ was used as a catalyst the yield of p-HOC$_6$H$_4$SCF$_3$ was increased, albeit only slightly (30%). A significant improvement in yield occurs (72%) when the reaction is conducted with pyridine in chloroform and at ambient temperatures (0–20 °C) [70,71]. Under these conditions and with electron-donating substituents in the phenol, two and even three perfluoroalkylthio groups can be introduced (Scheme 7).

Forcing conditions are required for the introduction of three CF$_3$S-groups. This can be achieved either by activation with iron powder under pressure (or by conduction the reaction in a...
steel autoclave) or by the presence of two donor groups in meta-positions [71].

For \( p \)-hydroquinone, reaction with \( CF_3SCl \) in the presence of pyridine results only in the formation of a chlorohydroquinone pyridinium species [72], and neutral conditions are required in this case [69]. For the synthesis of poly(SCF\(_3\)) substituted \( p \)-hydroquinones, Scribner oxidized 2,6-bis(SCF\(_3\))-4-methoxyphenol to generate 2,6-bis(SCF\(_3\))-1,4-benzoquinone. The addition of CF\(_3\)SH in the presence of pyridine to the bis-compound gave 2,3,5-tris(SCF\(_3\))hydroquinone [72] which could be subsequently converted into tetrakis(SCF\(_3\))-1,4-hydroquinone (Scheme 8).

Unlike \( p \)-hydroquinone, resorcinols and phloroglucinols perhaps surprisingly react with \( R_2SCl \) [75] to generate mono-perfluoroalkyl thio derivatives. With iron powder as a catalyst bis(SR\(_2\))-derivatives can be obtained (Scheme 9).

The introduction of strong electron-donating meta groups significantly activates the aromatic nuclei not only for \( N,N \)-bis-substituted anilines but also for \( N \)-monosubstituted substrates and even those with a free NH\(_2\) group (Scheme 11).

Similarly, methyl benzoates and benzaldehydes with two and especially three hydroxyl groups form bis(CF\(_3\)S)-substituted derivatives without of catalyst.

Analogous reactions are observed with aniline. However, since reaction takes place in the first instance on the amino group [74,76], for the introduction of SCF\(_3\) group into the aromatic ring the amino function must be protected. Mono-\( N \)-substitution is insufficient: \( N \)-methyl aniline, \( N \)-(SCF\(_3\))aniline and \( N \)-(Ac)-\( m \)-toluidine all yield mainly \( N \)-(SCF\(_3\))-derivatives, and only a small amount of aromatic CF\(_3\)S-substitution is observed [74]. The best results are achieved [70,74] with \( N,N \)-bis-substituted aniline (Scheme 10).

**Scheme 7:** Introduction of SCF\(_3\) groups into activated phenols [71-74].

**Scheme 8:** Preparation of tetrakis(SCF\(_3\))-4-methoxyphenol [72].
In naphthalene and benzophenone derivatives only those rings containing hydroxy or amino groups undergo perfluoroalkylsulfanylation [74,75]. Other electron-donating substituents on the aromatic ring are not so activating for reaction with CF₃SCl. For example, thiophenol [76] forms only phenyltrifluoromethyl disulfide [70]. The presence of a methyl group and halogens requires high temperatures (100–200 °C) and the presence of catalysts (HF or BF₃) for reaction and yields of the corresponding aryltrifluoromethyl sulfides are only 25–60%. Both toluene and halobenzenes lead to mixtures of isomers [70].

Benzene undergoes trifluoromethylsulfanylation with trifluoromethanesulfonic acid as a catalyst even at 20 °C. However, further reaction of the resultant phenyltrifluoromethyl sulfide leads mainly to chlorination with only minor amounts of bis-(CF₃S) products (Scheme 12).

Aryl magnesium [78] and -mercury [79] compounds have been employed for the introduction of CF₃S groups. Such reactions proceed in ether or THF at low temperatures; however, the yields of aryltrifluoromethyl sulfides do not exceed 50–60% and are accompanied with halogenated side-products (Scheme 13).

Among heterocyclic systems, pyrroles are the best substrates for reaction with trifluoromethyl-, difluorochloro- and dichlorofluoromethyl sulfenyl chlorides. Their reactivity exceeds that of benzene and its organometallic derivatives [80]. An excess of reagent gives bis- (SCF₃) pyrrole derivatives as shown in Scheme 14.
Condensed pyrroles also react readily with CF₃SCI. Indole undergoes substitution, as expected, at the 3-position [80], while indolizine and some of its derivatives give 1,3-bis (SCF₃)-substituted products, in some cases, in quantitative yield [81]. It is interesting to note that not only hydrogen, but also an acetyl group in the 1-position is substituted (Scheme 15).

However, no reaction occurs when there are two electron-withdrawing groups in the five-membered indolizine ring (e.g. R = Ph, and X = COPh or NO₂). By contrast, in the case of 1-benzyl-2-methyl indolizine [81] both the pyrrole and the aromatic ring of the benzyl group undergo trifluoromethylsulfanylation. Only N-substitution occurs in the case of carbazole [80].

N-Methylpyrrole can be variously substituted depending on the conditions as illustrated in Scheme 16.

Heating N-methylpyrrole in CHCl₃/Py affords the 2-SCF₃ derivative along with a small amount of 3-SCF₃-N-methylpyrrole [83]. Attempted selective introduction of the second SCF₃ group at −30 °C with C₄F₉SO₃H to 2-trifluoromethylsulfanylpiperidine was unsuccessful and gave a mixture of 2,4- and 2,5-isomers [87].

Unlike pyrroles, furan, thiophene and selenophene react with CF₃SCI only in the presence of catalysts. For selenophene [84] and thiophenes [85] SnCl₄ is sufficient, whilst furans require more forcing conditions usually involving prolonged heating.
(20 h at 60 °C) and in pyridine for activation [83,84] (Scheme 17).

Similarly, some five membered heterocycles with two heteroatoms [N-Ac- and N-(SO₂Alk)-thiazoles, 1-Me-2-SCH₂CF₃- and 1,2-Me₂-imidazoles] undergo single trifluoromethylsulfanylation on heating (60 °C) with CF₃SCl in a pyridine-chloroform mixture [83]. Interestingly, unlike 1,2-dimethylimidazole, the sulfanylation of 2,4-dimethylthiazole under the same conditions occurs twice on the same 2-methyl group (Scheme 18).

Pyridine is too deactivated for trifluoromethylsulfanylation under classical conditions and to achieve substitution it is first of all necessary to convert pyridine to an anionic hydride σ-complex by reduction with LiAlH₄ [86]. The reaction with CF₃SCl then proceeds with difficulty [84] and mono-substituted 3-trifluoromethylsulfanyl pyridine is formed in low yield along with small amounts of the 3,5-bis(SCF₃) derivative (~1%) (Scheme 19).

Introduction of additional RsS-groups into heterocyclic compounds (except for pyrrole and its derivatives) occurs in the presence of perfluoroalkanesulfonic acids (Scheme 20). Incorporation of the second fluoroalkylsulfanyl group into thiophenes [85] and selenophene [84] is possible in the presence of CF₃SO₃H. However, reaction of CF₃SCl with 2,5-bis(SCF₃) thiophene in presence of CF₃SO₃H gives the 3-chloro-derivative as the major product. 2,3,5-Tris(SCF₃) thiophene is accessible if CF₃SO₃H is added as its Ag-salt [77]. Such reactions can also be successfully carried out on pyrroles (Scheme 21).
Prolonged reaction times lead to chlorinated products as well as products that arise from migration of the CF$_3$S-groups (Scheme 22).

Thus, the reaction of perfluoroalkanesulfenyl chlorides with electron-rich aromatic and heterocyclic compounds offers an effective and comparatively straightforward method for the introduction of one or more SR$_F$ groups. The reactions are more problematic however, for electron deficient substrates where competing halogenation, reduction and isomerization products often result from perfluoroalkylthiolation reactions.

### 3.2. Nucleophilic introduction of SR$_F$ groups

Anionic salts of type R$_F$S$^-$/M$^+$ and their heavy metal complexes have been known for many years [88], however their application to the synthesis of aromatic perfluoroalkyl sulfides is comparatively recent. For example, trifluoromethylthiomercuroand trifluoromethylthiosilver react with aliphatic halogenides to generate aliphatic and benzylic trifluoromethyl sulfides [89-92].

It is well known that the reaction of non-activated aryl halides with phenols, thiophenols and amines are catalyzed effectively by copper (Ullmann reaction). L. M. Yagupol’skii [93-97] developed a related protocol for trifluoromethylsulfanylation of aromatic and heterocyclic compounds using trifluoromethylthiocopper (Scheme 23).

The reaction is carried out by heating in a polar solvent (e.g. DMF, quinoline or N-methyl pyrrolidone) and the substrate can...
contain electron-donating or electron-withdrawing groups. Electron-withdrawing groups activate the iodo atom and consequently, give better yields (70–75%). 2-Trifluoromethylsulfanylpyridine, 6-trifluoromethylsulfanylquinoline [93] and 1-trifluoromethylsulfanylnapthalene [97] are obtained in good yields (60–70%) by this method. Multiple aromatic iodine substituents result in multiple substitution by SCF₃ (Scheme 24).

![Scheme 24: Reaction of aromatic iodides with RₓZCu (Z = S, Se), Rₓ = CF₃, C₆F₅ [93,95,96].](image)

In the cases of triiodo derivatives, the yields generally do not exceed 30%. Thus, the synthesis of 1,3,5-tris(SCF₃)benzene is more efficient via 3,5-bis(SCF₃)-iodobenzene [93]. Hexaiodobenzene reacts with CuSCF₃ to form hexakis(trifluoromethylsulfanyl)benzene in modest yield (41%). However, with CuSC₆F₅ and CuSeCF₃ the corresponding hexa-substituted thio- and seleno-derivatives are obtained in yields of 70–90% [96].

![Scheme 25: Side reactions during trifluoromethylsulfanylation of aromatic iodides with CF₃SCu [98].](image)

It should be noted that the interaction of CuSCF₃ with aromatic iodides is sometimes accompanied by side-reactions. For example, the introduction of CF₃S groups into 2,6-diodo-4-nitrochlorobenzene and 2,6-diodo-4-nitroanisole involve simultaneous reduction and substitution (Scheme 25).

Trifluoromethylthiocopper is obtained by reaction of CuBr with AgSCF₃ [93,99], the latter is generated from silver fluoride and carbon disulfide [90,100]. To simplify the process, Remy [101,102] suggested carrying out the synthesis of aryltrifluoromethyl sulfides by generation CuSCF₃ (from trifluoromethylthio mercury and -copper) in situ with the aryl halides. This not only reduces the number of steps but also increases the overall efficiency (Scheme 26).

Aryl bromides can also be used but require higher temperatures (150–190 °C) and more polar solvents. Under such forcing conditions compounds containing both electron-withdrawing and electron-donating groups can now be used effectively. In the case of p-bromo-Ν,Ν-dimethylaniline an excess (3 equiv) of the reagent was used. Aromatic chlorides do not react under...
A variety of perfluoroalkyl- and perfluoroarylcopper mercaptides and selenides have become more accessible, prepared by cleavage of the corresponding disulfides and diselenides with copper powder [94]. The resultant R<sub>f</sub>ZCu reagents complexed with DMF or N-methylpyrrolidone are quite stable and can be stored without decomposition, can be used for the production of aryltrifluoromethyl-, arylpentafluorophenyl sulfides and -selenides from the corresponding iodobenzenes (Scheme 28) [94].

![Scheme 26: Reactions with in situ generated CuSCF<sub>3</sub>](image1)

these conditions. Thus, this method allows the selective substitution of different halogens by varying the temperature.

Since the original work on trifluoromethylthiocopper and trifluoromethylthiomercury [93, 95, 96, 101, 102], other nucleophilic reagents and new methods have been developed. For example, Clark et al. have used CuSCF<sub>3</sub> adsorbed onto Al<sub>2</sub>O<sub>3</sub> [100], whilst Munavalli et al. have employed the acetonitrile adduct CF<sub>3</sub>SCu·CH<sub>3</sub>CN [103] for the reaction with m-iodobenzoic acid and its methyl ester [104].

Bulky perfluoroalkylthiocopper reagents, derived from 2,2,4,4-tetrakis(CF<sub>3</sub>)-1,3-dithietane, hexafluoropropene and alcohols in the presence of KF or CuBr, have been also used for reaction with substituted iodobenzenes (Scheme 27).

![Scheme 27: Perfluoroalkylthiolation of aryl iodides with bulky R<sub>f</sub>SCu](image2)

The compounds shown in Figure 6 have been synthesized by this method.

An alternative approach for the generation of CF<sub>3</sub>SCu involves heating of methyl fluorosulfonyl difluoroacetate in polar aprotic solvents to generate difluorocarbene, which in the presence of CuI and sulfur, forms trifluoromethylthiocopper [106]. Subsequent reaction with aryl halides results in the corresponding trifluoromethylsulfanyl derivatives (Scheme 29).

![Scheme 28: In situ formation and reaction of R<sub>f</sub>ZCu with aryl iodides](image3)
Reduction of bis(perfluoroalkyl)disulfides with tetrakis(dimethylamino)ethylene produces tetrakis(dimethylamino)ethylene dication stabilized perfluoroalkyl thiolates. In contrast to the corresponding potassium and tetramethylammonium salts [29], this compound is stable and can be isolated in a pure state [107], and reacts with activated aryl halides to form the corresponding trifluoromethyl sulfides often in quantitative yields (Scheme 30).

Dmowski and Haas used the reaction of thiocarbonyl difluoride with metal fluorides, to generate the trifluoromethylthiolate anion [108] for introduction into activated perfluoroheterocyclic compounds. Thus, reaction of CF₂=S/CsF with pentafluoropyridine under mild conditions gave the 4-substituted product [109]. However, for the subsequent introduction of additional SCF₃ groups this system is not suitable due to effective self-condensation of thiocarbonyl difluoride (CF₂=S) at higher concentrations. For this purpose the trimer of thiocarbonyl difluoride, bis(trifluoromethyl)trithiocarbonate (CF₃S)₂C=S, is more stable and reacts with CsF in sulfolane to generate CF₃S⁻ anions [110]. However, the use of this reagent leads to mixtures of products (Scheme 31).

Whilst reaction of CF₂=S/CsF (or its trimer) with tetrafluoropyridazine allows for the selective formation of mono-, di- and tri-(SCF₃) substituted products, the analogous reaction with tetra-
fluoropyrimidine results in a mixture of polyfluoropyrimidine derivatives [111] (Scheme 31). Interestingly, the reaction of \((\text{CF}_3\text{S})_2\text{C}=\text{S}/\text{CsF}\) with \(\text{C},\text{N}\)-bis(pentafluorophenyl) imidoyl chloride leads to introduction of the \(\text{SCF}_3\) group into the pentafluorophenyl ring along with substitution of the imidoylic chlorine atom [112].

A considerable improvement of this method was developed by Clark et al. [113]: No preliminary preparation of difluorothio phosphogene or its trimer is necessary, the required reagents being generated in situ (from thiophosgene and KF). The reaction with activated aromatic compounds is shown in Scheme 32.

The less reactive \(2\text{-Cl-5-NO}_2\text{benzonitrile}\) forms the \(\text{CF}_3\text{S}^-\) derivative in only 49% yield after many hours reflux and \(2\text{-F-5-NO}_2\text{benzonitrile}\) is a by-product despite the use of a 100% excess of thiophosgene.

The use of \(\text{Me}_4\text{NF}\) in place of KF for the generation of the \(\text{CF}_3\text{S}^-\) anion in reactions with \(2,4\text{-dinitrofluorobenzene}\) and pentafluoropyridine increases the yields of the corresponding trifluoromethyl sulfides to 90–96% [29,114]. However, with other substrates this method can be problematic due to competing side reactions.

A new method for the preparation of trifluoromethylthiolate anion involves the reaction of \(\text{Me}_3\text{SiCF}_3\) with sulfur in the presence of a fluoride ion source [115]. The salts obtained by this method are considerably more thermally stable than those previously reported [29,110,114]. They can be treated with boiling ether or \(\text{CS}_2\) to remove excess sulfur and readily react at room temperature with inorganic, aliphatic and activated aromatic halides with the formation of trifluoromethyl sulfides (Scheme 33).

It has already been noted that trifluoromethylthiomercury and trifluoromethylthiosilver cannot be used for the preparation of aryltrifluoromethyl sulfides, as they react only with aliphatic halides [89-92]. However, it is known [116,117], that \(\text{Hg(SCF}_3\text{)}_2\) forms a complex with KI which decomposes with the formation of an unstable anion “∗SCF₃”. Based on this observation, Adams and Clark used a mixture of trifluoromethylthiosilver and KI (or \(\text{Bu}_4\text{NI}\)) as a source of trifluoromethylthiolate anion for nucleophilic introduction of the trifluoromethylsulfanyl moiety into aromatic molecules [118]. Of the metal halides investigated for this reaction, the best results were obtained with KI and \(\text{Bu}_4\text{NI}\), whilst NaI, NaBr, and KF were ineffective. Some of these reactions are illustrated in Scheme 34.

This reagent can displace a range of activated halides, particularly bromides and iodides. For the reaction of 2,4-
(NO₂)₂C₆H₃X with AgSCF₃/KI, the reactivity of the halogens occurs in the reverse sequence: F (26%) < Cl (52%) < Br (85%) < I (97%) [118]. Presumably, coordination of the complex anionic nucleophile K⁺[Ag(SCF₃)I]⁻ with aryl halide accelerates the reaction.

Trifluoromethylthiocopper and trifluoromethylthiomercury also participate in analogous reactions, CuSCF₃ is less active than AgSCF₃ whilst Hg(SCF₃)₂ displays increased reactivity as indicated in Scheme 35 [118].

It should be noted that the tellurium reagent, Me₃SnTeCF₃, is capable of introducing the TeCF₃ group into activated heteroaromatics [119]. In the reaction shown (Scheme 36) the use of three equivalents resulted in the introduction of only two TeCF₃ groups.

The Sandmeyer reaction is used widely to introduce functionality into aromatic compounds. However, early attempts using trifluoromethylthiosilver as the nucleophile were not encouraging [120] with yields below 30% accompanied with deammoniated side products (up to 38%). The use of trifluoromethylthiocopper was rather unsuccessful. However, with diazonium salts generated with tert-butyl nitrite in acetonitrile in the presence of CuSCF₃ and BF₃ better results were obtained [121]. Yields of the resulting aryltrifluoromethyl sulfides improved (~40–70%). The best results were observed with isolated tetrafluoroborate diazonium salts (Scheme 37), although the presence of electron-donating and bulky ortho-substituents in the aromatic ring led to reduced yields.

4. Perfluoroalkylation of aromatic sulfur compounds

Perfluoroalkyl iodides have not generally been considered as alkylating agents. Unlike R-X they show anomalous behavior in their reactions with nucleophiles. For example, the reaction of CF₃I with alkali gives fluoroform (CHF₃) and potassium hypoiodide (KIO) [122]. The interaction of organolithium compounds with perfluoroalkyl iodides [123-126] does not result in combination of the two alkyl species (R₃F and R), but in transmetallation (Scheme 38).

Such reactivity has been explained by the reverse polarization of the C–I bond in the fluorinated substrates. Because of the greater electronegativity of CF₃ over iodine (3.3 for CF₃ and
fluoroalkyl sulfides, -selenides and -tellurides. The original under UV irradiation with formation of corresponding per-
[143-146], and with seleno- [147] and tellurophenols [148] react readily with aliphatic, aromatic and heterocyclic thiols under UV irradiation, it is probable that they are also able to react with thiolate anions by a similar mechanism. Indeed, they react readily with aliphatic, aromatic and heterocyclic thiols [143-146], and with seleno- [147] and tellurophenols [148] under UV irradiation with formation of corresponding perfluoroalkyl sulfides, -selenides and -tellurides. The original method required liquid ammonia as the solvent and Pyrex glassware. Thiophenol and its derivatives containing both, electron-donating and electron-withdrawing substituents are easily transformed to the corresponding arylperfluoroalkyl sulfides in high yields (Table 1).$$a_o, o$-Diiodoperfluoroalkanes react at both reaction centers with the formation of bis(SAr)-derivatives containing perfluoroalkylene bridges [144,146] in yields of 80–96%.

With the exception of 4-nitrothiophenol, the reactions are independent of the type of substituents. Unlike many thiophenoxides which bear electron-withdrawing substituents ($p$-Cl, 2,4-Cl$_2$, $o$-SO$_2$CH$_2$ and even $p$-SO$_2$CF$_3$), sodium 4-nitrothiopheno-xide affords 4,4'-dinitrodiphenyl disulfide under these conditions. Conversion to 4-nitrophenyl trifluoromethyl sulfide (60% yield) requires prolonged irradiation in a quartz ampoule at 30–45 °C [143]. The length of the perfluoroalkyl iodide chain has no influence, although lower yields were observed using CF$_3$I in comparison with other iodoperfluoroalkanes. A branching R$_2$I chain results in lower yields of the corresponding sulfides (10–15%). In the case of tertiary perfluoro-butyli iodide, thiophenol are quantitatively transformed into diaryl disulfides. Such behavior of branched perfluoroalkyl iodides can be explained by the facile generation of the $I^*$ radical both as a consequence of their homolytic decomposition [155] and the decomposition of in situ generated radical anions [156]: $i$-R$_2$I$^\bullet$ $\rightarrow$ $i$-R$_2$I$^-$. The radical $I^*$ (or I$_2$) oxidizes the Ar$S^-$ anion to disulfide.

Diaryl disulfides may also be used as substrates. Although they can be trifluoromethylated directly [157], unlike dialkyl disulfides [130,131] the yields generally do not exceed 40% (except for nitro derivatives 4-NO$_2$, 2-NO$_2$–72%). The preliminary breaking of the S–S bond can be carried out very mildly and selectively [9], without affecting other functional groups (Scheme 39).

Perfluoroalkylthioanilines are accessible in a one-pot perfluoroalkylation reaction of dinitrodiphenyl disulfides [158,159] (Scheme 40). This method gives good yields of the desired products, higher than those from the perfluoroalkylation of amino thiophenols. Seleno- [147] and telluro phenols [148] also react with perfluoroalkyl iodides under UV irradiation. Subsequently, it was shown that ArSeNa and ArTeNa react with perfluoroalkyl halides without irradiation to generate R$_2$$S^-$ radicals which react with olefins [160,161]. Irradiation of polymercapto derivatives of benzene and CF$_3$I in liquid ammonia gives poly(trifluoro-methylsulfanyl) compounds in high yields (Table 2).

![Scheme 38: Reactions of perfluoroalkyl iodides with alkali and organolithium reagents](image_url)

2.5 for the atom of iodine [127,128]), the iodine acquires a partial positive charge:

$$\delta- \cdot \delta+ \cdot \delta-$$
$$F_3C-I \quad H_3C-I$$

Nevertheless, Haszeldine et al., were able to carry out the perfluoroalkylations of alkylthiols. Prolonged heating of polyfluoroalkyl iodides with the sodium methylthiolate at 100–110 °C in DMSO lead to the formation of methyl polyfluoroalkyl sulfides [129]. The halophilic generated carbanion (R$_F^-$) in turn reacted with the sulfonyl iodide to generate a thioether. However, R$_F$CH$_3$ and R$_F$H, are also obtained as by-products, which may be a result of homolytic decomposition of the perfluoroalkyl iodides at high temperature [130,131]. Similarly, reactions of R$_F$I with sodium thiophenoxide (like other aromatics such as halogenated benzenes [132] or aromatic heterocycles [133]) resulted in the introduction of the perfluoroalkyl radical into aromatic rings with the formation of a mixture of isomer R$_F$-compounds.

4.1. Ion-radical perfluoroalkylation

4.1.1. Interaction of S-, Se- and Te-phenols, and diaryl disulfides with perfluoroalkyl iodides in liquid ammonia under UV irradiation

Kornblum’s work on nucleophilic substitution in alkyl halides [134-137] and Bunnett’s reactions with non-activated aromatic substrates [138-142] (under UV irradiation) introduced the concept of the nucleophilic radical substitution mechanism (S$_{RN1}$). The essence of this approach consists of the generation of the anionic radical RHg$^\bullet$, its decomposition to a radical R$^*$ (Alk$^*$ or Ar$^*$) followed by reaction with a nucleophile.

Although perfluoroalkyl iodides have a reversed polarity, and in spite of their tendency to undergo homolytic decomposition under UV irradiation, it is probable that they are also able to react with thiolate anions by a similar mechanism. Indeed, they react readily with aliphatic, aromatic and heterocyclic thiols [143-146], and with seleno- [147] and tellurophenols [148] under UV irradiation with formation of corresponding perfluoroalkyl sulfides, -selenides and -tellurides. The original
### Table 1: Interaction of thiophenols with perfluoroalkyl iodides in liquid ammonia under UV irradiation.

![Diagram of reaction](image)

| R        | R_F          | Yields of ArSR_F, % | Ref. |
|----------|--------------|---------------------|------|
| H        | CF_3         | 76                  | [143]|
| 4-NH_2   | CF_3, n-C_3F_7, iso-C_3F_7 | 84, 81, 76          | [144]|
| 2-NH_2   | CF_3         | 71                  | [143]|
| 4-OH     | CF_3         | 69.5                | [143]|
| 2-OCH_3  | CF_3         | 86                  | [98] |
| 4-Cl     | CF_3         | 72                  | [146]|
| 2-SO_2CHF_2 | CF_3   | 69                 | [143,146]|
| 4-SO_2CF_3 | CF_3   | 78                 | [143,146]|
| 4-NO_2   | CF_3         | 2.7^a               | [143,146]|
| 2,4-Cl_2 | CF_3, C_3F_7 | 87                  | [149]|
| 2-COOH   | CF_3         | 89                  | [149]|
| 3- and 4-COOCH_3 | CF_3, n-C_3F_7, iso-C_3F_7 | 70–80          | [151]|
| 3- and 4-F | CF_3, n-C_3F_7 | 80–90            | [152]|
| 4-NHCOCH_3 | CF_3   | 96                  | [153]|
| 4-NHCOOCH_3 | CF_3, n-C_3F_7 | 88 (92^c), 82 (93^c) | [9]  |
|           | C_2F_5, C_4F_9 | 62, 55            | [154]|

^a In a quartz flask.
^b In a quartz ampoule at 30–45 °C.
^c With preliminary reduction of 4,4′-bis(MeOCONH)diaryl disulfide and without the isolation of corresponding thiophenol.

However, the reaction of 2,4,6-trimercaptochlorobenzene with CF_3I generates a mixture of compounds A, B, C and D as illustrated in Scheme 41. Reducing the irradiation time from 30 to 5 min does not change the product composition.

Control experiments indicate that aniline (B) is not derived from either chloro- (A) and iodo- (C)-sulfides, and iodo-product (C) is not formed from chlorosulfide (A). It is known [164] that photochemical nucleophilic aromatic substitution is promoted by electron-donating groups. Therefore, it appears most likely that the sulfides (B), (C) and (D) are produced as a consequence of loss of chloride from the intermediate radical anion as shown in Scheme 42.

Such side reactions explain the decrease of trifluoromethylation efficiency with the number of thiol groups present in a series of thiolated chlorobenzenes. The yields are 72% for 4-SH- [146], 64% for 2,4-(SH)_2- [143] and 37% for 2,4,6-(SH)_3- [163].

![Scheme 39: Perfluoroalkylation with preliminary breaking of the disulfide bond](image)
4.1.2. Perfluoroalkylation of heterocyclic thiols

Heterocyclic thiol form S-perfluoroalkyl derivatives when irradiated in liquid ammonia in the presence of iodoperfluoralkanes. The type of heterocyclic ring and the position of the thiol group influences the reaction. More electron-deficient heterocycles require longer irradiation times (Table 3).

It appears that 4-hydroxypyrimidine-2-thiol does not react with CF$_3$I under standard conditions. Similar to the reaction of 4-nitrothiophenol noted above [143,146], this reaction requires more forcing conditions. Other 4-hydroxypyrimidine-2-thiols behave similarly. The irradiation of an ammoniacal solution of 2-mercapto-4-oxy-6-trifluoromethylpyrimidine with CF$_3$I must be conducted in a Pyrex ampoule at 30–45 °C to produce the S-trifluoromethyl derivative (Scheme 43).

Table 2: UV irradiation of polymeric apt benzences with CF$_3$I in liquid NH$_3$.

| R   | Position of (SH)$_n$ and (SCF$_3$)$_n$ | Yield | Ref. |
|-----|----------------------------------|-------|------|
| Cl  | 2,4-                             | 64%   | [143]|
| COOH | 3,5-                             | 89%   | [162]|
| CH$_3$ | 2,4,6-                       | 90%   | [163]|
| NH$_2$ | 2,4,6-                       | 88%   | [163]|
| OH  | 2,4,6-                           | 69%   | [163]|

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| Cl  | 2,4-                             | 64%   | [143]|
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| CH$_3$ | 2,4,6-                       | 90%   | [163]|
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Scheme 43: Trifluoromethylation of 2-mercapto-4-hydroxy-6-trifluoromethylpyrimidine [145].

Apparently, the reaction of these hydroxymercapto heterocyclic derivatives is complicated by stabilization of sulfur centred radicals as illustrated in Scheme 44.

Scheme 43: Trifluoromethylation of 2-mercapto-4-hydroxy-6-trifluoromethylpyrimidine [145].

4.1.2. Perfluoroalkylation of heterocyclic thiols

Heterocyclic thiol form S-perfluoroalkyl derivatives when irradiated in liquid ammonia in the presence of iodoperfluoralkanes. The type of heterocyclic ring and the position of the thiol group influences the reaction. More electron-deficient heterocycles require longer irradiation times (Table 3).

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Table 2: UV irradiation of polymeric apt benzences with CF$_3$I in liquid NH$_3$.

| R   | Position of (SH)$_n$ and (SCF$_3$)$_n$ | Yield | Ref. |
|-----|----------------------------------|-------|------|
| Cl  | 2,4-                             | 64%   | [143]|
| COOH | 3,5-                             | 89%   | [162]|
| CH$_3$ | 2,4,6-                       | 90%   | [163]|
| NH$_2$ | 2,4,6-                       | 88%   | [163]|
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Table 2: UV irradiation of polymeric apt benzences with CF$_3$I in liquid NH$_3$.

| R   | Position of (SH)$_n$ and (SCF$_3$)$_n$ | Yield | Ref. |
|-----|----------------------------------|-------|------|
| Cl  | 2,4-                             | 64%   | [143]|
| COOH | 3,5-                             | 89%   | [162]|
| CH$_3$ | 2,4,6-                       | 90%   | [163]|
| NH$_2$ | 2,4,6-                       | 88%   | [163]|
| OH  | 2,4,6-                           | 69%   | [163]|

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Table 2: UV irradiation of polymeric apt benzences with CF$_3$I in liquid NH$_3$.

| R   | Position of (SH)$_n$ and (SCF$_3$)$_n$ | Yield | Ref. |
|-----|----------------------------------|-------|------|
| Cl  | 2,4-                             | 64%   | [143]|
| COOH | 3,5-                             | 89%   | [162]|
| CH$_3$ | 2,4,6-                       | 90%   | [163]|
| NH$_2$ | 2,4,6-                       | 88%   | [163]|
| OH  | 2,4,6-                           | 69%   | [163]|

It appears that 4-hydroxypyrimidine-2-thiol does not react with CF$_3$I under standard conditions. Similar to the reaction of 4-nitrothiophenol noted above [143,146], this reaction requires more forcing conditions. Other 4-hydroxypyrimidine-2-thiols behave similarly. The irradiation of an ammoniacal solution of 2-mercapto-4-oxy-6-trifluoromethylpyrimidine with CF$_3$I must be conducted in a Pyrex ampoule at 30–45 °C to produce the S-trifluoromethyl derivative (Scheme 43).

Table 2: UV irradiation of polymeric apt benzences with CF$_3$I in liquid NH$_3$.
Table 3: S-Perfluoroalkylation of heterocyclic compounds under UV irradiation of heterocycles thiol R-Het-SH in liquid ammonia.

| R      | R$_F$ | Reaction conditions                        | Yield of products, % | Ref. |
|--------|-------|--------------------------------------------|----------------------|------|
|        |       |                                           |                      |      |
| 2-(SCF$_3$)$_2$-Benzothiazole |       |                                           |                      |      |
| H      | CF$_3$ | −60 to −33 °C, 90 min                      | 87.5                 | [143]|
|        |       |                                           |                      |      |
|        |       | 2-(SR$_F$)$_2$-Benimidazoles               |                      |      |
| H      | CF$_3$ | −60 to −33 °C, 4 h                         | 51                   | [165]|
| 5-Cl   |       | Pyrex ampoule, 30 °C, 5 h                  | 63–80                | [154]|
|        | C$_2$F$_5$ | liquid NH$_3$, THF, 10 h                | 56                   | [166]|
| 5-(SR$_F$)$_2$-Benimidazoles$^a$ |       |                                           |                      |      |
| 2-Bu   | CF$_3$ | liquid NH$_3$, ampoule, 25–40 °C, 10 h     | 20–39                | [154]|
|        | C$_3$F$_7$ |                                           |                      |      |
| 5-(SR$_F$)$_2$-6-Azauracil |       |                                           |                      |      |
| H      | CF$_3$ | −33 °C, 45 min                             | 77                   | [10] |
|        | C$_3$F$_7$ |                                           | 76                   |      |
| 2-(SCF$_3$)$_2$-Pyrimidines  |       |                                           |                      |      |
| 4,6-(CH$_3$)$_2$ | CF$_3$ | −33 °C, 60 min                             | 82                   | [154]|
| 4-SH   | CF$_3$ |                                           | 61$^b$               | [154]|
| 4-SH-6-CH$_3$ | CF$_3$ |                                           | 58$^b$               | [154]|
| 4-OH-6-CF$_3$ | CF$_3$ | Pyrex ampoule, 30–45 °C, 5 h               | 59                   | [154]|
| 4,6-Me$_2$-5-OH | CF$_3$ | −30 °C, 4 h                                | 89                   | [154]|

$^a$Received from 5-SZn salts, poorly soluble in liquid ammonia.

$^b$The 2,4-bis(SCF$_3$)$_2$-derivatives.

In the case of 2-mercapto-5-hydroxy-4-hydroxypyrimidines, no tautomeric keto form such as that shown in Scheme 44 is possible and consequently, are perfluoroalkylated without any problems, e.g., 2-mercapto-5-hydroxy-4,6-dimethyl pyrimidine [145].

In summary, heterocyclic thiols react with perfluoroalkyl iodides with considerably more difficulty than aromatic thiols.

4.1.3. Photochemical perfluoroalkylation in organic solvents under phase transfer conditions

Liquid NH$_3$ is a key reaction medium for the reaction of organic thiols with perfluoroalkyl iodides under UV irradiation. However, other solvents have been investigated including alcohols, acetone, acetonitrile, dioxane, THF, DMF, DMSO, HMPA and so on. Polar aprotic solvents emerge as the best. Biphasic reactions with water work well, particularly with diethyl ether and benzene (Table 4).

Heterocyclic thiolates react more slowly with perfluoroalkyl iodides than thiophenoxides both in liquid ammonia and in organic solvents. Besides, in reactions with heterocyclic thiolates, as well as with thiophenoxides, CF$_3$I is a poorer electrophile than C$_3$F$_7$I - even under biphasic conditions.

4.1.4. Interaction of thiols with perfluoroalkyl bromides

Although brominated perfluoroalkanes are cheaper and more readily available than the corresponding iodides, they react more slowly in thioether forming reactions. In general, monobrominated perfluoroalkanes do not react. However, dibromodifluoromethane, bromochlorodifluoromethane as well as 1,2-dibromotetrafluoroethane [170,171] do react with metal phenoxides and thiophenoxides via halophilic mechanisms [64], and almost always lead to mixtures of bromo and chloro containing products of mono- and di-substitution.
Trifluoromethyl sulfides. Reactions between thiophenoxides and

Wakselman et al., have shown [176] that liquid C–Br bond is lower than C–I bond (55 kcal/mol for CF₃Br) compared to C–I (28 kcal/mol for CF₃I) [172]. In addition, CF₃Br has a higher dissociation energy than CF₃I and prefers to react with thiolate without any irradiation, whereas bubbling gaseous CF₃Br through a DMF solution of thiolates at 20 °C or heating such mixtures in an autoclave (80 °C) does not produce trifluoromethyl sulfide in low yield (~3–5%), although better yields are obtained when iodide salts are used as catalysts [175].

Nevertheless, it was found [174] that UV irradiation of thiophenols in liquid ammonia or dimethylformamide with perfluoroalkyl bromides does result in the formation of the corresponding perfluoroalkyl sulfides as shown in Scheme 45.

Thiols with electron-donating substituents give reasonable yields, whilst p-chlorothiophenol produces the corresponding trifluoromethyl sulfide in low yield (~3–5%), although better yields are obtained when iodide salts are used as catalysts [175].

Wakselman et al., have shown [176] that liquid CF₃F₁₃Br reacts with thiolates without any irradiation, whereas bubbling gaseous CF₃Br through a DMF solution of thiolates at 20 °C or heating such mixtures in an autoclave (80 °C) does not produce trifluoromethyl sulfides. Reactions between thiophenoxides and

Table 4: Reaction of thiophenols RC₆H₅SH and mercapto heterocycles with RpI under UV irradiation in organic solvents and biphasic conditions.

| R    | Rp    | Base     | Solvent | Conditions               | Yields of ArSR₉, % | Ref. |
|------|-------|----------|---------|-------------------------|-------------------|------|
|      |       |          |         |                         |                   |      |
| H    | CF₃   | PhSNa    | CH₃OH or acetone | 0–5 °C, 30 min | 57.5 or 79 | [143] |
|      |       | NaOH     | CH₃CN   | 0–5 °C, 30 min | 43 or 49 | [143] |
| CF(CF₃)₂ | Et₂N  | CH₃CN   | 0 °C, 30 min | 88 | [104] |
| CF₃  | NaOH  | Et₂O/H₂O | (Et₂)BzN⁺Cl⁻, 20–25 °C, 30 min | 54 | [167] |
| C₆F₁₃ |       |          |         |                         |                   |      |
| 4-Cl | CF₃   | ArSNa    | CH₃OH or CH₃CN | 20 °C, 30 min | 61 or 81 | [144] |
|      |       | NaOH     | Et₂O/H₂O | (Et₂)BzN⁺Cl⁻, 20–25 °C, 30 min | 61 | [167] |
|      |       | i-C₆F₁₃  |          |                         |                   |      |
|      |       | C₂F₇     |          |                         |                   |      |
| 4-CH₃| CF₃, C₆F₁₃ | NaOH | Et₂O/H₂O | (Et₂)BzN⁺Cl⁻, 20–25 °C, 30 min | 58, 83 | [167] |
|      |       | C₂F₇     |          |                         | 67 | [167] |
|      |       | i-C₆F₁₃  |          |                         | 60 | [167] |
|      |       | C₆F₁₃    |          |                         | 71 | [167] |
| 4-OCH₃| CF₃  | NaOH     | Et₂O/H₂O |             | 52 | [167] |
| 4-CO₂CH₃| CF₃  | NaOH     | Et₂O/H₂O |             | 71 | [167] |
| 4-NH₂| CF₃   | NH₂OH    | NH₂OH   | −60 to 25 °C | 95 | [168] |

2-Mercapto heterocycles

| Heterocycle | Rp    | Base | Solvent | Conditions | Yield | Ref. |
|-------------|-------|------|---------|------------|-------|------|
| Benzothiazole | Cl(CF₂)₄ | NaH | DMF | 70 °C, 10 h | 41.2 | [169] |
|             | Cl(CF₂)₆ | NaH | DMF | 70 °C, 10 h | 61.6[^d] | [169] |
|             | C₆F₁₃   | NaH | DMF | 70 °C, 10 h | 53.6 | [169] |
|             | C₆F₁₇   | NaH | DMF | 71.6 | [169] |
| Benzimidazole | Cl(CF₂)₄ | NaH | DMF | 70 °C, 10 h | 40.6[^d] | [169] |
|              | Cl(CF₂)₆ | NaH | DMF | 70 °C, 10 h | 38.2 | [169] |
|              | C₆F₁₃, C₆F₁₇ | NaH | DMF | 70 °C, 10 h | 77.6, 78.2 | [169] |
| Benzoxazole | Cl(CF₂)₆ | NaH | DMF | 70 °C, 10 h | 15.0 | [169] |

[^a]: At ArSH + Et₂NH or Et₂N for 3 h, the yields are 37% and 28%, respectively.
[^b]: In CH₃Cl/CH₂Cl₂ or CHCl₃/H₂O the yields are 50% and 55%, respectively.
[^c]: Yields of products are tabulated in Table 4, taking into account a conversion of RpI.
[^d]: In presence of (t-Bu)₂N-O' the yield is 8.6%.

Beilstein J. Org. Chem. 2010, 6, 880–921.
**Scheme 45:** Perfluoroalkylation of thiolates with CF$_3$Br under UV irradiation.

CF$_3$Br are successful if carried out under pressure (CF$_3$Br 2–3 atm) in DMF at 20 °C [176-178]. However, even under these conditions only thiols containing electron-donating groups in the para-position give high yields. All ethers (Table 5), even those with electron-donating groups in the ortho- and meta-positions show very poor reactivity.

The best results arise from a combination of two factors – a pressure of CF$_3$Br and UV irradiation [158,179]. Results are given in Tables 6–8. In these cases the influence of the solvent is obvious. For example, p-chlorothiophenol reacts poorly with CF$_3$Br and 4-chloro-4’-trifluoromethylsulfanyldiphenyl sulfide is obtained as a byproduct presumably as the result of photo-substitution of chlorine in 4-trifluoromethylsulfanylchlorobenzene by an S$_{RN}$1 mechanism. HMPA suppressed this side-reaction (similar to iodobenzene with potassium diethyl phosphite [180]) and promoted trifluromethylation (Table 6).

The reaction solvent is important and the yield of the trifluoromethylated product decreases in the following sequence: HMPA > DMF > CH$_3$CN > N-methyl pyrrolidone > sulfolane [179] (Table 6). The efficiency of the combined influence of irradiation and pressure of CF$_3$Br is presented in Table 7.

As can be seen from the data in (Table 6 and Table 7), in spite of increased product yields in general, the selectivity remains

---

**Table 5:** Yields of CF$_3$Br reaction with thiophenoxides in DMF at 20 °C under pressure (2–3 atm) [176].

| Substituents in thiophenol | H | 4-CH$_3$ | 4-OCH$_3$ | 3-OCH$_3$ | 2-OCH$_3$ | 3-NH$_2$ | 4-Cl | 3-CF$_3$ | 4-NHAc |
|---------------------------|---|---------|----------|----------|----------|---------|------|---------|--------|
| Yields of ArSCF$_3$, %    | 62| 75      | 83       | 40       | 7        | 23      | 34   | 13      | 9      |

**Table 6:** Reactions of thiophenoxides with CF$_3$Br under UV irradiation and pressure of reaction gas [179].

| R   | Solvent  | Base       | $p$ (atm) | $T$ (°C) | Irradiation time, (h) | Conversion of ArSH, (%) | Isolated yields of ArSCF$_3$ (%) |
|-----|----------|------------|-----------|----------|-----------------------|-------------------------|----------------------------------|
| 4-CH$_3$ | DMF | Et$_3$N   | 4–5       | 10–13    | 1.5                   | 82                      |                                  |
| 4-NH$_2$ | DMF | Et$_3$N   | 4.5–6     | 10–20    | 2                     | 76.4                    |                                  |
| 3-NH$_2$ | HMPA | morpholine | 3–4       | 17–19    | 3.25                  | 63.5$^a$                |                                  |
| 4-NHCOMe | DMF | Et$_3$N   | 3.5       | 19       | 2.7                   | 69                      |                                  |
| 4-NHCO$_2$Me | DMF | Et$_3$N   | 4.5–5     | 15–25    | 1.2                   | 63                      | 55.5                             |
| 4-NHCO$_2$Me | HMPA | morpholine | 2–5       | 8–10     | 2.5                   | 73                      | 83.6                             |
|       | CH$_3$CN | Et$_3$N   | 3–3.5     | 15–18    | 2.8                   | 53                      | 43$^a$                           |
| 4-Cl  | DMF | Et$_3$N   | 3–3.5     | 14       | 1.2                   | 100                     | 48$^a$                           |
|       | HMPA | Et$_3$N   | 4         | 8–10     | 1                     | 100                     | 69                               |
|       | HMPA | morpholine | 3–4       | 14–16    | 3.5                   | 97                      | 62.5                             |
|       | HMPA | morpholine | 3–4.5     | 29–30    | 3                     | 36                      | 46                               |
|       | Sulfolane | morpholine | 3.5       | 23       | 2                     | 19.5                    | 5.4                              |
|       | N-Methyl pyrrolidone | morpholine | 3.5       | 17       | 2.2                   | 35.5                    | 14.3                             |

$^a$Determined by GLC.
about the same. The best results are found with thiophenols, containing electron-donating substituents in the para-position. It is possible to increase the effectiveness of the p-chlorothiophenol reaction to ~70% by suppression of by-product formation (4-Cl-CF$_3$SCF$_3$-4) and by using HMPA as solvent.

Trifluoromethylation of easily oxidizable aminothiophenols can be conducted by a modified procedure. The required thiophenoxides are prepared directly prior to irradiation by reduction of the corresponding dinitrophenyl disulfides with Li/liquid NH$_3$ (Table 8, in much the same way as the described above for R$_3$I [158,159].

Table 8: Preparation of aminophenyl trifluoromethyl sulfides with CF$_3$Br (3–7 atm) and UV irradiation with preliminary reduction of dinitrophenyl disulfides [179].

| Location of | Solvents | p (atm) | T (°C) | Irradiation time, h | Yields of products, % |
|-------------|----------|---------|--------|---------------------|----------------------|
| NO$_2$      | DMF      | 4.6–6   | 10–13  | 7.75                | 40.9                 |
|             | DMF      | 3–3.5   | 8–10   | 2.2                 | 56$^a$               |
|             | DMF      | 3–6     | 10–14  | 4                    | 72.5$^a$             |
|             | DMF      | 4–6     | 12–19  | 6.8                  | 80.8                 |
|             | HMPA     | 3–5     | 8–10   | 3                    | 71.8$^a$             |
|             | DMF      | 5–6     | 15–20  | 5                    | 80.3                 |

$^a$ Isolated as the acetyl derivative.

Due to greater UV stability of CF$_3$Br compared to CF$_3$I, it is possible to increase the irradiation time, with a beneficial effect on the product yield.

4.1.5. Other methods of initiating

From the knowledge that the reaction mechanism is a single-electron transfer process involving R$_F^+$ radicals, alternative methods to photochemical initiation have been developed (see sections 4.1.1.–4.1.4.), e.g., the electrochemical reduction of perfluoroalkyl halogenides [173,181]. In the presence of thiolate anions the resulting electrophilic radicals react [182,183] to give aryl perfluoroalkyl sulfides (Table 9).

Table 9: Formation of aryl perfluoroalkyl sulfides by electrochemical initiated reactions of ArS$^+$ with R$_F$Hlg.

| ArS$^+$  | R$_F$Hlg | Yield of ArSR$_F$, % |
|----------|----------|----------------------|
| p-CH$_3$C$_6$H$_4$S$^+$ | CF$_3$I | 55  |
| p-CH$_3$C$_6$H$_4$S$^+$ | CF$_3$I | 77  |
| p-CH$_3$C$_6$H$_4$S$^+$ | CF$_3$I | 63  |
| p-CIF$_6$H$_4$I$^+$ | CF$_3$I | 75  |
| p-CIF$_3$H$_4$S$^+$ | CF$_3$I | 61$^b$ |
| p-CIF$_3$H$_4$I$^+$ | CF$_3$I | 82  |
| p-CF$_3$I$^+$ | CF$_3$I | 60  |
| p-CH$_3$OCONHC$_6$H$_4$S$^+$ | CF$_3$I | 33  |
| Thiazole-2-$^-$ | CF$_3$I | 64$^c$ |

$^a$ With a carbon-glass electrode a yield is 77%.

$^b$ With a carbon-glass electrode.

$^c$ In the presence of p-O$_2$NC$_6$H$_4$CN.

The good yields for electrochemical perfluoroalkylation (especially > 100% electrochemical yield) are consistent with a radical-chain process.

Perfluoroalkyl iodides are better substrates than the bromides which give lower yields in these electrochemical reactions (Table 9). Such electrochemically initiated reactions are described in detail in a review [35].

Another method of catalytic generation of R$_F^+$ radicals involves electron-transfer from a nucleophile to a perfluoroalkyl halide, in this case using the dimethyl dipyrinidium salt (methylviologen, MV$^{2+}$) as a catalyst. This dication is initially reduced to a radical cation, which then transfers an electron to a perfluoroalkyl iodide [185] to generate R$_F^+$ (Scheme 46). A small amount of MV$^{2+}$ (7% relative to ArSH) is sufficient for quantitative transformation of thios to aryl perfluoroalkyl sulfides (Table 10).
It should be noted that over-reduction of such halides will generate R_F\(^{-}\) anions rather than the desired R_F\(^{•}\) radicals. For example, tetrakis(dimethylamino)ethylene reacts with R_F\(^{-}\)I to form the perfluoroalkyl anion which acts as a nucleophilic R_F\(^{-}\)-alkylation agent for organic and inorganic substrates [187].

The use of any catalyst in the case of perfluoroalkyl iodides is of more theoretical interest, although the method can be applied in the case of poorly reactive thiophenols. In general these reactions work well (see section 4.1.6.) in common organic solvents or under biphasic conditions [188,189]. Reactions with perfluoroalkyl bromides are more sluggish. Only compounds with long perfluoroalkyl chains such as C_6F_{13}Br [178] react readily with thiolates. In the reaction of gaseous CF_3Br with thiophenols special procedures are required (see section 4.1.4.): UV irradiation [174], pressure [178] and electrochemical stimulation [182]. Moreover, thiophenols with electron-donating substituents in the para-position give the best results. Combined pressure and irradiation [158,179] improved yields only slightly and requires special equipment. A detailed study of catalytic stimulation in reactions of bromo- and chloro-containing freons R_FX with thiols is necessary.

The decreased reactivity of CF_3Br as compared to CF_3I can be explained, first of all, by the higher reduction potential (≈2.07 V against −1.52 V for CF_3I on a glass-carbon cathode), and secondly, by the fact that the CF_3\(^{•}\) radical has a reduction potential (≈1.80 V) close to that of CF_3Br [173]. Thus trifluoromethyl bromide in reactions with nucleophiles or on a cathode surface accepts two electrons and is transformed to CF_3\(^{-}\) and therefore does not react with thiolates. The SO_2\(^{-}\) radical anion can act as an electron mediator in such reactions. This radical anion, generated by chemical [190-193] or electrochemical [194,195] methods, causes a single-electron reduction of CF_3Br with the formation of the necessary trifluoromethyl radical. Thus, the use of SO_2\(^{-}\) sources (Na_2S_2O_4, HOCH_2SO_3Na or SO_2 in presence Zn and Na_2HPO_4 or HCOONa) on trifluoromethyl bromide in DMF in the presence of diaryl disulfides [193,196] leads to the formation of the corresponding trifluoromethyl sulfides, often in high yields (Scheme 47).

Related transformations with various SO_2\(^{-}\) sources involving R_F\(^{-}\)I and CF_3ClBr, CFC_12-CF_2Cl in the reactions with diaryl disulfides [197] and diselenides have been reported [198]. Electrochemical studies involving the SO_2\(^{-}\) radical anion prove that the electron transfer to CF_3Br takes place at a reduction potential of the mediator between −0.9 and −1.0 V which prevents the transfer of a second electron to CF_3\(^{•}\) and the generation of CF_3\(^{-}\) [199]. Therefore electrochemical reduction in the presence of sulfur dioxide allows the trifluoromethylation of thiophenols with the less reactive, but more readily available trifluoromethyl bromide (Scheme 48).

Although 4-nitrothiophenol is a very poor substrate (see section 4.1.1. and Table 11), it reacts with perfluoroalkyl iodides to afford 4-perfluoralkylsulfonylnitrobenzenes in presence of...
NaH in DMF in almost quantitative yields [201], presumably via “hydride” catalysis.

The catalytic influence of SO$_2$ on the reaction of ArS$^-$ with CF$_3$Br is not limited to the activation of the initial bromide. Sulfur dioxide can oxidize the radical anion ArSCF$_3$•, i.e., it can affect the rate determining step of the process [189] (Scheme 49).

![Scheme 49: Participation of SO$_2$ in the oxidation of ArSCF$_3$•.](image)

This dual influence of sulfur dioxide contributes to the overall efficiency of these reactions.

By comparing the possibility of two mediators (SO$_2$ and MV), Koshechko et al., [202] have shown that the radical cation MV$^{+}$ (E$_p$ = -0.4 V) easily reduces SO$_2$ (E$_p$ = -0.9 V) to its radical anion which in turn activates CF$_3$Br. Thus, a combination of both mediators generates an electron transfer cascade (Scheme 50).

![Scheme 50: Electron transfer cascade involving SO$_2$ and MV.](image)

Thus, bubbling CF$_3$Br into a solution of thiophenol or thiocresol in DMF containing pyridine, SO$_2$ and a catalytic amount of MV$^{2+}$, results in the formation of the corresponding aryl trifluoromethyl sulfides in moderate to good yields (40–70%) [202].

Similar reactions with SO$_2$, where KI or I$_2$ were used instead of MV$^{2+}$ have been carried out [202], however, the yields of PhSCF$_3$ were reduced. The catalytic effect of iodide ion was discovered from UV irradiation of a reaction mixture of p-chlorothiophenol with CF$_3$Br in different solvents [175].

The MV$^{2+}$/SO$_2$ system is effective for reactions with Freons, particularly those with C–Cl bonds such as Freon-113 (CF$_2$ClCFCl$_2$) [202].

A good example of the catalytic properties of SO$_2$ has recently been shown in the reaction of 1,2-dibromotetrafluoroethane with thiophenoxides [203]. It is known that these reactions ArSCF$_2$CF$_2$Br and a significant amount of ArSCF$_2$CF$_2$H are produced. The presence of SO$_2$ in the reaction promotes a S$_{N}2$1 process which results in quantitative yields of ArSCF$_2$CF$_2$Br without the byproduct ArSCF$_2$CF$_2$H.

### 4.1.6. Spontaneous perfluoroalkylation of thiols without initiators

Since Feiring reported in 1984 that reactions of thiolate anions and perfluoroalkyl iodides can occur spontaneously without any initiator [188], the method has been extensively investigated and the reaction conditions optimized (Table 11 and Table 12). Reactions times, for example, are shortened with heating (60–70 °C) [204].

Later it was found that these types of reaction can be made to proceed considerably easier and quicker (Table 12). In acetonitrile or DMF the majority of thiophenolates react rapidly with C$_3$F$_7$I at room temperature (from 10–15 min to 2–3 h). However, for spontaneous reaction many factors are involved such as carrying out the reaction in the dark, temperature, solvent etc. This is discussed in more detail in section 4.1.7.

### 4.1.7. Reaction mechanism

The stages of S-perfluoroalkylation [22,35,143,188,208] can be represented as follows (Scheme 51):

![Scheme 51: Four stages of the S$_{N}$1 mechanism for thiol perfluoroalkylation.](image)

The peculiar behavior of 4-nitrothiophenol [143,146] and 4-hydroxy pyrimidine-2-thiol [145] unlike the more electronegative p-SO$_2$CF$_3$- and o-SO$_2$CHF$_2$-thiophenols [143] is presumably related to the ability of the nitro- and carbonyl groups to stabilize the mercapto-radicals in the radical ion pairs ['O$_2$NArS + R$_F$] and ['O=CArS + R$_F$]. As a result, these radicals are less reactive, although at higher temperatures an increase in their activity is observed.

The participation of radicals is supported by the fact that the addition of nitrobenzene [178] or di-tert-butynitroxide [169]
Table 11: Reactions of thiols R(C₈H₄SH) and HetArSH with R₇F in organic solvents and in biphasic conditions without initiators.

| R     | SH, (SCat⁺) | R₇F | Base | Reaction conditions | Yields of ArSR₇F, % | Ref. |
|-------|-------------|-----|------|---------------------|---------------------|------|
|       |             |     |      |                     |                     |      |
|       |             |     |      |                     |                     |      |
| H     | SNa         | C₈F₁₇ | —    | DMF, 25 °C, 17 h    | 90                  | [188]|
| H     | SNa         | C₈F₁₇ | —    | DMF, 25 °C, 17 h + norbornene | 77                  | [188]|
| H     | SNa         | C₈F₁₇ | —    | DMF, 25 °C, 17 h + styrene | 0                   | [188]|
| H     | SNa         | CF₂(CF₃)₂ | —    | DMF, 25 °C, 17 h    | 76                  | [188]|
| H     | SNa         | C₈F₁₃ | —    | CH₂Cl₂/H₂O, 40 °C, 4 h | 48                  | [188]|
| H     | SNa         | C₈F₁₃ | —    | C₆H₆/H₂O, 25 °C, 2.5 h  | 76⁺                 | [188]|
| H     | SNa         | R(CF₂)ₙ | —    | DMF, conditions are not presented | 56–87              | [205]|
| 4-NH₂ | SH          | C₉F₅  | K₂CO₃ | DMF, 10 °C      | 84                  | [206]|
| 4-F   | SNa         | C₈F₂₄ | —    | DMF, 70 °C, 1 h     | 97                  | [204]|
| 4-F   | SNa         | CF₂I₄ | —    | DMF, 25 °C, 12 h, 60 °C, 1 h | 86ᵇ              | [201]|
| 4-Cl  | SNa         | (CF₂)₄I | —    | DMF, 50 °C, 6 h    |                     |      |
| H     | SH          | C₈F₉  | NaH  | DMF, 20–25 °C, 17–18 h | 66                  | [201]|
| 4-CH₃ | SH          | C₈F₉  | NaH  | DMF, 20–25 °C, 17–18 h | 77                  | [201]|
| 4-OH  | SH          | C₈F₉  | NaH  | DMF, 20–25 °C, 17–18 h | 30                  | [201]|
| 4-Cl  | SH          | C₈F₉  | NaH  | DMF, 20–25 °C, 17–18 h | 83                  | [201]|
| 4-NO₂ | SH          | C₄-C₆ | NaH  | DMF, 20–25 °C, 17–18 h | 93–99               | [201]|
| F₅    | SCu         | CF₂=CF | —    | DME, 70 °C, 20 h    | 65                  | [207]|
| F₅    | SCu         | C₈F₁₇ | —    | DME, 70 °C, 20 h    | 0                   | [207]|
| H     | SeNa        | CF₃Br | —    | ETOH, 20 °C, 2 h, olefins | 2–60               | [160]|
| H     | SeNa        | C₄F₃I–C₈F₁₇I | —   | ETOH, 20 °C, 2 h, olefins | 2–60               | [160]|

| Heterocycle | R₇F | Base | Reaction conditions | Yields | Ref. |
|-------------|-----|------|---------------------|--------|------|
| 2-SH-benzothiazole | C₃F₇ | NEt₃ | DMF, 55–60 °C, 3–48 h | Traces | [189]|
|              | C₃F₇ | NEt₃ | DMF, 20–22 °C, 120 h | 59     | [189]|
|              | CI(CF₂)₄–6 | NaH | DMF, 70 °C, 10 h | 0–4.5ᶜ | [169]|
| 2-SH-benzimidazole | CI(CF₂)₄–6 | NaH | DMF, 70 °C, 10 h | 0–3ᵈ | [169]|
| 8-SCu-quinoline | C₃F₇ | NEt₃ | DMF, 20–22 °C, 24 h | 72     | [189]|

|ᵃ| In the presence of norbornene and styrene the yields are 30% and 0%, respectively. |  
|ᵇ| α, ω-Bis(SH)/perfluoroalkanes. |  
|ᶜ| 8.5% conver, RxF. |  
|ᵈ| 3% conver, RxF. |

inhibits the reaction. The addition of olefins such as norbornene or styrene [188] has a similar effect and perfluoroalkyl derivatives of these olefins have been identified in the reaction products. The formation of radicals in the reaction of PhSeNa with perfluoroalkyl halides (PhSe⁺ and R₇F⁻) has been firmly established from their interception by unsaturated compounds [160].

Further confirmation of a radical mechanism was obtained by studying the reaction without an initiator (Table 12 and Table 13). The decrease of reaction temperature, carrying out the reaction in the absence of light, the presence of electron-withdrawing substituents in the thiol ring and use of low-polar solvents all led to lower ArSR₇F yields. Also replacement of C₃F₇I for CF₃I leads to a slower reaction and reduced yields of aryl perfluoroalkyl sulfides. In spite of heptafluoropropyl iodide being a stronger oxidant than CF₃I [182,209], greater amounts of diaryl disulfides are obtained only with CF₃I. The factors listed above influence the yields of diaryl disulfides in a different way. They either do not change (in darkness), or they even slightly increase (from 3–4 to 12–13%).

These observations point towards the rate determining step of the reaction [189]. Two steps (Scheme 51), i.e., the rapid fragmentation of the radical anion R₇F⁻ (Equation 2) [173] and recombination of the electrophilic radical R₇⁺ with the ArS⁻ anion (Equation 3) are fast and cannot therefore be rate limiting.
Table 12: Reaction conditions of thiophenoxides $R_2C_6H_4S^{-}$ $Et_3NH^+$ with $RF$ without irradiation [189].

| Entry | $R$       | $RF$      | Solvent | $T$ (°C) | $t$ (h) | Yields (%) |
|-------|-----------|-----------|---------|---------|--------|------------|
| 1     | H         | $C_3F_7$  | DMF     | 19–20   | 2      | 83         | 3          |
| 2     | 4-NHCO$_2$H$_3$ | $C_3F_7$  | DMF     | 21–22   | 20 min | 89         | 3          |
| 3     | 4-NHCO$_2$H$_3$ | $C_3F_7$  | DMF     | 21–22   | 1      | 60         | 4          |
| 4     | 4-NHCO$_2$H$_3$ | $C_3F_7$  | DMF     | 21–22   | 1      | 70         | 9          |
| 5     | 4-NHCO$_2$H$_3$ | $C_3F_7$  | DMF     | 0–5     | 3      | 17         | 12         |
| 6     | 4-NHCO$_2$H$_3$ | $C_3F_7$  | DMF     | 0–22    | 5      | 30         | 7          |
| 7     | 4-NHCO$_2$H$_3$ | $C_3F_7$  | HMPA    | 0–5     | 3      | 0          | 12         |
| 8     | 4-NHCO$_2$H$_3$ | $C_3F_7$  | HMPA    | 21–22   | 2      | 75         | 3          |
| 9     | 4-NHCO$_2$H$_3$ | $C_3F_7$  | CH$_3$CN | 21–22 | 0.5 | 98         | —          |
| 10    | 4-NHCO$_2$H$_3$ | $C_3F_7$  | dioxane | 21–22   | 2      | 82         | 2          |
| 11    | 4-NHCO$_2$H$_3$ | $C_3F_7$  | THF     | 21–22   | 1.5    | 64         | 10         |
| 12    | 2-NH$_2$  | $C_3F_7$  | CH$_3$CN | 21–30$^b$ | 10 min | 84         | —          |
| 13    | 2-NH$_2$  | $C_3F_7$  | CH$_3$CN | 23–24   | 1      | 66         | 7          |
| 14    | 4-OCH$_3$ | $C_3F_7$  | CH$_3$CN | 22–40$^b$ | 10 min | 88         | 6          |
| 15    | 4-Cl      | $C_3F_7$  | DMF     | 22      | 2      | 72         | 3          |
| 16    | 4-Cl      | $C_3F_7$  | CH$_3$CN | 21–22 | 3      | 40         | 12         |
| 17    | 4-COOH    | $C_3F_7$  | DMF     | 22–30$^b$ | 10 min | 72         | Traces     |
| 18    | 4-COOCH$_3$ | $C_3F_7$  | DMF     | 20      | 3      | 39         | 13         |
| 19    | 4-NO$_2$  | $C_3F_7$  | DMF     | 50–55   | 5      | Traces     | 6          |

$a$ In the dark.

$b$ Spontaneous warming.

$c$ Sodium thiophenoxide.

Since all experimental factors (light, temperature, solvent etc.) have an inverted influence on the yields of disulfides, it can be assumed that Equation 1, the generation of ArS$^-$ is also not limiting. Therefore electron transfer from the radical anion [ArSR$_F$]$^-$, Equation 4, seems to be the most likely.

Homogeneous catalysis by the methyl viologen (MV) [186] supports this. This catalyst can oxidize the radical anion [ArSR$_F$]$^-$ via its dication (MV$^{2+}$) [200,202], accelerating the last step (Scheme 52).

4.2. Radical perfluoroalkylation

Synthetic methods for aryl perfluoroalkyl sulfides via $R_F^-$ radicals are now described. Prolonged UV irradiation of CF$_3$I solutions with diaryl disulfides in liquid ammonia results in the formation of the corresponding aryl trifluoromethyl sulfides (Table 13).

For diaryl disulfides the CF$_3^-$ radical can attack either the sulfur atom or the aromatic ring, [132,210] and thus give rise to undesired side products. Aryl perfluoroalkyl sulfides are formed also in a reverse strategy from aliphatic disulfides and aryl radicals. For example, during irradiation of bis(trifluoromethyl) disul-
The $N$-trifluoromethylnitrososulfonamide of trifluoromethane sulfonic acid reacts similarly with aliphatic disulfides [214]. Interaction of CF$_3$NO with aryl sulfonamides generates relatively stable trifluoromethyl azosulfonyl arenes ArSO$_2$N=NCF$_3$, which decomposed on heating to CF$_3$• radicals which react with organic disulfides to form trifluoromethyl sulfides [215] (Scheme 55).

### Scheme 55: Radical trifluoromethylation of organic disulfides with ArSO$_2$N=NCF$_3$.

Decarboxylation of non-fluorinated carboxylic acid esters proceeds in a similar manner to afford 2-pyridyl sulfides. However, in the presence of C$_6$F$_{13}$I the reaction follows a different course where the perfluorinated radical attacks sulfur with the formation of the fluorinated sulfide [217] (Scheme 57).

Barton has shown [216] that the irradiation of thiohydroxamic esters of perfluorocarboxylic acids generates RF• radicals which in the presence of olefins give addition products. However, in the absence of radical traps they attack the sulfur to yield, for example, S-perfluoroalkyl derivatives of pyridine (Scheme 56).

### Scheme 56: Radical trifluoromethylation of organic disulfides with ArSO$_2$N=NCF$_3$.

N-Trifluoromethyl-$N$-nitrosobenzene sulfonamide has been used as a source of CF$_3$• radicals. This reagent (obtained by reaction of CF$_3$NO, NH$_2$OH and benzenesulfonic acid chloride) reacts with organic disulfides under irradiation or on mild heating to give the corresponding trifluoromethyl sulfides (Scheme 54).

### Scheme 54: $N$-Trifluoromethyl-$N$-nitrosobenzene sulfonamide – a source of CF$_3$• radicals [212,213].

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**Table 13:** UV irradiation of CF$_3$I with diaryl disulfides in a sealed quartz tube [157].

| R  | t (h) | Yield of ArSCF$_3$ (%) |
|----|------|------------------------|
| H  | 12$^a$ | 36.5$^a$ |
| 4-Cl | 5   | 36.5$^a$ |
| 4-NO$_2$ | 25  | 58   |
| 2-NO$_2$ | 12  | 72   |

$^a$Extracted from mixtures.

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The photochemical reaction of pentafluoroiodobenzene with trifluoromethyl disulfide.

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The irradiation of thioesters of trifluoroacetic and trifluoromethanesulfonic acids in refluxing methylene chloride results in their decarbonylation (or desulfonation in the case of CF$_3$SO$_2$SR) with the production of CF$_3$• radicals, which then react with diaryl- or dialkyl disulfides (Scheme 58).

The formation of aryl trifluoromethyl sulfides from thioesters of trifluoroacetic acid occurs in rather better yields (30–40%) than from the corresponding esters of trifluoromethanesulfonic acid (20–30%). Alkyl thioesters of trifluoroacetic and trifluoromethanesulfonic acids form AlkSCF$_3$ in higher yields (up to 80%). As shown in Scheme 58, the CF$_3$• radical can attack at several sites. Phenyl selenide esters of trifluoromethanesulfonic acid react analogously [218].

The photochemical decomposition of trifluoromethanesulfonic and carboxylic thioesters affords CF$_3$• radicals which can be used to prepare trifluoromethyl sulfides [219].

Xenon difluoride has been used to initiate oxidative decarboxylation of perfluorocarboxylic acids for R$_F$•, generation and with aromatic and heterocyclic compounds the perfluoroalkyl groups can also become incorporated into the aromatic ring [220]. Nevertheless, Sipyagin et al., have employed this method for the perfluoroalkylation of thiols such as polychloropyridine thiols [221]. Two different methods were used: the action of preformed xenon carboxylates (method A) or treatment of a pyridinethiol solution in R$_F$COOH directly with xenon difluoride (method B). A range of isomeric perfluoroalkyl sulfides was obtained (Scheme 59).

Similar reactions have been carried out with tetrafluoropyridine 4-thiol [224] and its corresponding disulfide [225,226] (40–50% yield). The formation of S-perfluoroalkyl derivatives with performed xenon carboxylates from nitro aromatic disulfides was also successful (Scheme 60).
Perfluoroalkylsulfinic acids can also be used for oxidative decomposition. For example, careful treatment of sodium trifluoromethanesulfinate with tert-butyl hydroperoxide in the presence of an organic disulfide gives the corresponding trifluoromethyl sulfide [228,229]. Aliphatic disulfides react well to give alkSCF₃ but problems arise with aromatic disulfides due to attack of the CF₃• radical on the aromatic rings. For example, diphenyl disulfide is converted only in 13% yield. The S/C ratio reflecting the amount of trifluoromethylation on sulfur and on the aryl ring depends on the solvent. In CH₃CN it is 36:64, while in aqueous CH₃CN it is 60:40. Dichlorodiphenyl disulfide gives the best ratio in favor of the sulfide in aqueous acetonitrile [228].

One final method of CF₃• radical generation involves the interaction of Bi(CF₃)₃/Cu(OCOCH₃)₂ with thiophenolate (Scheme 61).

The above methods for the synthesis of aryl perfluoroalkyl sulfides all generate electrophilic R₄⁺ radicals which prefers to react at nucleophilic reaction centers such as S⁻ or C=S or S•. In the case of diaryl disulfides [228] the regioselectivity of attack is less controlled due to ring delocalization.

4.3. Anionic perfluoroalkylation

This method of perfluoroalkylation involves the reaction of aromatic or heterocyclic sulfur compounds with perfluoroalkyl anions, stabilized by suitable ligands, or with a reagent that generates such an anion.

Perfluoroalkyl anions are extremely unstable. For example, the CF₃⁻ anion decomposes at −100 °C with the elimination of F⁻ and formation of difluorocarbene, which reacts further or dimerizes [123]. Nevertheless, in the last two decades nucleophilic perfluoroalkylation of organic compounds has expanded. The problem of R₄⁻ stability in synthesis has been reviewed [24]. Trifluoromethylated reagents of heavy metals and their application in organic synthesis were considered by Barton [25], whilst perfluoroalkylated [31,32] and trifluoromethylated [27,28,30] organosilicon compounds have attracted considerable interest. However, despite the large body of literature involving the use of such reagents, the synthesis of aryl perfluoroalkyl sulfides is restricted to anionic attack on sulfenyl chlorides and thiocyanates.

Various methods for the synthesis of aryl perfluoroalkyl sulfides, depending on the mode of generation of the perfluoroalkyl anion, are described below.

4.3.1. “R₄⁻” from a perfluorinated olefin

Relatively stable tertiary perfluoroalkyl carbanions can be prepared by addition of fluoride ion to fluoroolefins [151,231-234] or by the deprotonation of monohydroperfluoroalkanes or their derivatives [235,236] as shown in Scheme 62. Most processes involve generating the hexafluoroisopropyl carbanions with a third stabilizing group such as CF₃ [151,231,232,236], C₃F₇ [233,234], as well as CN, COC₂F₅, COOMe [232,236]. Reactions of the resulting salts with aryl
Scheme 62: Reaction of fluorinated carbanions with aryl sulfonyl chlorides.

**4.3.2. “R F−” from perfluoroalkyl halogenides**

In a similar manner to alkyl halides, perfluorinated alkyl halides also form organometallic derivatives which can be used for the synthesis of perfluoroalkyl sulfides. The effectiveness of such reagents depends largely on the counterion which is illustrated below for reactions with organic thiocyanates (Scheme 64). Potassium perfluoroisopropyl (generated from CF2=CFCF3 and KF) reacts with phenyl- and p-nitrophenyl thiocyanates in sulfolane at 100 °C, whilst the Grignard reagent (n-C4F9MgI) reacts at subzero temperatures.

Cuprates react with benzyl thiocyanate but require more forcing conditions, i.e., 100 °C [237], whereas in situ generated zinc reagents R3ZnX react with thiocyanates at 20 °C in pyridine [238].

Scheme 63: Reaction of methyl perfluoromethacrylate with PhSCl in the presence of fluoride.

sulfonyl (or aryl selenyl) chlorides yield perfluoro- or polyfluoroalkyl sulfides (selenides).

The [C3F7(CF3)2C]− anion, obtained from isomeric dimers of perfluoropropylene in the presence of KF or CsF, reacts with sulfonyl chlorides and selenyl chlorides to afford the corresponding sulfides and selenides bearing a tertiary perfluorohexyl group [233].

In the reaction of R F− carbanions with sulfonyl chlorides high yields of sulfides are obtained when either electron-withdrawing or electron-donating substituents are present on the aryl ring. The yields of isomeric nitrophenyl perfluoro-tert-butyl sulfides decrease, the closer the nitro group is to the sulfur atom: p-NO2 – 86%, m-NO2 – 78% [231] and o-NO2 – 68% [232]. Both secondary and tertiary anions react [236] but nature of the counter ion is important. Thus, cesium or potassium perfluoro tert-butyl alkyls obtained by the addition of CsF or KF to perfluoroisobutene, give high yields of ArSC(CF3)3 [151,231,232], while the same anion, generated by deprotonation of nonfluoroisobutane (CF3)3CH with NEt3 gives PhSC(CF3)3 in low yield ~20% [236].

In the reaction of methyl perfluoromethacrylate with PhSCl in the presence of fluoride ion, prolonged stirring gave two sulfides as shown in Scheme 63, illustrating the competition between halides (F− and Cl−) for fluoroolefin addition [232].
Recently, it has been shown that tetrakis(dimethylamino)ethylene (TDAE) can undergo a two-electron transfer to perfluoroalkyl iodides to generate $\text{R}_2\text{F}^-$ anions [187] which react with organic disulfides to afford perfluoroalkyl sulfides in high yields [239,240]. The economy of this method, as distinct from previous methods [196,241-248], lies in the fact that the thiolate released by the first nucleophilic attack on the disulfide reacts directly with a second equivalent of perfluoroalkyl iodide, to form a second equivalent of the desired perfluoroalkyl sulfide (Scheme 65). This approach thus combines two principles of trifluoromethylation, i.e., nucleophilic attack of the $\text{R}_2\text{F}^-$ anion on the disulfide and reaction of a radical anion with a thiol as noted in section 4.1.

4.3.3. “$\text{R}_2\text{F}^-$” from perfluorocarboxylic acids

A simple method for the generation of metal derivatives of perfluoroalkyl carbanions by the decarboxylation of alkali salts of perfluorocarboxylic acids, has also been used. For example, heating potassium perfluoroalkyl carboxylates in the presence of diaryl disulfides in DMF or sulfolane leads to the formation of the corresponding aryl perfluoroalkyl sulfides as summarized in Table 14.

Disulfides of pyridine [242], pyrimidine and naphthalene [249] have also been used in such reactions. The use of this method for longer perfluorocarboxylic acids leads to product mixtures that result from chain isomerism and cyclisation [250,251] (Scheme 66).

### Table 14: Perfluoroalkylation of aryl disulfides by decarboxylation of perfluorocarboxylates.

| $\text{R}$ | $\text{R}_2\text{F}$ | Solvent | $T$ (°C) | Yield of $\text{ArSR}_2\text{F}$ % | Ref. |
|------------|----------------------|---------|---------|-------------------------------|------|
| H          | CF$_3$               | DMF     | 140     | 84                            | [245]|
| H          | CF$_3$               | sulfolane | 180–230 | 56                            | [242]|
| 4-Me       | CF$_3$               | sulfolane | 180–230 | 51                            | [242]|
| 4-Cl       | CF$_3$               | sulfolane | 180–230 | 56                            | [242]|
| 4-F        | CF$_3$               | sulfolane | 180–230 | 82                            | [242]|
| 2-Br       | CF$_3$               | sulfolane | 180–230 | 48                            | [242]|
| 4-OMe      | CF$_3$               | sulfolane | 180–230 | 50                            | [242]|
| H          | C$_2$F$_5$           | DMF     | 145     | 70                            | [245]|
| 4-Me       | C$_2$F$_5$           | DMF     | 145     | 50                            | [245]|
| 4-NO$_2$   | C$_2$F$_5$           | DMF     | 145     | 42                            | [245]|

Polyhalogenated carboxylic acids containing fluorine together with other halogens can also alkylate disulfides. However, the results strongly depend on the structure of halogenated alkyl group. The method is successful for potassium trichloroacetate but not for difluorochloroacetate. In the latter case the corresponding sulfide PhSCF$_2$Cl was found but only in trace amounts whilst PhSCCl$_3$ is obtained in 80% yield [245]. The mixed haloalkyl anions appear to be less stable.

The stability and reactivity of perfluoroalkyl anions largely depend on the solvents used. For example, CF$_3$Mgl [252-254] and CF$_3$Li [123,255-258] in diethyl ether are unstable even at low temperatures, but in coordinating solvents such as sulfolane, N-methylpyrrolidone, HMPA and especially, in DMF, the CF$_3^-$ anion does not decompose so readily and can be used as a nucleophilic reagent [259].

4.3.4. “CF$_3^-$” from trifluoromethane (fluoroform)

Trifluoromethane (fluoroform) has been used as a source of the trifluoromethyl anion. Trifluoromethane is a waste product of
Teflon manufacture and it is of interest as a raw material for organofluorine chemistry [260]. However, its application has been restricted by the low stability of the CF$_3^-$ anion [123,252-255].

The CF$_3^-$ anion has greater stability when the counter ion is a bulky ammonium ion, and in the presence of pyrrolidone it reacts with aldehydes and ketones [261]. This suggests that an intermediate gem-aminoalcoholate is involved. The method is improved with DMF, which is also thought to form a stable aminoalcoholate intermediate (Scheme 67) [243,262,263].

This mechanism is supported by the observation that equivalent reactions do not occur in THF or DMSO [263]. Furthermore, the intermediate CF$_3$ aminoalcoholate has been trapped in its protonated form and as hydrated trifluoroacetaldehyde by the action of acids, as well as trapped as a silyl ether [243]. The deprotonation of fluoroform has been applied successfully for the synthesis of aromatic trifluoromethyl sulfides and selenides, as summarized in Table 15.

Langlois et al. have used silylated amines in the presence of fluoride ion to promote fluoroform deprotonation [244]. For example, with (Me$_3$Si)$_3$N such reactions were possible in both DMF and THF. In the latter case stabilization of the CF$_3^-$ anion and its reaction with disulfide probably involves a transition state complex such as that depicted in Scheme 68.

In the case of trifluoromethylation of aliphatic disulfides, silazanes are the preferred reagents. However, in the case of diaryl disulfides, e.g., diphenyl disulfide, the significant formation of byproducts occurs and, PhSN(TMS)$_2$ (46%) and PhSCHF$_2$ (23%) are main reaction products. Other CF$_3$ aminomethanols have been synthesized by Langlois et al. [264] (Figure 7).

Table 15: Reaction of the CF$_3^-$ anion derived from fluoroform with S-derivatives of thiophenols.

| B           | R  | X   | Yield, % | Ref. |
|-------------|----|-----|----------|------|
| t-BuOK      | H  | SPh | 80       | [243]|
| t-BuOK      | H  | SO$_2$Ph | 90     | [243]|
| t-BuOK      | 4-NO$_2$Cl |     | 80       | [243]|
| LiN(TMS)$_2$(NH(TMS)$_2$ | H  | SPh | 4        | [244]|
| N(TMS)$_2$/Me$_4$NF | H  | SPh | 6        | [244]|
| t-BuOK      | H  | SPh | 82       | [244]|
| N(TMS)$_2$/Me$_4$NF | H  | SePh | 61$^a$  | [244]|
| t-BuOK      | H  | SePh| 77$^a$   | [244]|

$^a$PhSeCF$_3$.

Trifluoromethylation of disulfides by the first of them was efficient, for example, 87% in the case of PhSCF$_3$ but less efficient for diselenides (PhSeCF$_3$ 45%) [246]. The reaction failed with bis(4-chlorophenyl) disulfide and dioctyl disulfide where only by-products were generated.

Silylated hemiaminals are more suitable for CF$_3^-$ transfer (Table 16), although high reaction temperatures (60–80 °C) are required.

The use of F$^-$ anion as an alkaline agent (De-Shopge reagent, Bu$_4$N$^+$ Ph$_3$SiF$_2^-$) in place of a strong base (t-BuOK) allows trifluoromethylation of aliphatic disulfides.
4.3.5. \( \text{CF}_3^- \) anion from trifluoromethyl silanes

Perfluoroalkyltrialkyl silanes in the presence of fluoride ion generate reactive \( R_3 \) carbanions which have been used widely in synthesis [27,28,30-32,265]. For example, Ruppert’s reagent, \( \text{CF}_3\text{SiMe}_3 \) [266] and its tin analogue \( \text{CF}_3\text{SnMe}_3 \) have been used for the nucleophilic introduction of a \( \text{CF}_3 \) group to electrophiles [274]. Prakash et al. have adapted this chemistry for the synthesis of aryl trifluoromethyl sulfides, it should be noted that these are prepared from sulfones, which are in turn synthesized from the same sulfides.

Reactions proceed easily in THF or light hydrocarbon solvents and the reaction can also be extended to aliphatic and heterocyclic [271] sulfur-trifluoromethylations. The data (Table 17) indicate that the source of the \( \text{F}^- \) anion exerts an important influence on the reaction of sulfonyl chlorides with \( \text{CF}_3\text{SiMe}_3 \) [267]. For example, in the presence of TASF \( p \)-nitrophenyl trifluoromethyl sulfide is formed in almost 70% yield, while the use of \( \text{Bu}_4\text{NF} \) (even 2 equiv) under identical conditions gives only a 14% yield. In addition, in the reaction of diaryl disulfides with \( \text{CF}_3\text{SiMe}_3 \) it has been shown [241] that the best results are obtained when the \( \text{Bu}_4\text{NF} \) is added with a syringe-pump rather than by ordinary dropwise addition.

Such trifluoromethylation reactions with \( \text{CF}_3\text{SiMe}_3 \) can also be catalysed with cyanide ion. However, this also results in competing side reactions where the cyanide attacks the sulfide directly and is especially problematic in the case of aliphatic disulfides [271].

4.3.6. \( \text{CF}_3^- \) anion from \( \text{ArSOCF}_3 \) and \( \text{ArSO}_2\text{CF}_3 \)

Aryl trifluoromethyl sulfones react with \( \text{CH}_3\text{ONa} \) to generate sodium arylsulfonates and fluoroorom [272], and with Grignard reagents to generate aryl alkyl- or diaryl sulfones [273]. Also nucleophilic substitution of the pentafluoroethyl group can be induced in bis(pentafluoroethyl) sulfone by various nucleophiles [274]. Prakash et al. have adapted this chemistry for nucleophilic trifluoromethylation. Both phenyl trifluoromethyl sulfone or the corresponding sulfoxide on treatment with \( t\)-BuOK in DMF generate a \( \text{CF}_3 \)-adduct similar to that formed during fluoroorom deprotonation [243,263], which is a useful trifluoromethylating agent for aldehydes, ketones and disulfides [248]. An example is shown in Scheme 69.

On the other hand, under the same reaction conditions methyl trifluoromethyl sulfone does not function as a trifluoromethylating agent, whilst esters and amides of trifluoromethane sulfonic acid are good trifluoromethyl transfer agents [247] (Scheme 70).

However, trifluoromethylation strategies with aryl trifluoromethyl sulfoxides, sulfones, -sulfinites, and amides have to compete with cheaper reagents such as fluoroorom, trifluoroacetic acid derivatives and trifluoromethyl halogenides. For the synthesis of aryl trifluoromethyl sulfides, it should be noted that these are prepared from sulfones, which are in turn synthesized from the same sulfides.

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**Table 16: Reactions of silylated hemiaminals with disulfides [246].**

![Image of Table 16]

| X     | \( F^- \) | R   | Yield, % | Ref. |
|-------|-----------|-----|----------|------|
| Cl    | TASF      | H   | 59       | [270]|
| Cl    | TASF      | 4-Cl| 72       | [270]|
| Cl    | TASF      | 4-NO_2| 69    | [270]|
| Cl    | \( \text{Bu}_4\text{NF} \) | 4-NO_2| 14     | [241]|
| SPh   | \( \text{Bu}_4\text{NF} \) | H   | 32 (43\(^b\)) | [241]|
| CN    | \( \text{Bu}_4\text{NF} \) | H   | 70 (58\(^b\)) | [271]|
| CN    | \( \text{Bu}_4\text{NF} \) | 4-NO_2| 58     | [271]|
| CN    | \( \text{Bu}_4\text{NF} \) | 2.4-(OMe)_2| 30    | [271]|

\(^a\)TASF = (Me_2N)_S\(^b\) (Me_3SiF)_2.

\(^b\)ArSeCF_3.

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**Table 17: Trifluoromethylation of sulfur and selenium compounds with Ruppert’s reagent.**

![Image of Table 17]

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**Scheme 69: Trifluoromethylation of diphenyl disulfide with PhSO_2CF_3/ \( t\)-BuOK.**

![Image of Scheme 69]

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**On the other hand, under the same reaction conditions methyl trifluoromethyl sulfone does not function as a trifluoromethylating agent, whilst esters and amides of trifluoromethane sulfonic acid are good trifluoromethyl transfer agents [247] (Scheme 70).**

However, trifluoromethylation strategies with aryl trifluoromethyl sulfoxides, sulfones, -sulfinites, and amides have to compete with cheaper reagents such as fluoroorom, trifluoroacetic acid derivatives and trifluoromethyl halogenides. For the synthesis of aryl trifluoromethyl sulfides, it should be noted that these are prepared from sulfones, which are in turn synthesized from the same sulfides.
4.4. Cationic perfluoroalkylation

Aryl perfluoroalkyl iodonium reagents as perfluoroalkylating agents were first developed by Yagupolski et al. [275]. Unlike perfluoroalkyl iodides, tolyl perfluoroalkyl iodonium chlorides react easily with sodium thiophenolates and selenophenolates at low temperature to form the corresponding aryl perfluoroalkyl sulfides and selenides as summarized in Table 18.

These iodonium salts even react with sodium p-nitrothiophenolate and while C$_3$F$_7$I does not react without some initiation [189] the C$_3$F$_7$ containing salts (Table 18) react readily. The yields of p-O$_2$NC$_6$H$_4$SR$_F$ (R$_F$ = C$_3$F$_7$ and C$_6$F$_{13}$) are increased to a quantitative level by the use of iodonium tetrafluoroborate salts [276] instead of chlorides.

Similarly, perfluoroalkyl phenyl iodonium trifluoromethanesulfonates (FITS reagents) react with thiolates [277]: Perfluoroalkylation is selective for sulfur even in the presence of other functional groups (e.g. OH, NHMe, COOH, COOAlk). The preparation and application of R$_F$ iodonium salts has been reviewed [33]. However, CF$_3$ iodonium salts were not discussed, presumably due to their low stability.

A “hyper-valent” iodine (III) compound containing a trifluoromethyl group, first synthesized in 2006 [278], appears to be quite stable. This moisture-sensitive reagent reacts with aromatic, heterocyclic and aliphatic thiols at low temperature (~78 °C) with the formation of the corresponding SCF$_3$ derivatives in high yields (Scheme 71).

**Table 18:** Interaction of tolyl perfluoroalkyl iodonium chlorides with sodium thiophenolates and selenophenolate [275].

| R   | R$_F$  | Yields, % (GLC) of ArSR$_F$ | ArSeOR$_F$ | a |
|-----|--------|-----------------------------|-------------|---|
| H   | C$_3$F$_7$ | 67 (81)                 | 87          |   |
| CH$_3$ | C$_3$F$_7$ | 71 (96)                  | -           |   |
| NO$_2$ | C$_3$F$_7$ | 34 (56)                  | -           |   |
| H   | C$_6$F$_{13}$ | 41 (56)                 | 45          |   |

aAfter chlorination and subsequent hydrolysis of corresponding selenides.

**Scheme 70:** Amides of trifluoromethane sulfonic acid are sources of CF$_3^-$ anion.

**Scheme 71:** Trifluoromethylation of various thiols using “hyper-valent” iodine (III) reagent [279].
However, this attractive methodology has some drawbacks in that its synthesis involves four steps and trifluoromethylation products must be purified by chromatography to remove a side-product – 2-iodophenyl dimethyl carbinoil.

Unlike iodonium salts, onium salts of the group VI elements appear to be more stable with CF$_3$ group. Diaryl R$_F$-sulfonium salts, where R$_F$ = CF$_3$, are readily synthesized from aryl trifluoromethyl sulfoxides [280]. Reaction of these reagents with sodium p-nitrothiophenolate affords the trifluoromethyl sulfide in good yield (Scheme 72).

Subsequently, diaryl thiophenium, -selenophenium and -tellurophenium reagents have been developed with perfluoroalkyl groups attached to S, Se and Te [33,281,282] which can transfer perfluoroalkyl fragments to nucleophilic centers. In particular, the dibenzo (CF$_3$)S-, (CF$_3$)Se- and (CF$_3$)Te-phenium systems have been investigated. For example, S(CF$_3$)dibenzo-thiophenium triflate (A = S) reacts with sodium thiolate in DMF to give the S-trifluoromethyl derivative in high yield. The related selenophenium salt (A = Se) appears to be more effective in trifluoromethyl transfer (Scheme 73).

The same general reactivity is also observed in reactions of these reagents with aliphatic thiols. Dibenzoselenophenium triflate (A = Se, R$_1$ and R$_2$ = H) reacts much better with sodium dodecyl thiolate (yield of C$_{12}$H$_{25}$SCF$_3$ is 87%) than the sulfur analogue (yield 47%) [283,284].

On the whole R$_F$ onium compounds are powerful perfluoroalkylating agents [33,281], however they are rather exotic reagents which require to be synthesized by multi-stage methods as illustrated in Scheme 74.

**Conclusion**

A summary of the known methods for the synthesis of aromatic and heterocyclic perfluoroalkyl sulfides are presented. These
involve perfluoroalkylation of thiols by single electron transfer, nucleophilic and electrophilic methods. The variety of methods reflects the level of interest chemists have given to generating this class of fluorine containing organic compounds. As a class of compounds, perfluoroalkyl sulfides find increasing utility in agrochemical and pharmaceutical applications.

A concise review concerning the preparation of selectively fluorinated ethers, thiocarbamates, amines and phosphines was published [285] during preparation of this manuscript.

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