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Chemistry of Pentafluorothio (SF$_5$) Alkyl Derivatives

Rolf Walter Winter

Portland State University

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CHEMISTRY OF PENTAFLUOROTHIO (SF₅) ALKYL DERIVATIVES

by

ROLF WALTER WINTER

A dissertation submitted in partial fulfillment of the requirements for the degree of

DOCTOR OF PHILOSOPHY

in

ENVIRONMENTAL SCIENCES AND RESOURCES/CHEMISTRY

Portland State University
1990
TO THE OFFICE OF GRADUATE STUDIES:

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AN ABSTRACT OF THE DISSERTATION OF ROLF WALTER WINTER FOR THE DOCTOR OF PHILOSOPHY in Environmental Sciences and Resources/Chemistry presented October 4, 1990.

Title: Chemistry of Pentafluorothio (SF₅) Alkyl Derivatives.

APPROVED BY MEMBERS OF THE DISSERTATION COMMITTEE:

Gary L. Gard, Chairman
Carl C. Wamser
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John Dash

This thesis is divided into four parts:

Part I describes the preparation of a number of useful intermediates from the sultone F₅SCHCF₂OSO₂, (1). These intermediates include the following: the ketene
F₅SC(SO₂)=C=O (21), the unique sultone F₄S=CCF₂(OSO₂) (23), and various ionic intermediates of the form Y⁺ [F₅SC(SO₂F)COX]⁻. These intermediates play an important role in providing new pathways for preparing fluorosulfonic acid precursors. Fluorosulfonic acids are excellent candidates to improve the efficiency of environmentally safe fuel cells. It was possible to obtain new derivatives of 1 by various reactions of 21, 23 and the ionic intermediates, but it was not possible to introduce an alkyl substituent with either of the intermediates.

Part II is concerned with the synthesis of SF₅ containing epoxides as possible monomers. When polymerized, materials of high dielectric strength and high chemical resistance should be obtained. The reaction chemistry of one of the epoxides was investigated and it was found that all reactions studied were accompanied by the loss of the SF₅ group. Thus, it was not possible to obtain any polymeric materials.

Part III deals with the synthetic accessibility of SF₅ malonic esters and thence of SF₅ acrylic esters. These compounds should serve as building blocks in the synthesis of SF₅ containing pyrimidines. The CF₃ analogue, 5-trifluoromethyl uracil, is used as an antiviral drug. In order to synthesize either SF₅ malonic ester or 2-SF₅ acrylic ester an SF₅ group has to be introduced at a secondary carbon. A secondary SF₅ compound could be
synthesized (F₅SCH(COOC₂H₅)(CHBrOAc)) but could not be converted to the respective malonic ester F₅SCH(COOC₂H₅)₂.

Part IV describes the reactions and preparations of part I-III.
To My Family
ACKNOWLEDGEMENT

I would like to thank Professor Gard for his support and kindness in the course of this work. I am grateful to the bright stars on their celestial path that guided me in the years past towards Arcadia.
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PART I

DERIVITIZATIONS OF $F_5SCHCF_2OSO_2$
CHAPTER I
INTRODUCTION

The investigation of compounds containing the SF$_5$ substituent is relatively new. It originated in 1934 with the discovery of disulfur decafluoride, S$_2$F$_{10}$, as a by-product of the fluorination of sulfur; the main product in this reaction is sulfur hexafluoride, SF$_6$. Sulfur hexafluoride can be considered as an SF$_5$ compound ("pentafluorosulfur fluoride"), but for chemical purposes this kinship cannot be employed due to the great inertness of this compound. On the other hand, S$_2$F$_{10}$ is useful as a starting material for making SF$_5$ compounds because the relatively weak S-S bond [55±5 kcal/mol, as derived from an empirical relationship between the bond energy and force constant] breaks easily upon heating and can be used in certain cases to obtain SF$_5$ derivatives directly, or more commonly, through the intermediate formation of SF$_5$Br, pentafluorosulfur bromide. SF$_5$Br was used as the chief starting material in this work and in the course of this investigation it was necessary to develop a new method of synthesis, due to the difficulty of obtaining sufficient quantities of S$_2$F$_{10}$. There are some remarkable similarities between the derivatives of SF$_6$ and CF$_4$, and there is the
possibility of the existence of a homologous series, analogous to \( \text{C}_n\text{F}_{2n+2} \), of the general composition \( \text{S}_n\text{F}_{4n+2} \), of which \( \text{SF}_6 \) and \( \text{S}_2\text{F}_{10} \) are the first and so-far only members. \( \text{SF}_5^- \) and \( \text{CF}_3^- \)-alkyl compounds usually show little tendency to react at their \( \text{CF}_3 \) and \( \text{SF}_5 \) sites, respectively. They are thermally very stable, form olefins with electrophilic \( \pi \)-bonds, and do not allow for nucleophilic displacement of halogens in the vicinity of the perfluoro groups. Such substitution reactions do occur for \( \text{CF}_3 \) (perfluoroalkyl) derivatives, but with a different mechanism (S_{RN}1); for similar \( \text{SF}_5 \) compounds a different course of reaction is usually observed, that is, loss of \( \text{SF}_5 \), although in some cases substitution reactions were successful.\(^3\) Here, and as will be seen later, some examples of the divergence in the behavior of the two types of compounds (\( \text{SF}_5 \) vs. \( \text{CF}_3 \)) are already seen.
CHAPTER II

SIGNIFICANCE AND ENVIRONMENTAL RELATIONSHIP

The development of electrochemical energy converters, or fuel cells\(^4\), as convenient and clean sources of electrical energy requires also the knowledge of suitable mobile charge carriers. Schematically, a fuel cell consists of two electrodes, separated by a diaphragm and connected by an external circuit. The charge transport within the solution is provided by the electrolyte. Oxygen is reduced at the cathode and hydrogen or another hydrocarbon fuel is oxidized at the anode; both electrodes act also as electrochemical catalysts by lowering the activation barrier for the two half reactions. It is especially advantageous to use platinum as part of the electrode material. The process of combustion is thus spatially separated, which allows one to make use of the electron flux stemming from the oxidation and reduction processes taking place. High efficiency and no product formation, other than water, carbon dioxide and nitrogen, depending on the fuel, is expected from such a device. There are only a certain number of fluorosulfonic acids available as possible replacements for phosphoric acid, which is now used as the fuel cell electrolyte of choice. Trifluoromethane sulfonic acid is comparable to
\( \text{H}_3\text{PO}_4 \) as a fuel cell electrolyte, but is too volatile at the required operating temperature. Nafion, a solid polymeric fluorosulfonic acid electrolyte, has been found to be useful. Therefore, fluorosulfonic acids offer promise as fuel cell electrolytes and the search for new and improved materials continues. Desirable properties of or changes brought about by a new fuel cell electrolyte are as follows:

a.) Chemical, electrochemical and thermal stability

b.) Good ionic conductivity

c.) High boiling point

d.) High capacity of oxygen solubility

e.) Improvement of the kinetics of the cell

Fuel cells increase in efficiency as the temperature rises, thus thermal stability and high boiling point, to minimize losses by evaporation, are a chief concern in choosing an electrolyte.

The above listed criteria (a-e) are largely met by fluorinated sulfonic acids; whereas chemical and thermal stability can be predicted as a property, this is not obvious for the electrochemical behavior. At that point, structural modifications of such acids can be quite arbitrary, but certain alterations of existing sulfonic acids are desirable. A high degree of fluorination conveys the property of chemical and thermal stability and increases the acid strength, due to the anion stabilizing inductive and hyperconjugative effect of the fluoroalkyl group,
particularly perfluoroalkyl residues. The introduction of alkyl-residues, particularly fluoroalkyl radicals, should be advantageous, accounting for greater branching of the acid, with the ensuing properties, such as the increase of $O_2$-solubility.

The present work was conducted with the aim towards modifying sulfonic acids that contain already the pentafluorosulfur, $SF_5^-$, group. Most fluorosulfonic acids are obtained from fluoro olefins via their intermediate $\beta$-sultones. Sultones are sulfonic acid analogues of lactones, i.e., they are internal esters of hydroxy sulfonic acids and in the present case of $\beta$-hydroxy sulfonic acids. The substitution pattern of the acid is determined by the structure of the olefin. Modification of the substituents of an already existing sulfonic acid or its precursor, the sulfonyl fluoride, is thus a possible method for the synthesis of new fluorosulfonic acids, bypassing the need for certain olefins. If the respective olefins cannot be synthesized, or cannot be sulfur trioxidized, then the need for such a general procedure, which involves either synthesis of sultones by a different route, and their subsequent rearrangement, or the chemical manipulation of the rearranged forms of the sultones and their secondary products, usually acyl fluorides or esters, becomes obvious. A further possibility for modifying sulfonic acids is by integrating them in a polymeric chain. This strategy has led
to the synthesis of Nafion (a tradename by E.I. DuPont de Nemours). Nafion is a polymeric perfluoro sulfonic acid which is obtained by sequential ionic co-polymerization of a fluorosulfonyl perfluoroalkoxide (obtained from the sultone \( \text{CF}_2\text{CF}_2\text{OSO}_2 \)) with hexafluoropropylene oxide, treatment with base, leading to salt formation, decarboxylation and elimination of NaF and the formation of a terminal fluoroolefin (a polyether). This olefin is then cross-linked with tetrafluoroethylene. The last step consists of hydrolyzing the \( \text{SO}_2\text{F} \) group, and the result is the formation of a strong-acid membrane. In order to obtain an "SF\(_5\)-Nafion" the synthesis of SF\(_5\)-epoxides or SF\(_5\)-sultones is necessary. In this context is also worthwhile to mention that these epoxides should provide dielectric materials because the superb electron capture capability of SF\(_6\) is carried over to its derivatives. It is generally believed that SF\(_5\)C=\( \text{N}^5 \) has a greater dielectric strength than the parent compound and CF\(_3\)SF\(_5\)\(^6\) and SF\(_5\)CF=CF\(_2\)\(^7\) actually do have a greater dielectric strength than SF\(_6\). Retaining the essential parts of these molecules, it is expected that a polymer of an epoxide should also retain the dielectric property. Epoxides like SF\(_5\)CF\(_2\)\(^3\) or SF\(_5\)CF\(_2\)CF\(_2\)\(^3\), when polymerized, would fulfill the structural requirement. Like CF\(_3\) analogues, which have also great dielectric strength, one would expect them to be high-boiling, chemically and thermally highly inert, non-poisonous compounds.\(^8\) This would
make them ideal substitutes of other dielectrics, which are usually noxious compounds. The synthesis and some reactions of two simple SF₅-epoxides are dealt with in PART II of this manuscript.
CHAPTER III

GENERAL DISCUSSION, DERIVITIZATION OF F5SCCHF2OSO2

One of the aims of this research was to investigate ways that would allow for the derivatization of the sultone SF₅CHCF₂OSO₂⁻. It will be most useful in this context to review related perfluoroalkyl systems. Fluorosultones are precursors of sulfonic acids via a base catalyzed rearrangement:

\[ \text{R}_{f}\text{CX-CF}_2 \rightarrow \rightarrow \rightarrow \text{R}_{f}\text{CXCFO} \rightarrow \rightarrow \rightarrow \text{R}_{f}\text{CXCFO} \]

\[ \text{O}_2\text{S}--\text{O} \rightarrow \text{SO}_2\text{F} \rightarrow \text{SO}_3\text{H} \]

X = H, halogen; Rₙ = fluoroalkyl

The carbonyl fluoride does not survive the basic conditions necessary for the sequence SO₂F → SO₃H, but can be either converted to esters, and thence to ethers (by treatment with SF₄, which causes the conversion C=O → CF₂), or hydrolyzed to a C-H derivative, as described subsequently.

Fluorinated sultones do exhibit, as investigated particularly thoroughly in Russia, a very diverse derivative chemistry. It was expected to encounter many similar reactions with the present sultone ¹, as were found with other fluoro sultones. They are obtained by the electrophilic addition of sulfur trioxide to a fluoro olefin. Hydrolysis leads then, in sulfuric acid, to the formation of a carboxylic acid – the sulfur from the SO₃ is lost – and in basic media to the formation of a carboxylate.
Alcoholysis results in the formation of SO₂F-containing esters. It is of interest to alter these secondary products structurally, as not all olefins that could add SO₃ are readily available or do not react in the wished-for manner. There is the possibility of the formation of reactive intermediates, like ketenes or carbanions. Of the reactions that were investigated it was generally found that it is especially simple to obtain stable anionic species derived from the sultone F₅SCHCF₂OSO₂ (1).

All reactions can be carried out with the respective sulfonyl fluorides, and they can be rather simply converted to sulfonic acids. A close kin to F₅SCHCF₂OSO₂ is F₃CCHCF₂OSO₂(2), the simplest member of the series RfCHCF₂OSO₂. Its reaction chemistry has been investigated by Knunyantz (references are given subsequently).

The sultone CF₃CHCF₂OSO₂ forms a ketene, F₃CC(SO₂F)=C=O (3), when treated with the base F₃B·N(C₂H₅)₃ in ≈69% yield; when its isomer CF₃CH(SO₂F)COF (4), obtained by base catalyzed rearrangement of 3, was treated with the same base, the ketene was obtained in nearly 90% yield; it could also be obtained by heating the esters CF₃CH(SO₂F)COOR (R=isopropyl, t-butyl, but not ethyl or methyl) with P₂O₅ (≈70%). Originally it was synthesized by sulfurtrioxidation of 3-methoxy-3,3-difluoro-2-trifluoromethylpropionic acid. This ketene reacts with
SO$_3$ at 150-160°C, in contrast to (CF$_3$)$_2$C=O (6), which reacts already at -30°C.$^{13}$ From this the much greater electrophilicity of CF$_3$C(SO$_2$F)=C=O, as compared to (CF$_3$)$_2$C=O, is inferred. Sulfur trioxidation is usually assumed to be an electrophilic reaction.$^{14}$

The rather harsh conditions that were necessary for the sulfur trioxidation of CF$_3$FC(SO$_2$F)=C=O demonstrate the strong electron withdrawing capability of the SO$_2$F-group. The reactivity of this ketene has not been established on a broader basis. The result of the sulfur trioxidation reaction implies that there is a substantial difference between the reactivity of CF$_3$C(SO$_2$F)=C=O and (CF$_3$)$_2$C=C=O, but a detailed comparison is not possible, due to the few experimental results from trifluoromethyl fluorosulfonyle ketene. The reactivity of bistrifluoromethyl ketene is well-investigated$^{15}$ and it is found that it is susceptible to the attack of electron rich olefins and acetylenes, and both double-bonds can take part in bond formation. For example, the reaction with vinylbenzoate leads to a mixture of

$$\text{(CF$_3$)$_2$C=C=O \quad 34\%} \quad \text{and} \quad \text{(CF$_3$)$_2$C=C=O \quad 42\%}$$

The attack of SO$_3$ on CF$_3$C(SO$_2$F)=C=O takes place exclusively at the C=O bond, while reaction of (CF$_3$)$_2$C=C=O with CH$_2$=O,$^{3}$
at 150°C (catalyzed by ZnCl₂) leads only to the cyclobutanone derivative. Nucleophilic attack of the oxygen in tropone on the C=O group in Cl₂C=C=O is probably responsible for the formation of a 1,3-cycloaddition product, and bond formation is between Cα, Cβ and tropone. This constitutes still the normal direction of addition, which leads here to a γ-lactone.

An important reaction of (CF₃)₂C=O is the anionotropic rearrangement to perfluoromethacryloyl fluoride (8):18

\[
(CF₃)₂C=O \xrightarrow{F^-} (CF₃)₂C^-=O \xrightarrow{F^-} CF₂=O + CF₃\text{ 9}
\]

Perfluoromethacryloyl fluoride is a useful intermediate; for example, it allows one to obtain rather simply derivatives of trifluoromethyl malonic acid:

\[
CF₂=O + CH₃COOH \rightarrow [CH₃COOCF₂CH(CF₃)COF \rightarrow CF₃CH(COF)₂ (78\%) + CH₃COF
\]

The formation of β-alkylperfluoromethacryloyl fluoride is even more favored for β,β-difluoro-α-trifluoromethyl carboxylic acids; the intermediate ketene that is formed upon heating the carboxylic acid with P₂O₅ must have rearranged to the β-alkylperfluoromethacryloyl fluoride.21

The dimer of (CF₃)₂C=O, (CF₃)₂C=C(CF₃)₂

\[
O-C=O (9),
\]

a β-lactone, which, after Krespan and England, is formed from perfluoromethacryloyl fluoride and bistrifluoromethyl ketene in an F⁻-catalyzed reaction, can also be obtained
from (CF₃)₂C=O through the catalysis by weak bases like triethyl phosphite, tetraalkylureas, diethylnitrosamine, or dimethylacetamide.²³ The formation of a β-lactone is unusual in that disubstituted ketenes form more commonly 1,3-cyclobutanones as dimers. From the dimer of (CF₃)₂C=O one can then synthesize derivatives that contain several CF₃-groups, e.g. the reaction with water results in 1,1,3,3-tetra trifluoromethylacetone²⁴, or pyrolysis or treatment with P(OC₂H₅)₃ leads to (CF₃)₂C=C=C(F₃)₂ (93 %), which can also be obtained directly from (CF₃)₂C=C=O and P(OC₂H₅)₃. Thus, there is a broad spectrum of reactions that give access to complicated compounds which would otherwise not be easily synthesized.

Addition of an alkoxide to the C=O carbon of ketenes results in ester anions (enolates). Such anions, which have to be regarded within the larger group of allene and heteroallene adducts²⁵, were not described for bistrifluoromethyl ketene until recently, when the salt [(CH₃)₂N)₃S]⁺ [(CF₃)₂CCOF]⁻ was obtained. The stability of these adducts should depend upon the electrophilic nature of ketene; possibly, other nucleophiles could also add. An example is the addition of dichloroketene to tropone (page 12), which is a regioselective reaction, or the addition of amines HNR₂, which leads to amides. For bistrifluoromethyl ketene several reactions are described in the literature, which, although not resulting in a ketene salt, must pass
through an ionic intermediate, formed by nucleophilic attack at the C=O group. NaNO₂ addition results, after hydrolysis, in hexafluoroacetone oxime (CO₂ is lost in the reaction) and dimethyl formamide leads to the formation of (CF₃)₂C=CHN(CH₃)₂. Alkoxide adducts are well-investigated for F₃CC(SO₂F)=C=O:

\[
\begin{align*}
F₃C\text{C}=C=O + RO^- & \rightarrow \left[ \begin{array}{c}
\text{F₃C} \\
\text{FO₂S}
\end{array} \right] ^- \text{C=O + RO}^-
\end{align*}
\]

In reality, these anions are obtained by proton abstraction from esters of the composition F₃CCH(SO₂F)COOR with suitable bases and they can be chlorinated and brominated:

\[
\begin{align*}
\text{SO₂F} & / \text{CF₃-C-H + N(C₂H₅)₃} \rightarrow \text{CF₃-C}^- \\
\text{COOR} & \text{HN(C₂H₅)₃}^+ \rightarrow \text{CF₃-C-Br(Cl)} \quad \text{(IIa,b)}
\end{align*}
\]

R= CH₃, C₂H₅

The bromo adduct IIa (with Br₂) was obtained in 60\% yield, and the chloro adduct IIb (with SO₂Cl₂) in 80\% (R=CH₃) and 82\% (R=C₂H₅) yield. Chlorination with N,N-dichlorobenzenesulfonamide led to a dichloro compound:

\[
\begin{align*}
\text{SO₂F} & / \text{CF₃-C}^- + \phi \text{SO₂NCl₂} \rightarrow \text{CF₃-C-SO₂F} (40\%).
\end{align*}
\]

The fate of the ester group was not investigated.

Since the esters CF₃CH(SO₂F)COOR (13) descend directly from the sultone CF₃CHCF₂OSO₂ (2), there is, at least formally, a
relationship between the sultone, the ketene and the enolates. This relationship was employed for the sultone \( \text{SF}_5\text{CHCF}_2\text{OSO}_2 \) and found upon study to go beyond formal considerations. It is also possible to synthesize the ketene \( (\text{FO}_2\text{S})_2\text{C}=:\text{C}=\text{O} \) (from \( \text{FO}_2\text{SCH}=\text{CF}=\text{F} \))\(^{13}\), which would probably be the most electrophilic one of this series. It reacts with methanol with loss of one \( \text{SO}_2\text{F} \)-group but has not been appreciably investigated.

A comparison with similar fluorosulfonyl carbanions delineates to some degree the reactivity of these species. Other fluorosulfonyl carbanions can be obtained as stable salts or are implicated as intermediates in chemical transformations. The bistrifluoromethyl fluorosulfonyl carbanion \(^{(14)}\) is obtained from \( (\text{CF}_3)_2\text{CHSO}_2\text{F} \) or \( [(\text{CF}_3)_2\text{C}=\text{SO}_2]_3 \) with either pyridine or triethylamine\(^{28}\) or from 2,2,4,4-tetratrifluoromethyl-1,3-dithietane-1,1 dioxide with \([((\text{CH}_3)_2\text{N})_3\text{S}]^+ [\text{Si}(\text{CH}_3)_3\text{F}_2]^−\).\(^{29}\) When boiled with pyridine for a long time, the compound \( (\text{CF}_3)_2\text{CHSO}_2\text{F} \) gave a sulfene complex\(^{26}\):

\[
(\text{CF}_3)_2\text{CHSO}_2\text{F} + \text{C}_5\text{H}_5\text{N} \rightarrow (\text{CF}_3)_2\text{C}=\text{SO}_2\cdot\text{C}_5\text{H}_5\text{N} (86%)
\]

The free sulfene was obtained by Smart and Middleton from their tri-diaminomethylsulphonium salt by treatment with \( \text{BF}_3, \text{BF}_3\cdot(\text{C}_2\text{H}_5)_2\text{O} \) or \( \text{SiF}_4 \), which form \( \text{BF}_4^- \) or \( \text{SiF}_5^- \). The sulfene can be intercepted by reaction with suitable captors. Without a reactive partner, the olefin \( \text{CF}_2=\text{C}(\text{CF}_3)\text{SO}_2\text{F} \) was obtained, apparently formed by rearrangement of the
sulfene. Substitution reactions of anion 14 were complicated in some cases, e.g. in the reaction with perfluoroacetic anhydride, a CF₃⁻ fluorine was replaced, leading to CF₃COOCF₂C⁻(CF₃)SO₂F HN(C₂H₅)₃⁺ and CF₃COF. The mechanism of this transformation was formulated with a consecution of substitutions, additions and eliminations, with the assumption that CF₂=C(CF₃)SO₂F (16), as the final intermediate, add CF₃COO⁻. Based on these considerations it was possible to obtain the cesium salt, Cs⁺ [(CF₃)₂CSO₂F]⁻ (17) by reacting CsF with CF₂=C(CF₃)SO₂F (16), which was obtained from (CF₃)₂CHSO₂F (15) and F₃B·Pyridine. The cesium salt underwent normal substitution reactions with bromine or sulfuryl chloride, but also with BrCN:

(CF₃)₂C⁻SO₂F + BrCN ----> (CF₃)₂C(SO₂F)CN (34%)

The fact that a carbon substituent could be introduced at the anionic center in this case is significant in light of the failure which was encountered in this study with Kat⁺ [SF₅C(SO₂F)COR]⁻ (R = F, Kat⁺ = Cs⁺ and Ag⁺; R = OCH₃, Kat⁺ = HN(C₂H₅)₃⁺; R = OCH(CH₃)₂, Kat⁺ = HN(C₂H₅)⁺, Na⁺).

The sultone F₃CC(CF₂O)SO₂ (2), when treated with the base F₃B·N(C₂H₅)₃ (18, "Kazmina's base"), a strong hydrogen fluoride acceptor, leads cleanly to the ketene F₃CC(SO₂F)=C=O (Ref.11), although there are two possible directions of HF-abstraction:

\[
\begin{align*}
\text{H} & \quad --X--> \quad \text{F}_2\text{C}=-\text{C}-\text{CF}_2\text{O}\text{SO}_2 \quad (19) \\
\text{F}_3\text{C}-\text{C}-\text{CF}_2 + \text{F}_3\text{B} \cdot \text{N(C}_2\text{H}_5)_3 & \quad --X--> \quad \text{F}_3\text{C}-\text{C}=-\text{CFOSO}_2 \quad (20) \\
\end{align*}
\]

-----> F₃CC(SO₂F)=C=O (3, ≈90%)
It is contended that the base causes first rearrangement to the isomeric acyl fluoride $F_3CCH(SO_2F)COF$ (4), which then upon loss of hydrogen fluoride yields trifluoromethyl fluorosulfonyl ketene. The fact that $F_2C=CCF_2OSO_2$ was not formed was to be expected from the reaction of tetrafluoroallene, $F_2C=C=CF_2$ with $SO_3$, which led to $F_2C=C=CF_2OSO_2OSO_2$; a difluoromethylene group next to a sultone ring has never been described. Compound 19 is also an isomer of 3; if formed, it might undergo in the reaction environment rearrangement to 3. $F_3C-CFOSO_2$ was not expected to be found because the addition addition of $SO_3$ to $(F_3C)_3CC=CF$ led also to $F_3CC(SO_2F)=C=O$ (3) and passed probably through $F_3C-C=CFOSO_2$ (20). In this latter reaction the oxathiaacyclobutene derivative was formulated as an intermediate, but these compounds might be incapable of existence under ordinary conditions. Indeed, they have never been observed.

In this study, the ketene $F_5S$

\[
\begin{array}{c}
\backslash \\
C=C=O \\
\end{array} \]

\[
\begin{array}{c}
/ \\
F_0_2S \\
\end{array}
\]

was expected to be the most useful intermediate for further, secondary, transformations of 1. As with other ketenes with two bulky substituents it was expected to be a stable compound, i.e., it should not be prone to dimerizing without catalysis. It is the internal anhydride of the acid $F_5SCH(SO_2F)COOH$, which is not isolable. A similar mechanism
as with malonic acid can be formulated for the decarboxylation:

\[
\text{F}_5\text{SC} \rightarrow \text{F}_5\text{SCH}_2\text{SO}_2\text{F}
\]

Some conceivable reactions of 21 are delineated below.

The ketene should be able to react with perfluoro alcohols and this would comprise an easy method for the synthesis of perfluoro esters. These esters can then be converted to perfluoro ethers by treatment with SF₄ + HF₃². The ether linkage in perfluoro ethers is noted for its chemical and thermal stability.⁸ The carbon-SF₅ bond is also difficult to attack but is weaker than the C-O-C linkage in perfluoroethers. An example is the pyrolysis of 1,4 perfluorothioxane, which forms perfluoro tetrahydrofuran + SF₄ at 475°C over NaF.³³ Cycloadditions with carbonyl compounds, dimerization or nucleophilic reactions of the secondary enolates should lead to compounds substituted in α-position of both the SF₅ and SO₂F group. Moreover, the enolates could be halogenated, and the resulting α-halogen
compounds should undergo reactions with olefins to form α-C substitution products; there is of course the risk that they lose the ester group, as was found with 10 in the reaction with \( \phi \text{NCl}_2 \). A similar ketene, \( \text{F}_5\text{SCH=O} \), was described by Seppelt et al.\(^{34}\) It formed a dimer upon heating (a \( \beta \)-lactone, \( \text{F}_5\text{SCH=C-O} \)), polymerized in the reaction with KF or CsF, underwent the usual addition reactions of ketenes with ethanol, hydrogen chloride, water and bromine, and was also subject to an anionotropic rearrangement when passed over glass (weakly basic) at \( \approx 290^\circ \text{C} \) at low pressure. This rearrangement is the analogue to the relationship between 5 and 8.

\[
\begin{align*}
\text{F}_5\text{S-C=C=O} \ (5a) \rightarrow \text{F}_4\text{S=C=O} \ (8a)
\end{align*}
\]

The \( \text{SF}_4 \) derivatives that are obtained are very reactive compounds (sulfur ylides) and are prone to undergoing polar additions, such that cis adducts are obtained.\(^{35}\) cis refers to the octahedral sulfur environment. There is thus a possibility to alter the \( \text{SF}_5 \) group by passing through intermediate sulfur ylides. Compound 8a can be hydrolyzed to give the compound (\( \text{F}_2\text{S}=\text{O} \))=CHCFO (8b), or it reacts with alcoholic base to form cis-(\( \text{F}_4\text{SOC}_2\text{H}_5 \))CH\(_2\text{COOC}_2\text{H}_5 \).\(^{36}\)

Darragh has shown that \( \text{R-SF}_4\text{Cl} \) (trans) reacts completely analogously to \( \text{SF}_5\text{Cl} \) with olefins and acetylenes to form, for example with ethylene, trans-\( \text{ClCH}_2\text{CH}_2\text{SF}_4\)-R.\(^{37}\) Complementary results were also obtained by Shreeve.\(^{38}\) There is thus the possibility that the cis-\( \text{RSF}_4\text{Cl} \) adducts react
similarly to the trans-adducts. With the latter, configurational stability was observed, trans ClSF₄₋ gave a product trans-ClC₂SF₄₋ (C₂ stands for a two-carbon fragment originating in a C=O or C≡C bond). For the cis-compounds no comparable results are available. The corresponding bromine adduct, cis-SF₄₋Br would, naturally, be more reactive than the chlorine adduct, as the S-halogene bond-strength in SF₅X falls in the order X = F > Cl > Br > I. For bromine no value is published, but the trend and the greater reactivity of SF₅Br as compared to SF₅Cl support this conclusion. A comparable sulfur-iodine bond has not yet been detected. When CH₂=SF₄ is allowed to react with HI, formation of iodine is observed. In only one case the existence of SF₅I has been conjectured, and in this case (the reaction: S₂F₁₀ + 2 C₂F₄ + I₂ → 2 F₅SCF₂CF₂I) it is easy to formulate an alternate mechanism. It is likely that iodine would conform to the general trend of falling S–X bond-strength.
CHAPTER IV

RESULTS AND DISCUSSION

SYNTHESIS AND REACTIONS OF F₅SC(SO₂F)=C=O

To prepare the ketene 21, F₅SC(SO₂F)=C=O, three possibilities were considered. As with 3, the ketene should be formed either from 1 (the sultone) or F₅SC(SO₂F)COF (22) and 18 or from an ester F₅SC(SO₂F)COOR (R = (CH₃)₂CH or (CH₃)₃C) and P₄O₁₀, analogously to 3. All three methods did indeed work, but only the dehydrofluorination of 22 was of practical value. 1 and 18 produced a mixture:

\[ \text{F}_5\text{SCHCF}_2\text{OSO}_2 \ + \text{F}_3\text{B} \cdot \text{N(C}_2\text{H}_5)_3 \ ----> 21 + \text{F}_4\text{S}=\text{CF}_2 \ (34\%) \]

\[ \text{Compound 23 is a novel sultone, and its reactions will be considered later (page 44). The separation of 21 and 23 was considerably difficult and made the method, besides the low yield for the ketene in the reaction, impractical.} \]

The second method employed dehydrofluorination of 22:

\[ \text{F}_5\text{SCH(SO}_2\text{F})\text{COF} + 18 ----> 21 \ (+ \text{BF}_4^- + \text{HN}^+(\text{C}_2\text{H}_5)_3) \]

22

This method worked very well, and yields of up to 95% of the ketene (of high purity) could be regularly obtained.

The third method consisted of heating an ester with phosphorus pentoxide:
Although the ketene may be synthesized in this way, there are several difficulties involved. For one, the yields in this reaction were not very consistent (~50-90%) from run to run and dropped when larger quantities were used (e.g. from 2.65 g of ester 1.6 g of crude 21 could be collected, but from 8.7 g of ester only ~1 g). Also, there was always a contamination with F₅SCH₂SO₂F, and the reaction was very time-consuming, even when only small quantities of 21 were made.

The synthesis of the ester from 1 is accomplished in the following way:

\[ \text{F₅SCH(SO₂F)COOCH(CH₃)₂ + P₄O₁₀ \rightarrow 21} \]

\[ 1 + (\text{CH₃})₂CHOH + NaF \rightarrow \text{F₅SCH(SO₂F)COOCH(CH₃)₂ (21a, 73%) + NaHF₂} \]

It is recommendable to have NaF present, as otherwise the yields were found to be lower, as for the methyl ester:

\[ 1 + \text{CH₃OH \rightarrow F₅SCH(SO₂F)COOCH₃ (21b, 35%)} \]

When one considers the yield of the (isopropyl) ester synthesis in the conversion of 1 \(\rightarrow\) 21 also, one has an overall yield of \(\approx 40\%\). This could be improved by converting only small quantities of the ester per run, but then one has an additional time-expenditure and even more phosphorous pentoxide consumption.

The dehydrofluorination of 22 requires first rearranging 1 to 22. Interestingly, 21 is found in this rearrangement as a by-product in varying ratios. The
comparatively weak base NaF acts apparently also as a dehydrofluorination reagent, but it was not possible to bring this reaction to completion. Sodium fluoride failed to give pure 21 even upon warming to 135°C (15 h); under these conditions some carbonyl fluoride and a 50:50 mixture of 21 and 22 were produced. It was also observed that the rearrangement of 1 to 22 only went well when a large excess of NaF was used; a further improvement could be achieved by powdering the NaF. In this case mixtures with more than 80% ketene contents could be obtained. The reaction vessel had to be heated in vacuo for several hours to ≈ 120°C in order to drive off the apparently absorbed 21. HF would transfer under the same conditions from NaHF₂ (a method to obtain HF of high purity).

When 21 was heated to 220°C (4h), it could be almost completely recovered (98%). It shares this property with other perfluoro ketenes.

Of interest were reactions of 21 with electrophiles and with nucleophiles; of the electrophilic reagents, SO₃ was particularly interesting, as this would not only allow one to compare the electrophilicity of 21 with 3 and 5, but could also lead to the sulfur ylide.

When 21 was subjected to sulfur trioxidation, the expected product F₄S=C(SO₂F)₂ (see reference above) was not observed:
Up to 240°C almost no reaction was observed in two days, but at 270°C rather quick formation of CO$_2$ occurred, accompanied by the formation of dark material. It is not unlikely that the SF$_4^-$ product 7 had been formed indeed but did not withstand the high temperature.

The order of electrophilicity (= the reverse order of nucleophilicity) is thus:

\((\text{CF}_3)_2\text{CCO} \ (-30^\circ\text{C}) < \text{CF}_3(\text{SO}_2\text{F})\text{CCO} \ (160^\circ\text{C}) < \ 21 \ (280^\circ\text{C})\)

The numbers in parentheses indicate the reaction temperature with SO$_3$ (p. 11).

Corresponding to this order, there is a great proneness to undergo nucleophilic additions. Whereas bistrifluoromethyl ketene undergoes F$^-$-catalyzed rearrangement (with NaF) and the intermediate (CF$_3$)$_2$C$^-$COF Na$^+$ is assumed, and a mercury compound, ((CF$_3$)$_2$C(CFO)$_2$Hg, was described in 1972 by Knunyantz et al., the corresponding anion could only recently be isolated with the counterion, ((CH$_3$)$_2$N)$_3$S$^+$, and no corresponding results are available for trifluoromethyl fluorosulfonyl ketene, it is found that anion formation is the predominant mode of reaction of pentafluorothio fluorosulfonyl ketene.

A pertinent observation was made in the NaF-catalyzed rearrangement of 1 to 22. Although both 21 and 22 are
volatile liquids, it was noticed that it took a long time
\((\approx 2 \, \text{h})\) and heating \((100-120^\circ \text{C})\) to collect most of the
product by vacuum transfer. From these observations one
assumes the formation of a complex between either 21 or 22
and NaF or NaHF$_2$. Such a complex was later indeed observed,
when the residual solid of the rearrangement 1 $\rightarrow$ 22 was
extracted with dry CH$_3$CN but could not be isolated;
according to $^{19}$F n.m.r. data it is likely that it was 21·NaF
(25e). Nuclear magnetic resonance data are given in tabular
form together with other identified complexes of the form
21·MF (Table 1, p.65). The yield of this rearrangement was
\(\approx 95\) per cent, and together with the conversion of the acyl
fluoride to 21, a total yield of \(\approx 90\%\) can be achieved in the
conversion 1 $\rightarrow$ 21. For most reactions 21, as obtained by
reaction of 22 and 18, did not require further purification.

Of the possible reactions of 21, its rearrangement to
24 did not occur, instead the compound that would have
normally been the intermediate in the reaction, 25, was
isolated as the sole product:

\[
\begin{align*}
F_5SC=CHC=O + CsF &\rightarrow \{ F_5SC^-=COF Cs^+ \} \rightarrow F_4S=CHC=O,
\end{align*}
\]

A similar intermediate was formulated in the anionotropic
re-arrangement of 5 to 8. Ketene 5a is converted to 8a with
glass, but not with CsF or KF:

\[
\begin{align*}
(CF_3)_2C=O (5) + NaF &\rightarrow CF_2=CF(CF_3)COF (8) \\
F_5SCH=CHC=O (5a) \text{ (glass)} &\rightarrow F_4S=CHCOF (8a)
\end{align*}
\]
The strongly electron withdrawing capability of the SO$_2$F group is evidenced by the formation of 25. According to an earlier investigation, the SF$_5$ group has a stronger charge-stabilizing effect than a CF$_3$ group$^{45}$; thus there are two highly electronegative substituents at an sp$^2$ carbon, which leads to the proneness for nucleophilic addition. A similar reaction was observed with silver fluoride, where the fluoride dissolved in a vigorous reaction, forming Ag$^+$ [F$_5$SC(SO$_2$F)COF]$^-$ (25d), which was light and heat sensitive. Compound 25 was investigated in more detail and will be discussed in a subsequent part.

The reactivity of 21 towards nucleophiles offered the possibility of several derivations. It should have been possible to generate an azide, which by the Curtius degradation should lead to an amine:

21 + NaN$_3$ $\rightarrow$ F$_5$SC$^-$(SO$_2$F)CON$_3$ Na$^+$ (46) $\rightarrow$ F$_5$SCH(SO$_2$F)CON$_3$

However, although it was found by $^{19}$F n.m.r. that the reaction passed through an anionic intermediate, only F$_5$SCH$_2$SO$_2$F was found as product. It must be assumed that the azide was hydrolyzed faster than it could have undergone loss of nitrogen and rearranged. Hydrolysis would have occurred when aqueous acid was added to convert the anion to the azide F$_5$SCH(SO$_2$F)CON$_3$. The carboxylic acid, formed from the azide, is not stable towards decarboxylation.

Of special interest would have been the formation of a
ketene dimer, which should have been, by analogy to 9 or the dimer of 5a, a β-lactone, with reactions similar to 9. To effect its formation, reaction with a weak base, tetramethyl urea, was tried. No dimer formation was observed, but instead a crystalline, very moisture-sensitive compound was isolated.

\[ \text{F}_5\text{SC(SO}_2\text{F)=C=O} + ((\text{CH}_3)_2\text{N})_2\text{C}=\text{O} \rightarrow 21\cdot((\text{CH}_3)_2\text{N})_2\text{C}=\text{O} \quad 26 \]

Pertinent features of this compound are the relatively low C=O stretch vibrations in the infrared spectrum and the typical chemical shifts of anionic SF₅ compounds (Table 1, p. 65), i.e., the S-F resonances in the SF₅ and SO₂F group are observed downfield from appropriate model compounds. The C=O resonance frequencies lie very close together, so that it is not possible to assign a certain infrared band to a certain carbonyl group. In principle one could infer a probable structure from that knowledge. The methyl protons are observed as a singlet in ¹H n.m.r. spectroscopy. The structure that accounts best for the properties of 26 has the oxygen bonded to the carbonyl group of ketene 21, resulting in an oxonium compound:

\[
\begin{array}{c}
\text{F}_5\text{S} \\
\text{C}=\text{C} \\
\text{O}^- \\
\text{PO}_2\text{S}
\end{array}
\]

The compound is hydrolyzed completely and instantly by water to F₅CH₂SO₂F, CO₂ and tetramethyl urea.

\[ 26 + \text{H}_2\text{O} \rightarrow \text{F}_5\text{SCH}_2\text{SO}_2\text{F} + \text{CO}_2 + \text{O}^=\text{C(N(CH}_3)_2\text{)}_2 \]
With triethyl amine and 21 a similar reaction seems to take place, but darkening occurs. The infrared spectrum of the dark residue has a C=O band at \( \approx 1700 \text{ cm}^{-1} \); in this case an ammonium salt should have been formed. The product was not isolated, however.

It is now clear that 21 could be made by the above dehydrofluorination of 22 only because \( \text{F}_3\text{B} \cdot \text{N(C}_2\text{H}_5)_3 \) is a base of low nucleophilicity, and did consequently not react with the ketene.

An attempt to add methoxide ion to 21 was partially successful; it was observed (by \( ^{19}\text{F} \) n.m.r. spectroscopy) that two anionic species were formed, one of them containing a COF group (probably \( \text{Na}^+ [\text{F}_5\text{S} \cdot \text{SC(SO}_2\text{F})\cdot \text{COF}]^- \)). The reaction was vigorous and the (CH\(_3\)CN) solution darkened at the same time. The second anion (the major product) that was formed must have been the sodium salt,

\[
\begin{align*}
\text{F}_5\text{S} & \quad \text{OCH}_3 \\
\text{C} = \text{C}=\text{O} + \text{NaOCH}_3 & \quad \text{C} = \text{C} \quad \text{Na}^+ \\
\text{FO}_2\text{S} & \quad \text{FO}_2\text{S}
\end{align*}
\]

The reaction could not be controlled so as to allow for the formation of the ester enolate alone, but these compounds can be obtained very efficiently by another route. A further reaction is the addition of \( \text{HgF}_2 \) to 21. This should provide for the compound \( \text{F}_5\text{S} \cdot \text{SC(SO}_2\text{F})\cdot \text{COF})_2\text{Hg} \) (45). Mercury compounds offer some interesting possibilities, as found with other perfluoromercurials\(^{46} \), especially the formation of sulfides.
These could then be converted to another SF₅ compound, viz:

\[(\text{F}_5\text{SC(SO}_2\text{F})(\text{COF})\text{CS})_2\text{Hg} \rightarrow (\text{F}_5\text{SC(SO}_2\text{F})(\text{COF})\text{S})_2\text{Hg} \rightarrow \]
\[(\text{F}_5\text{S(SO}_2\text{F})(\text{COF})\text{CS})_2 \rightarrow (\text{F}_5\text{S(SO}_2\text{F})(\text{COF})\text{CS})_2 \rightarrow \]
\[(\text{F}_5\text{S}_2\text{C(SO}_2\text{F})\text{COF}), \text{or a bis-sulfuryl derivative, e.g.,} \]
\[(\text{F}_5\text{SCCOF(SO}_2\text{F})_2), \text{could be obtained by oxidation of the mercurial.} \]

The ¹⁹F n.m.r spectrum of the product of this reaction, in which HgF₂ dissolves in CH₃CN in the presence of 2₁, is indicative of salt formation (see table 1 p. 65), and it can be isolated as a white amorphous solid by pumping off the solvent (CH₃CN). The i.r. spectrum indicates that there is a CH₃C=N - 4₅ complex, and the melting behavior shows gradual softening with final melting at ≈ 80°C.

Further reactions of 2₁ involved the addition of other nucleophiles. With water, the following reaction took place:

\[2₁ + \text{H}_2\text{O} \rightarrow (\text{F}_5\text{SCH(SO}_2\text{F})\text{COOH}) \rightarrow \text{F}_5\text{SCH}_2\text{SO}_2\text{F} + \text{CO}_2 \]

When 2₁ was exposed to air, a solid formed, which, when pressed on KBr plates, evolved a gas. This solid must have been the intermediate carboxylic acid. It is often observed that carboxylic acids that contain two strongly electronegative substituents at Cα are not stable towards decarboxylation. Malonic acid behaves in this manner, as do other β-carbonyl carboxylic acids, as well as the half ester of trifluoromethyl malonic acid.⁴⁷

Alcohols combined with 2₁ to form esters:

\[2₁ + (\text{CH}_₃)_₂\text{CHOH} \rightarrow \text{F}_5\text{SCH(SO}_2\text{F})\text{COOCH(CH}_₃)_₂ \text{2₁a} \]
\[ 21 + \text{CF}_3\text{CH}_2\text{OH} \longrightarrow F_5\text{SCH(SO}_2\text{F)}\text{COOCH}_2\text{CF}_3 \]
\[ 21 + (\text{CH}_3)_2\text{CHOH} \longrightarrow F_5\text{SCH(SO}_2\text{F)}\text{COO(CH(CH}_3)_2} \]
\[ 21 + (\text{CH}_3)_3\text{COH} \longrightarrow F_5\text{SCH(SO}_2\text{F)}\text{COO(CH}_3)_3 \]

Interestingly, the t-butyl ester was not a stable compound; it decomposed upon distillation, and \( F_5\text{SCH}_2\text{SO}_2\text{F} \) was found as a product. This decomposition occurred also at lower temperature, and a purple product resulted at room temperature after a few days.

With \( \text{P(OCH}_2\text{H}_5)_3 \) only a complex mixture was found, and a major product was an acyl fluoride. The products could, however, not be separated nor identified. A similar reaction with \( 5 \) and \( \text{P(OCH}_2\text{H}_5)_3 \) led to tetrakistrifluoromethyl allene\(^{48} \).

\[ 21 + \text{P(OCH}_2\text{H}_5)_3 \longrightarrow \text{complex mixture} \]

Diethyl amine formed the amide:

\[ 21 + \text{HN(C}_2\text{H}_5)_2 \longrightarrow F_5\text{SCH(SO}_2\text{F)}\text{CON(C}_2\text{H}_5)_2 \quad (47\%) \]

The low yield is related to the vigorous nature of the reaction; although the reactants were brought together at low temperature in \( \text{CH}_2\text{Cl}_2 \) solution, the formation of a lot of dark tarry product could not be prevented. The amide \( 27 \) shows two \( \text{CH}_3 \) groups in the \( ^1\text{H} \) n.m.r. spectrum, and the methylene protons appear as a multiplet. This is certainly an indication of the hindered rotation that is found around the C-N bond. The structure of this solid was determined by X-ray crystallography (p.71).

\[ 21 \] reacted readily and cleanly with trimethyl silane
upon warming to room temperature to give a white solid that was highly susceptible to moisture and exhibited a repugnant odor. After sublimation it was obtained as colorless regular crystals. The structure of this material is not certain. Addition of the silicon could have taken place either at the oxygen or the carbon of the carbonyl group, resulting in the following compositions:

\[
\begin{align*}
39a: & \quad \text{F}_5\text{S} \quad \text{C}==\text{C} \quad \text{H} \quad \text{OSi(CH}_3)_3 \\
39b: & \quad \text{F}_5\text{S} \quad \text{C}==\text{C} \quad \text{Si(CH}_3)_3 \\
\end{align*}
\]

A distinction between these two structures is difficult. As mentioned before, some SF₅ olefins have very low i.r. frequencies and intensities. Trimethylsilylketones have also unusually low i.r. frequencies (1600-1640 cm⁻¹) (see reference below). A distinction between 39a and 39b is therefore not possible on the basis of the infrared spectrum. Most trimethylsilyl ketones are yellow⁴⁹, and the adduct 21·HSi(CH₃)₃ is colorless. One distinctive property of trimethylsilyl ketones is the low field position of the carbonyl group in the ¹³C n.m.r. spectrum⁵⁰, as compared to normal carbonyl compounds. The C=O resonance is usually observed between 220 and 240 ppm (Si(CH₃)₄ = 0). In the ¹³C n.m.r. spectrum of 21·SiH(CH₃)₃ no such resonance is seen. One has to assign the structure of the adduct to 39a.

Other reactions of 21 were investigated. With SF₅Br a transformation was observed that resulted mainly in BrF addition:
21 + SF₅Br \rightarrow F₅SCBr(SO₂F)COF (28) + [SF₄]

28 could not be obtained in pure form from this reaction; all attempts to free it from attendant compounds failed, but there is another route leading to it that yields a purer product. This will be described in the chapter dealing with 25. An intermediate formation of a crystalline compound was observed; although not identified, this could correspond to the intermediate formation of F₅SCBr(SO₂F)C(O)SF₅.

Trioxane, used as a source of formaldehyde, reacts with 21 (at 150°C) to give initially a primary product. The colorless liquid that is obtained is characterized by a strong, although not unpleasant, pungent odor. Normally, by analogy to 5, this product should be a β-lactone,

\[
\begin{array}{c}
\text{F}_5\text{S} \\
\text{FO}_2\text{S} --\text{C}=\text{O} \\
\text{H}_2\text{C} --\text{O},
\end{array}
\]

but the absence of a carbonyl band in the i.r. spectrum does not support this composition. On the other hand, there is a very weak band at 1616 cm⁻¹. If this is taken as an indication for a double bond (some SF₅ olefins show very weak or no C=C bands at all⁵¹) together with the information from the proton n.m.r spectrum (two resonances of equal intensity but different couplings at ≈ 7 ppm) and the mass spectrum, and taking into account its rather high volatility (i.e., not a polymer), one is led to assuming the following structure:

\[
\begin{array}{c}
\text{F}_5\text{S} \\
\text{FO}_2\text{S} \\
\text{C} = \text{C} --\text{CH}_2 \\
\end{array}
\]
This formulation would also fall into line with the reactions that are observed with other reagents, i.e. nucleophilic attack at C=O:

\[ \text{F}_5\text{S(SO}_2\text{F})=\text{C}=\text{O} + \text{CH}_2^+\text{-O}^- \rightarrow \text{F}_5\text{S(SO}_2\text{F})=\text{C}-\text{O}^- \rightarrow \text{28} \]

\[ \text{O-CH}_2^+ \]

The compound exhibits a very unusual instability. The elemental analysis could not be obtained; a white solid separates from the liquid after some time. When the liquid is stored in a capillary, the whole volume of the liquid appears to be replaced by the white solid within weeks, even when stored at -12°C. The solid is barely soluble in CDC\text{\textsubscript{3}}, but is soluble in acetone and in CD\text{\textsubscript{3}}CN. The amorphous material was converted to crystals by very slow evaporation of an acetone solution. A crystal was used for X-ray analysis and the compound was found to have the following structure:

\[ \text{F}_5\text{S-CH-CH}_2\text{-O-CH}_2\text{-CH-SF}_5 \]

\[ \text{F}_2\text{OS} \quad \text{SO}_2\text{F} \]

\[ \text{29} \]

A mechanism for this conversion \text{28} \rightarrow \text{29} is not yet formulated.

Support for this structure comes also from the \textsuperscript{1}H n.m.r spectrum, with a multiplet at \( \approx 5.5 \) ppm, and an AB system at \( \approx 4.4 \) ppm (ratio \( \approx 1:2 \)).

When \text{21} and acetone were allowed to react at room temperature, several products were formed. The main product seems to be an ester:

\[ \text{21} + \text{CH}_3\text{COCH}_3 \rightarrow \text{F}_5\text{SCH(SO}_2\text{F})\text{COOCH(CH}_3\text{)}=\text{CH}_2 \]

\[ \text{30} \]
The product has a C=C (1683 cm\(^{-1}\)) and a C=O (1776 cm\(^{-1}\)) band in the i.r. spectrum. The remainder of the spectrum looks almost identical to the spectrum of \(\text{F}_{5}\text{SCH(SO}_{2}\text{F})\text{COOCH(CH}_{3}\text{)}_{2}\), \text{21a}.

A further similarity is seen in the \(^{19}\text{F}\) n.m.r. spectrum, with a collapsed AB\(_4\) resonance (SF\(_5\)) and an SO\(_2\text{F}\) signal with nearly the same chemical shifts. The \(^{1}\text{H}\) n.m.r spectrum has a multiplet at 5.74 ppm, a vinyl resonance at 4.89 ppm and a methyl band at 2.00 ppm.

This reaction must be brought about through the keto-enol equilibrium of acetone, where the enol would be constantly removed from the mixture. \text{30} could be isolated only in trace amounts.

Despite the close kinship of CH\(_2\)=O and (CH\(_3\))\(_2\)C=O, two completely different reactions are observed. One reason could be that formaldehyde does not exist in a keto-enol equilibrium. From the \(^{19}\text{F}\) n.m.r. spectrum of the crude product, there is no indication that the dimethyl anlaogue of \text{28} is present. If the primary product of \text{21} and CH\(_2\text{O}\) is indeed \text{28}, then it is surprising not to obtain the analogous product with acetone. If \text{28} were formed by nucleophilic addition to \text{21}, then acetone should react more rapidly, due to the two +I substituents at the carbonyl group. This course must then be slowed down due to either steric hindrance or the much faster addition of OH (CH\(_2\)=CHOH(CH\(_3\))) to the ketene. No reaction between \text{21} and hexafluoroacetone.
was observed.

THE FORMAL ADDITION OF RO⁻ to F₅SC(SO₂F)=C=O; SYNTHESES:

AND REACTIONS OF F₅SC⁻(SO₂F)COOR

If one applies now the formal relationship between ketenes and enolates,

\[ RR'\ C\!\!\ =\ C=O + X^- \rightarrow RR'\ C\!\!\ C=\!\!\ X, \]

one arrives at an extension of the reactions of 21.

As described above, the synthesis of these enolates from the ketene is not practical, but there is a very convenient route, analogous to 10.

The methyl and isopropylester of the nonisolable acid F₅SCH(SO₂F)COOH easily form salts with triethylamine (in ether or acetone):

\[ F₅SCH(SO₂F)COOR + N(C₂H₅)₃ \rightarrow [HN(C₂H₅)₃]^+ [F₅SC(SO₂F)COOR]^- \]

32a,b

32a, R = CH(CH₃)₂; 32b, R = CH₃

The triethylammonium salts are collected as white amorphous solids and can be recrystallized from either CH₂Cl₂ or CHCl₃, giving large platelike crystals. It was possible to determine the crystal structure of the methyl ester. The solutions of these compounds are yellowish.

These triethylammonium salts show only low reactivity. Protonation takes place with strong acid immediately, but
only slowly with water:
\[ 32a + HCl \rightarrow F_5SCH(SO_2F)COOCH(CH_3)_2 + ([HN(C_2H_5)_3]^+Cl^-) \]

Two days of refluxing the isopropylester salt with n-pentyl bromide in acetone left the \(^{19}\)F n.m.r. spectrum of the reaction mixture unchanged. An attempt to brominate the same salt led to the isolation of the starting ester. It is clear now from experiments with the sodium salt that the bromo ester is unstable towards hydrolysis. The formation of the starting ester in this case can thus be attributed to the work-up procedure, which involved at one point precipitation with water.

An ammonium salt, \(NH_4^+ [F_5SC(SO_2F)COOCH(CH_3)_2]^- \), 34, was also obtained by treating the isopropyl ester with gaseous ammonia in diethyl ether at lowered temperatures. The sulfonamide could not be obtained by this method, but in dioxane solution, at \(\approx 50-60^\circ\)C (by the heat of reaction), the sulfonyl fluoride was converted to the sulfonamide:

\[ F_5SCH(SO_2F)COOCH(CH_3)_2 + NH_3 \rightarrow F_5SC^- (SO_2F)COOCH(CH_3)_2 NH_4^+ \]

\[ \rightarrow F_5SC(SO_2NH_2)COOCH(CH_3)_2 + (NH_4F) \]

This conversion describes one of the few reactions these pentafluorosulfur fluorosulfonyl alkyl acetates undergo. It was also thought that by having a different cation it should be possible to induce reaction with halogen-bearing
compounds.

Of the three ways to obtain these anions with metal cations, only one was practicable. The keten \( \text{F}_5\text{SC(SO}_2\text{F)=C=O} \) reacted too vigorously with \( \text{NaOCH}_3 \), as described above:

\[
\begin{align*}
\text{F}_5\text{S} & \quad \text{C}=\text{C}=\text{O} + \text{NaOCH}_3 \quad \text{--CH}_3\text{CN}----> \quad \text{C}^- \text{C} \quad \text{Na}^+ \\
\text{FO}_2\text{S} & \quad \text{FO}_2\text{S} \\
\end{align*}
\]

A second route would have led, by analogy to \( \text{CF}_2=\text{C(CF}_3\text{)SO}_2\text{F} \) (16), to \( \text{F}_4\text{S}=\text{C(SO}_2\text{F)COOR} \) \( \text{R} = \text{CH(CH}_3\text{)}_2, \text{CH}_3 \), by treating the ester with the base \( \text{F}_3\text{B} \cdot \text{N(C}_2\text{H}_5\text{)}_3 \).

\[
21\text{a} + \text{F}_3\text{B} \cdot \text{N(C}_2\text{H}_5\text{)}_3 \quad \text{---X----> F}_4\text{S}=\text{C(SO}_2\text{F)COOCH(CH}_3\text{)}_2 \quad 21\text{c}
\]

Treatment of this \( \text{SF}_4= \) compound with \( \text{CsF} \) should have led to addition of \( \text{F}^- \) at the sulfur, analogously to 17. That the sulfur is an electrophilic center in sulfur \( \text{(SF}_4= \) ylides has been shown by Seppelt et al. earlier29, and such an addition direction is now quite well established. This reaction led from the very outset to a completely different product and is thus not a possible route to the ester anion salts. In fact, the isolated product was \( \text{(F}_2\text{S}=\text{O})=\text{C(SO}_2\text{F)COF} \)

\[
\text{F}_5\text{SCH(SO}_2\text{F)COOCH}_3 + \text{F}_3\text{B} \cdot \text{N(C}_2\text{H}_5\text{)}_3 \quad \text{--> (F}_2\text{SO})=\text{C(SO}_2\text{F)COF} \quad 44 \quad + \quad \text{CH}_3\text{F}
\]

It was possible to obtain easily the sodium salt without attendant by-products from the isopropyl ester with sodium isopropoxide in diethyl ether:
Isolation of this salt was not possible, and it showed a reluctance to react, similar to the triethyl ammonium salts. After each reaction, except for chlorine, where a different product was isolated, about 50% of the original ester was recovered. This was the case for the reactants BrCN, CH₂=CHCH₂Br, CH₃COCl (after removal of the isopropanol formed in the reaction with sodium isopropoxide), Cl₂, Br₂, and ClCOOCH₃. The reaction with CH₂Br₂ was studied in an n.m.r. tube experiment. The reaction conditions varied in case to case from stirring at room-temperature to refluxing.

The recovery of 21a from the reaction of 33 with BrCN, ClCOOCH₃, CH₂=CHCH₂Br, CH₃COCl and its detection by ¹⁹F n.m.r. spectroscopy in the case of CH₂Br₂ is of some interest. In all cases the formation of a white precipitation was observed. It must be assumed, that part of the anion decays with the release of HF, which would then protonate the anion. It was shown in a separate experiment, that the reversion to the ester takes place even without any substrate being present:

\[ 33 \rightarrow \text{F}_5\text{SCH(SO}_2\text{F})\text{COOCH(CH}_3\text{)}_2 \ (53\%) \]

Isolated yields of the ester were 40-60%, respectively in the reactions with and the reaction without a substrate. How this degradation of the anion would take place remains an open question. There could be a similarity to the related
anion \((\text{CF}_3)_2\text{C}^-\text{SO}_2\text{F}\) (14), which can lose \(\text{F}^-\) from the \(\text{SO}_2\text{F}\) group\(^{27}\).

All of these salts were characterized by the now-familiar low-field position of the \(\text{SF}_5\) and \(\text{SO}_2\text{F}\) group in the \(^{19}\text{F}\) n.m.r. spectrum and the yellowish color of their solutions.

A ureide 31, obtained by reacting 1 with urea,

\[
\begin{align*}
1 + \text{H}_2\text{NCONH}_2 & \rightarrow \underset{\text{FO}_2\text{S}}{\text{F}_5\text{SCHCONHCONH}_2} 31,
\end{align*}
\]

reacted also with triethyl amine to give a salt 36. It was, however, only of marginal stability and could only be characterized by i.r. and fluorine n.m.r. spectroscopy. Again, the low CO frequency and the low-field position of the fluorines indicate the presence of an anion (table 1). Upon storing its solutions at room temperature other (SF\(_5\)) products were formed, but their nature was not investigated. It was because of the formation of 36 that it was not possible to cyclize 31:

\[
\begin{align*}
31 + \text{N(\text{C}_2\text{H}_5)_3} & \rightarrow \underset{\text{CO}}{\text{\text{F}_5\text{SCH}_2\text{NH}}},
\end{align*}
\]

" " " \(\rightarrow \underset{\text{\text{CO}}}{\text{\text{F}_5\text{SC}^-\text{(SO}_2\text{F})\text{COONHCONH}_2}}\text{HN(\text{C}_2\text{H}_5)_3} + 36\)

On the other hand, both the chlorination and bromination of 33 led to the formation of new compounds:

\[
\begin{align*}
33 + \text{Cl}_2 & \rightarrow \underset{\text{\text{Cl}}}{{\text{F}_5\text{SCCOOCH(CH}_3)_2}} 37 + \text{NaCl},
\end{align*}
\]

\[
\begin{align*}
33 + \text{Br}_2 & \rightarrow \underset{\text{\text{Br}}}{{\text{F}_5\text{SCCOOCH(CH}_3)_2}} 38 + \text{NaBr}.
\end{align*}
\]
While the chloroester 37 could be isolated and is chemically stable, the bromo derivative 38 is highly sensitive towards hydrolysis. When it was tried to remove excess bromine by stirring with Na$_2$S$_2$O$_3$ solution, it was completely consumed and only a compound with the spectral characteristics of F$_5$SCH(SO$_2$F)CH(CH$_3$)$_2$ was isolated. Because this conversion could not be completely suppressed, even by using non-aqueous conditions, it was not possible to obtain 38 in pure form. This peculiar instability must be related to bond-polarization by the strongly electron withdrawing fluoroalkyl residue. A positively polarized bromine atom should then be easily attackable by water:

\[
\begin{align*}
\text{F}_5\text{S} & \quad \text{Br}^+ \\
\text{FO}_2\text{S} & \quad \text{C} \quad \text{OR} \\
\rightarrow & \\
\text{F}_5\text{S} & \quad \text{C} \quad \text{OR} \\
\text{FO}_2\text{S} & \quad \text{Br}^-, [\text{HOBr}]
\end{align*}
\]

The CF$_3$ analogue of 38, 11a, is, on the other hand, perfectly stable.

One can assume that the environment of the α-carbon in 33 is so crowded that an $S_N2$ reaction is not normally possible, except for chlorine and bromine, where attack can occur along the X-X axis. This could also in principle occur for BrCN, with the consequent formation of an isonitrile, but the reaction might go via an addition-elimination mechanism:

\[
\begin{align*}
\text{X}^- + \text{C}≡\text{N} & \longrightarrow \text{X}-\text{C}≡\text{N}^- \\
\text{Br} & \quad \text{Br}
\end{align*}
\]

which is probably sterically too demanding.
The sterically more amenable O-substitution, resulting in the formation of ketene acetals, was not observed.

\[
\text{F}_5\text{S} \overset{\text{C}==\text{C}}{\underset{\text{O}^-}{\longrightarrow}} \text{F}_5\text{S} \overset{\text{C}==\text{C}}{\underset{\text{OR'}}{\longrightarrow}} \text{F}_5\text{S} \overset{\text{C}==\text{C}}{\underset{\text{OR}}{\longrightarrow}}
\]

It was now of interest to compare the ease of formation of various anionic species under the same conditions. The compounds \(\text{F}_5\text{SCH}_2\text{SO}_2\text{F}\)\(^{52}\), \(\text{F}_5\text{SCH}_2\text{COOCH}_3\)\(^{53}\), \(\text{F}_5\text{SC}H\text{FSO}_2\text{F}\)\(^{54}\), \(\text{F}_5\text{SCH(SO}_2\text{F})\text{COOC}(\text{CH}_3)_2\), \(\text{F}_5\text{SCH}(\text{SO}_2\text{NH}_2)\text{COOCH(}\text{CH}_3)_2\) and \(\text{F}_5\text{SCH(SO}_2\text{F})\text{CONHCONH}_2\) were dissolved in acetone in an n.m.r. tube and a drop of triethyl amine was added, causing compounds with the three functional characteristics of an \(\text{SO}_2\text{F}, \text{SF}_5\) and \(\text{COR} (R = \text{OCH(}\text{CH}_3)_2, \text{NHCONH}_2)\) group to exhibit the downfield shift of the \(\text{SF}_5\) and the \(\text{SO}_2\text{F}\) fluorine resonances in anions. In the case of \(\text{F}_5\text{SCHFSO}_2\text{F}\) the fluorine signals were diminished after base addition, and slow darkening occurred, but no signal attributable to an anion could be detected. Indeed, the sequence for CH-acidity in the analogous CF\(_3\) series has CF\(_3\)CHFSO\(_2\)F as its least acidic member\(^{10}\):

\[
\text{CF}_3\text{CHFSO}_2\text{F} < \text{CF}_3\text{CH}_2\text{SO}_2\text{F} < \text{CF}_3\text{CH(SO}_2\text{F})\text{COOC}\text{H}_3, \text{CF}_3\text{CHClSO}_2\text{F}.
\]

Fluorine as a +M,-I substituent should stabilize the anionic site less than chlorine, a weaker (-M,-I) substituent\(^{10}\).

Neither with \(\text{F}_5\text{SCH}_2\text{COOCH}_3\) nor with \(\text{F}_5\text{SCH(SO}_2\text{NH}_2)\text{COOCH(}\text{CH}_3)_2\)\(^{35}\) was any reaction observed. In \(35\), the \(\text{SO}_2\text{F}\)-fluorine was replaced by another +M,-I substituent. Apparently, nitrogen is a stronger +M and a
weaker -I substituent which leads to the decrease in charge stabilizing capability. With the ester $F_5SCH_2COOCH_3$ it was also not possible to induce reaction with formaldehyde in the presence of diethylamine (Tollens reaction), showing again that it has only a poor ability to form an anion. The extraordinary stability of 25 is additional evidence that the substituent at the C=O group is of minor importance and that the central carbon just needs to be flanked by the $SF_5$, the $SO_2F$ and the C=O group in order to form salts.

The synthetic usefulness of these ester anions is small, because of their great inertness. They make it possible, however, to obtain derivatives of the sultone

$$\begin{align*}
\text{Cl} & \quad F_5SC-\text{CF}_2 39 \\
\text{O}_2S-\text{O} & \quad \text{F}_5\text{CCl=CF}_2
\end{align*}$$

which could not be obtained by the usual sulfur trioxidation of the olefin $F_5\text{CCl=CF}_2$, because loss of $SF_5$ and the formation of unknown products ensued. As described below, it is possible to synthesize 39 by a different and overall tedious route, which is of little synthetic value. It is, however, very easy to prepare derivatives of 39 by chlorination of 33, with possible further transformations. The bromo ester 38 might be of similar usefulness, as it should be possible to couple it with olefins in a radical-type reaction.
The reaction of 21 with metal fluorides results in stable complexes. The scope of this reaction has not been completely charted, but with cesium fluoride, silver fluoride and mercury fluoride, vigorous reactions occur when carried out in CH₃CN, causing the fluorides to dissolve with the boiling of the liquid mixture:

\[ \text{Me} = \text{Ag} \ (25\text{d}), \ 
\text{Cs} \ (25), \ 
\text{Na} \ (25\text{e}), \ 
\text{Hg} \ (\text{HgF}_2) \ (45) \]

It has been found that NaF will also form such a complex (25e); when the sodium fluoride that was used in the rearrangement of 1 to 22 was extracted with CH₃CN after removal of the product, a soluble SF₅ compound was obtained that had a very similar \(^{19}\text{F} \) n.m.r. spectrum as 25 (F₅SC(SO₂F)COF-Cs⁺) (table 1). The cesium salt can also be obtained by heating 21 and CsF without solvent; it can then be obtained by extracting the solid residue with acetone. 25 is of an extraordinary stability. It can be heated for hours in the vacuum to > 200°C without significant weight loss, although darkening occurs. The trace of material that distills off is not 21, showing the irreversibility of the reaction of 21 and CsF.

Compound 25 was obtained in quantitative yield, when 21 reacted in excess with cesium fluoride. The initial vivid
reaction (in CH₃CN) slowed down very considerably with time. While 25 could be freed from solvent by vacuum pumping, the sodium and silver salts assumed a glassy consistence and never reached the predetermined weight loss. It must therefore be assumed that they retain solvent. The same observation is made for the Hg(II) complex.

Acetone and acetonitrile were excellent solvents for 25. It was possible to precipitate it from concentrated acetone solutions by addition of methylene chloride. It was not necessary to observe any special precautions; a sample of 25 was kept in the open air for several weeks at circa 30°C, and its i.r. spectrum was unchanged after this time. Although it could be slowly converted to cesium fluoride by reaction with water and the formation of the volatile SF₅CH₂SO₂F, it is clearly demonstrated that it has a very unusual stability for a carbanion and an acyl fluoride. Recrystallization was possible from both acetone and acetonitrile, where conglomerates of crystallites were formed. Unfortunately, these crystals, as far as they could be separated from the conglomerate, were twinned, which excluded an X-ray crystallographic investigation. The ¹⁹F n.m.r. spectrum will be discussed in the n.m.r. section, and the i.r. spectrum showed clearly the presence of an anion, with a comparatively low C=O stretch frequency (1766 cm⁻¹), about 90 cm⁻¹ lower than in the neutral 22.

Although it was possible that the silver salt could
undergo a facile coupling reaction with alkyl halides, it showed only low reactivity and only complicated product mixtures could be obtained. Particularly conspicuous was the disappearance of the COF signal that was observed when the reactions were followed by $^{19}$F n.m.r. spectroscopy. Although a copious amount of AgI was formed with n-propyl iodide, the respective SF$_5$ products could not be identified. The reactions of the silver salt were not further investigated.

The cesium salt showed a similar reluctance to react; for example, when 25 was refluxed in acetone with BrCN, only SF$_5$CH$_2$SO$_2$F could be identified as a reaction product.

$$25 + \text{BrCN} \rightarrow\rightarrow \text{F}_5\text{SC}_{\text{H}_2}\text{SO}_2\text{F}$$

This must be a consequence of slow hydrolysis due to small amounts of water being present. Simpler reactants did, however, react well with it. Chlorine formed F$_5$SCCl(SO$_2$F)COF in very good yield and high purity, F$_5$SCBr(SO$_2$F)COF was obtained in only fair yield and F$_5$SCI(SO$_2$F)COF could not be detected:

$$\text{F}_5\text{SC} = \text{Cs}^+ + \text{Cl}_2 \rightarrow \text{F}_5\text{SC} - \text{Cl}^- \rightarrow >90\% + [\text{CsCl}]$$

$$\text{F}_5\text{SC} = \text{Br}_2 \rightarrow \text{F}_5\text{S} - \text{Br}^- \approx 50\% \text{ (impure)} + [\text{CsBr}]$$

$$\text{F}_5\text{SC} = \text{I}_2 \rightarrow \text{F}_5\text{S} - \text{I}^- \rightarrow + [\text{CsI}]$$

Whereas the reactions with chlorine and bromine were run without solvent, the reaction with iodine was tried with and
without the presence of CH$_3$CN. Running the reaction without solvent simplifies the isolation process. An interesting aspect is that 25 reacts with bromine in a different way in CH$_3$CN than it does without solvent. When Br$_2$ is added to a concentrated (to simplify removal of solvent) CH$_3$CN solution of 25, quick discoloration of the brown Br$_2$ color is observed, accompanied by the formation of gas bubbles and a white precipitate. The solution can be titrated in this fashion until the brown bromine color persists. The product that is obtained has no more CO group (i.r.), and shows a conspicuous upfield shift in the $^{19}$F n.m.r. spectrum. Such a shift is usually observed when electronegative elements are attached to the SF$_5$-bearing carbon atom. Further characteristics of this compound, a colorless solid when pure, are its high volatility (it distills with acetonitrile), its halogen-like odor and its low melting point (~40°C); it dissipates rapidly in air. This compound was identified as:

\[
\begin{array}{c}
\text{F}_5\text{S}-\text{C}-\text{SO}_2\text{F} \\
\text{Br}
\end{array}
\]

It also seems to react very readily with mercury; when a sample was shaken with mercury, warming was noticed and only a small amount of material could be retrieved. In a similar mode, destruction was observed with AgF.

This rather interesting result offered the
possibility of synthesizing the compounds $F_5SCCl_2SO_2F$ and $F_5SCF_2SO_2F$ by a similar route. With chlorine, a different result was obtained; while the reaction stops at the stage of $F_5SCCl(SO_2F)COF$ without irradiation, it continues under the irradiation of a sunlamp to give one major product, which is an acyl chloride. This product did not seem to undergo any further changes. The solvent was necessary for the reaction; without CH$_3$CN present only $F_5SCCl(SO_2F)COF$ was formed, even after long irradiation. The progress of the reaction without solvent could also be conveniently monitored by the drop in chlorine pressure: no further pressure drop was observed after 1 equivalent of Cl$_2$ was consumed: a dichloro compound should have consumed two equivalents and produced either CO or COClF as a by-product. Neither was detected:

\[
\begin{align*}
\text{--hv, no solvent} & \rightarrow F_5SCl(SO_2F)COF \\
25 + Cl_2 & \rightarrow \text{hv, CH}_3\text{CN} \rightarrow F_5SCCl(SO_2F)COCl \ (25g) \\
& \rightarrow \text{no hv, CH}_3\text{CN} \rightarrow F_5SCCl(SO_2F)COF
\end{align*}
\]

According to n.m.r. spectral comparisons, one by-product in the reaction is $F_5SCHClSO_2F$, and another minor product has an n.m.r. spectrum very similar to $SF_5CBr_2SO_2F$, so that the formation of $F_5SCCl_2SO_2F$, which should also have a similar $^{19}F$ n.m.r. spectrum cannot be excluded entirely. From the analogous reaction with fluorine one would have expected the compound $F_5SCF_2SO_2F$. The compound $F_5SCHFSO_2F$ shows base sensitivity (see above), and recently a similar
sensitivity of $\text{F}_5\text{SCH}_2\text{CF}_2\text{SO}_3\text{H}$ towards aqueous NaOH was found with the loss of the SF$_5$ group and evolution of gas. This is most certainly connected to the hydrogen at C$_a$, which should be easily removable, and the resulting anion, not having the possibilities of charge dispersal like $32a$, $32b$ or $33$, could then decompose. $\text{F}_5\text{SCF}_2\text{SO}_2\text{F}$ would not undergo such a base attack and it is expected to be convertible into a strong and very stable acid. The different course of this reaction with chlorine makes it unlikely that the reaction with fluorine would go in the desired direction. This would have furnished a particularly easy route to $\text{F}_5\text{SCF}_2\text{SO}_2\text{F}$.\textsuperscript{55}

The compounds $\text{F}_5\text{SCCl}(\text{SO}_2\text{F})\text{CFO}$ and $\text{F}_5\text{SCBr}(\text{SO}_2\text{F})\text{CFO}$ hydrolyzed slowly in water:

$$\text{F}_5\text{SC}(\text{Cl,Br})(\text{SO}_2\text{F})\text{CFO} + \text{H}_2\text{O} \rightarrow \text{F}_5\text{SCH}(\text{Cl,Br})\text{SO}_2\text{F} + \text{HF} + \text{CO}_2$$

Br = $25j$, Cl = $25h$

It was found that $\text{F}_5\text{SCBr}(\text{SO}_2\text{F})\text{CFO}$ does react with $\text{C}_2\text{H}_4$ in the presence of a free radical initiator, but loss of the COF group was observed:

$$\text{F}_5\text{SCBr}(\text{SO}_2\text{F})\text{COF} + \text{C}_2\text{H}_4 + (\phi\text{COO})_2 \rightarrow \text{F}_5\text{SCH}_2\text{SO}_2\text{F} \text{ (chiefly)} + \text{F}_5\text{SCH}_2\text{CH}_2\text{Br} \text{ SO}_2\text{F}^\prime$$

(25i)

The system avoids further substitution at C$_a$ by decarboxylation.

The chemical behavior with other cations was tried and reaction with $\text{NO}^+\text{BF}_4^-$ and $\text{NO}_2^+\text{BF}_4^-$ was observed. Nitrosonium tetrafluoroborate reacted rather cleanly to one major product, but it was not possible to isolate it. With CH$_3$CN
codistillation occurred, and the product appeared to have been only marginally stable, as a slow change of the n.m.r. spectrum of the acetonitrile solution suggested. This solution had a faint blue color in the cold. A blue color is typical for nitroso compounds.

A downfield shift of the COF and an upfield shift of the SF₅ and SO₂F resonances were observed, as compared to 25. From these observations one can safely assume that the product that was obtained was a nitroso compound indeed:

\[
25 + \text{NO}^+ \text{BF}_4^- \rightarrow \begin{array}{c}
\text{F}_5\text{S} \\
\text{FO}_2\text{S} \\
\end{array} \text{NO} \rightarrow \begin{array}{c}
\text{F}_5\text{S} \\
\text{FO}_2\text{S} \\
\end{array} \text{CNO}
\]

The latter decarbonylated product (which might exist as the tautomeric oxime) is assumed to be the compound that was observed when the initial adduct, which showed a COF resonance in the \(^{19}\text{F}\) n.m.r. spectrum, was kept in CH\(_3\)CN solution for several hours. Addition of water to the solution resulted in the formation of SF\(_5\)CH\(_2\)SO\(_2\)F. The oxime or the nitroso compound must have been hydrolytically unstable. This is in line with the compound (F\(_3\)C\(_2\))C(NO)COF, obtained from (CF\(_3\))\(_2\)C=C=O and NOF.\(^{56}\) Hydrolysis leads in this case to (CF\(_3\))\(_2\)CHCOOH (plus traces of (CF\(_3\))\(_2\)C=NOH). Hydrolysis of 40 should also result in the formation of SF\(_5\)CH\(_2\)SO\(_2\)F, as the intermediate acid F\(_5\)SC(NO)(SO\(_2\)F)COOH would probably be unstable too (like (F\(_3\)C\(_2\))C(NO)COOH), and, after hydrolysis of the NO group, decarboxylate, while (CF\(_3\))\(_2\)CHCOOH is stable and does not undergo loss of CO\(_2\).
The reaction of 25 and nitronium tetrafluoroborate was much slower and resulted in a mixture of compounds as evidenced by the fluorine n.m.r. spectrum of the product. The slowness of the reaction could be attributed to the apparently low solubility of NO$_2^+$BF$_4^-$ in CH$_3$CN. Nevertheless, the $^{19}$F n.m.r. spectrum leads to the conclusion that at least one COF resonance is present; this should correspond to compound 41:

\[ 25 + \text{NO}_2^+ \text{BF}_4^- \rightarrow \text{F}_5\text{S}\text{C}--\text{NO}_2\text{O} \quad \text{41} + \text{Cs}^+\text{BF}_4^- \]

Prolonged reaction resulted in an almost clean product that lacked a COF group and had an inverted AB$_4$ resonance, i.e., $\phi_B$ is downfield of $\phi_A$. However, further characterization was not possible, as the material apparently exploded when injected into a gas chromatograph.

A further interesting result was obtained with p-$\text{O}_2\text{N}\phi\text{N}_2^+$ BF$_4^-$, which did not react with 25. All that was observed was the very slow conversion of 25 into SF$_5$CH$_2$SO$_2$F (within weeks) when the reaction was run in CH$_3$CN:

\[ 25 + \text{p-}\text{O}_2\text{N}\phi\text{N}_2^+ \text{BF}_4^- \rightarrow \text{SF}_5\text{CH}_2\text{SO}_2\text{F} \]

That SF$_5$CH$_2$SO$_2$F was formed can be attributed to the presence of trace quantities of water, which would, in a catalytic cycle, lead to hydrolysis of 21, accompanied by HF evolution, which in turn would react with the glass container, again forming H$_2$O.
SYNTHESIS AND PROPERTIES OF $F_4S=C\text{CF}_2\cdots 0$SO$_2$

When 1 was treated with 18, a two-directional elimination was observed:

\[
F_5S\text{CF}_2\text{SO}_2 \quad 1 + F_3B\cdot N(C_2H_5)_3 \quad \rightarrow \quad F_4S=C\text{CF}_2\text{SO}_2 \quad 23 \ (\approx 63\%) \\
F_5S=C=C=0 \quad 21 \ (\approx 32\%)
\]

Two facts are remarkable about this reaction:

1: The formation of an unsaturated sultone
2: The formation of ketene 21

As with the formation of 21 from 18 and 22, the weak basicity of 18 allowed this reaction to be carried out. Ketene 21, as was said above, adds even comparatively weak bases very readily to form ionic adducts. Other bases, like triethylamine (which reacts vigorously with the ketene) or NaF (which forms an adduct with it) would not have allowed for this reaction to take place.

The ratio 21:23 remains nearly constant under a variety of conditions. When sultone 1 is heated with various excesses of base 18 (0.1-0.4-fold), heated longer, reacted by slow addition of hot base to its melt, or reacted in chloroform, the reaction produces a mixture of 30-35% 21 and 65-70% of 23 (19F n.m.r.).

While 23 could be expected as a normal, although unprecedented, elimination product of 1, the formation of 21 was, in light of the formation of 23, surprising. As with 2, where the analogous reaction had led to 3, it was expected
to obtain only \( \text{21} \) from this reaction. It was suggested that the conversion \( \text{2} \rightarrow \text{3} \) led through the rearranged form of \( \text{2} \), the acyl fluoride \( \text{4} \). Although no evidence for this was put forward, it must have been concluded from the fact that 1. the reaction of \( \text{2} \) and \( \text{18} \) was much more vigorous than the reaction of \( \text{4} \) and \( \text{18} \) and 2. that only a keten product was observed, but no \( \text{F}_2\text{C}=\text{CCF}_2\text{OSO}_2 \), the analogue to \( \text{23} \). There is no a priori reason as to why an exomethylene group at oxathiacyclobutane should not be capable of existing. In the present case a partial re-arrangement of \( \text{1} \) to \( \text{22} \) cannot be excluded, but it is also possible that a two-directional elimination of HF from \( \text{1} \) took place, leading to a ring-unsaturated sultone \( \text{41} \) and \( \text{23} \):

![Chemical Diagram]

Compound \( \text{41} \) would not be stable towards rearrangement to \( \text{21} \), as only the latter is observed. A similar course of reaction has been suggested in the conversion of \((\text{F}_3\text{C})_3\text{CC}=\text{CF}\) to \((\text{CF}_3)_3\text{CC(SO}_2\text{F})=\text{C}=\text{O}\), when reacted with \( \text{SO}_3 \) (page 13).

The first member of the alkylidene sulfur tetrafluorides, \( \text{CH}_2=\text{SF}_4 \), was obtained by Kleemann and Seppelt.\(^{57a-d}\) This compound undergoes facile addition
reactions with the polar agents HF, HCl, HBr, ICl, HgF₂, ASF₅, and with chlorine.⁵⁸ These additions are such that cis SF₄X systems are obtained, in contrast to the trans-SF₄X compounds from the fluorination of fluoroalkyl sulfenylchlorides.⁵⁹ Additional members of this series include FC(O)CH=SF₄⁶⁰, H₃CCH=SF₄⁶¹, F₃CCH=SF₄, and F₃C(CF₃)=SF₄.⁶²,⁶³

Separation of 2₁ and 2₃ is difficult, as their boiling points lie close together. A small quantity of pure 2₃ may be collected at 92°C and of pure 2₁ at 94°C, however, most of the material decomposes during the distillation. It was possible to utilize the high affinity of 2₁ towards bases to effect selective compound formation, leaving pure 2₃ behind. Treatment of the mixture of 2₁ and 2₃ with CsF removes 2₁, which forms the adduct 2₅.

\[ 2₁ + 2₃ + \text{CsF} \rightarrow 2₅ + 2₃ \]

It is necessary to keep the temperature close to room temperature, as otherwise a reaction of 2₃ with CsF ensues and also to use very finely ground and thoroughly ground CsF in order to effect reaction. One finds, however, always that more Cs⁺ F₅SC(SO₂F)COF⁻ is formed than could arise from the amount of 2₁ in the mixture. Formation of 2₅ from 2₃ is confirmed in a separate experiment where pure 2₃ is treated with CsF:

\[
\begin{array}{c}
\text{F₄S=O} \quad \text{Cs⁺} \\
\text{F²S⁻O} \quad -\rightarrow \quad \text{F₅SC=O⁻} \quad -\rightarrow \quad \text{F₂S⁻O} \quad \text{Cs⁺} \\
\end{array}
\]
Compound 23 is a stable colorless liquid with an unpleasant odor; it can be handled in air for short times without decomposition.

The addition of other polar species besides $F^-$, and in analogy to $CH_2=SF_4$, was expected, with the negatively polarized end attacking the sulfur. Hydrogen fluoride adds to 23, giving sultone 1 in high yields:

$$F_4S=\text{[cyclic structure]} + HF \rightarrow F_5S-\text{[cyclic structure]}_1$$

It is much more difficult to effect addition of $HCl$ to 23. At room temperature no reaction occurs, only after heating with an excess of gaseous $HCl$ for nearly 100 hours ($68^\circ C$) is reaction complete without significant by-product formation:

$$F_4S=\text{[cyclic structure]} + HCl \rightarrow F_4S-\text{[cyclic structure]}_2$$

This contrasts very strongly with the ease of $HCl$ addition to $CH_2=SF_4$. One reason for this could be that in 23 there is a better charge-dispersal as compared to CH$_2$-SF$_4$:

$$F_4S=\text{[cyclic structure]} \leftrightarrow F_4S-\text{[cyclic structure]} \leftrightarrow F_4S-\text{[cyclic structure]} \leftrightarrow F_4S=\text{[cyclic structure]} \text{ etc}$$

and

$$F_4S=CH_2 \leftrightarrow F_4S^+-CH_2^-$$

which would lower the polarity of the S-C bond in 23. Another possibility is increased steric hindrance due to the comparatively large sultone ring. Compound 42 bears some
interesting possibilities. In analogy to trans R-SF$_4$-Cl, which reacts with olefins and acetylenes to adducts, there was the hope to see a similar reaction. Upon attempted reaction with CF$_2$=CF$_2$, only decomposition was seen. A small amount of unreacted 42 was detected in the n.m.r. spectrum of the reaction mixture:

$$42 + C_2F_4 \text{ (hv)} \rightarrow 4SC_2F_4Cl$$

Similarly, the re-arrangement of 42 with sodium fluoride, which is normally an unproblematic reaction, led to a complicated mixture, and it cannot be said whether the reaction took place at all:

$$42 + NaF \rightarrow \text{complicated mixture}$$

Methanol led to the decomposition of the SF$_4$Cl group (in an n.m.r. experiment); thus, at this time, corresponding esters F$_4$SClCH(SO$_2$F)COOR have not been obtained from 42.

42 is a colorless liquid with a repugnant odor. Its $^{19}$F n.m.r. spectrum will be discussed in the n.m.r. chapter. The preparation of 42 is quite lengthy, so that only small amounts of it were available; consequently, its reactivity was not extensively investigated.

A further reaction of 43 is the specific addition of methanol, to yield another cis-SF$_4$-R compound as a white low-melting solid:

$$43 + CH_3OH \rightarrow F_4S-C-CF_2$$
The reaction of 23 and isopropanol led to a mixture of what was probably the cis-SF$_4$(OCH(CH$_3$)$_2$) derivative of the sultone and the ester F$_5$SCH(SO$_2$F)COOCH(CH$_3$)$_2$ 21a. Attempts to isolate the SF$_4$-derivative failed, as the compound was apparently not stable. The result with (CH$_3$)$_3$COH was even more pronounced, as no more cis-SF$_4$OR compound could be detected in the $^{19}$F n.m.r. spectrum. What was found was largely the ester F$_5$SCH(SO$_2$F)COOC(CH$_3$)$_3$ 21d, and $^{19}$F n.m.r. results suggest also initial formation of 1, and later F$_5$SCH$_2$SO$_2$F. In order to explain this result, one has to consider the two possible sites of attack at 23:

\[ \begin{align*}
23 & \quad \xrightarrow{\text{k}_2} \quad 21c \\
\text{HF} + 23 & \quad \rightarrow \quad 1
\end{align*} \]

It was described earlier, that an intermediate 21c should have been present leading to the compound (F$_2$S=O)=C(SO$_2$F)COF (44) by a re-arrangement in a F$_3$B·N(C$_2$H$_5$)$_3$ + BF$_4^-$ + NH(C$_2$H$_5$)$_3^+$ medium; HF would be first absorbed by 18, and no HF addition to the double-bond is expected in this case. In the present case, no 44 could be detected. This is not surprising, as neither the F$_2$S=O, nor the S=C, nor the COF moieties are expected to withstand an alcoholic environment. This re-arrangement to 44 is also expected to compete with
the addition of HF to the S=C double-bond, and HF addition appears to be quite faster than the rearrangement, as concluded from the product-distribution (mainly the SF$_5$-ester, but also 1), $k_1 > k_2$.

The relationship of $k_1$ and $k_3$ seems to be dictated by the alcohol used. The most obvious difference between methanol and t-butanol is the steric bulk, and the fact that with isopropanol an intermediate result is obtained allows one to draw the conclusion that steric reasons are responsible for the different course of the reactions of 23 and various aliphatic alcohols.

The preferred attack of CH$_3$OH at the doubly bonded sulfur shows that the S=C bond is more reactive than the sultone ring, but that this increased reactivity is easily offset by steric demands. It is here also pertinent that 23 failed to react with either C$_6$F$_5$OH, CF$_3$CH$_2$OH, (CF$_3$)$_2$CHOH or (CF$_3$)$_2$(C$_2$F$_5$)COH. In all cases a broad new feature was observed at $\approx$ 73 ppm, which is also observed as a decomposition product of pure 23, accompanied by the slow disappearance of 23.

Furthermore, 23 did not react in the expected fashion with either CF$_3$OCl or CF$_3$O$^-$Cs$^+$. Familiar products were obtained in these cases:

$$23 + \text{CF}_3\text{OCl} \rightarrow \text{F}_6\text{S}C(\text{Cl})\text{CF}_2 \text{O}_2\text{S}$$

$$23 + \text{CF}_3\text{O}^- \text{Cs}^+ \rightarrow 25 + (\text{CF}_2\text{O})$$

The addition of ClF instead of CF$_3$OCl in reactions of CF$_3$OCl
is not uncommon. The addition of CF₃OC₁ is now considered a polar addition, rather than a free radical one. This conclusion has been reached from the product distribution in several reactions.⁶⁴

Compound 39 can be obtained from 2₃ and ClF, although not without difficulty: an excess of ClF and several days of reaction time were necessary in order to have complete reaction.

\[
\begin{array}{c}
\text{Cl} \\
F₅S-C-CF₂ \\
O₂S-O \quad 39
\end{array}
\]

It could not be obtained by sulfur trioxidation of SF₅CCl=CF₂ (p.42). Complete degradation of the SF₅ group was observed in this case. This process demanded apparently more than one equivalent of SO₃, and the product that was obtained in this fashion could not be analyzed.

39 is a reactive compound, as is evident from the formation of F₅SCCl(SO₂F)COF (2₅a) as a major by-product in its formation from 2₃ and ClF; only when acid-washed apparatus was used throughout, formation of 2₅a could be mostly avoided. The conversion of 39 into 2₅a occurred on the potassium bromide plates (apparently catalyzed by Br⁻) that were used for i.r. analysis: the initially absent COF band increased slowly. It is much easier to obtain 2₅a from the cesium salt 2₅ than from 39, so that the latter route is of no synthetic value. The conversion of 39 into 2₅a is part of a general reaction of fluorosultones of the form RₓCXCF₂OSO₂
(R_f = fluoro radical, X = H or halogen) with bases that results in the opening of the four-membered ring to give an acyl fluoride.

23 seems to be completely unreactive towards SF_5Br.

THE CONVERSION OF AN SF_5 GROUP INTO AN (F_2SO)= GROUP

A peculiar reaction occurs when esters 21a or 21b are treated with 18. In a vigorous reaction, the complete degradation of the SF_5 group is observed. Instead of obtaining the expected F_4S=C derivative it is found, that, in the case of the isopropylester, a mixture of gaseous products and a high-boiling liquid is produced. The gas product was tentatively identified as a mixture of propylene and 2-fluoropropane, whereas the spectral and analytical data of the high-boiling liquid are only consistent with a composition of the form:

\[
\begin{align*}
\text{F}_2\text{S}=\text{O} & \quad \text{O} \\
\text{C}-\text{C} & \quad \text{F}.
\end{align*}
\]

It seemed likely, that isopropyl fluoride was formed first as an elimination product of the ester and underwent then HF loss by reaction with the strong HF-acceptor F_3B·N(C_2H_5)_3 to produce propylene as the secondary product.

In order to facilitate the analysis of the gaseous products, and to support the surmise of primary alkyl fluoride formation, the ester F_5SCH(SO_2F)COOCH_3 was synthesized from F_5SCHCF_2OSO_2 (1) and CH_3OH. Treatment of
the methylester with $F_3B\cdot N(C_2H_5)_3$, (18), should give methyl fluoride as the only volatile product, since HF elimination would not lead to a stable product. When the ester was treated with $F_3B\cdot N(C_2H_5)_3$, $CH_3F$ was produced in a high yield and as the sole gaseous product:

$$F_5SCH(SO_2F)COOCH_3 + F_3B\cdot N(C_2H_5)_3 \rightarrow (F_2SO)=CCOF + CH_3F$$

The new trifunctional compound 44 is the first of its kind. There is only one other example that contains the COF and $F_2SO$ functional groups bonded to carbon. This compound, $(F_2SO)=CHCOF$, was obtained by hydrolysis of $F_4S=CHCOF$ and seems to exist in a number of rotation isomers, as shown by variable temperature n.m.r. spectroscopy.

The first order $^{19}F$ n.m.r. spectrum of 44 at room temperature is very similar to that of $(F_2SO)=CHCOF^2$. Chemical shifts and splitting patterns are comparable, with the exception that one has the additional large coupling to the $SO_2F$ fluorine. The coupling constants are similar to the ones found in $SF_5CX(SO_2F)COF$, where $X = H$ (11), $F^6$, $Cl$ (25a), Br (25b).

The main characteristics of the infrared spectrum include the strong C=O stretching vibration (1825 cm$^{-1}$), various $S=O$ stretching vibrations, i.e., $v_{assSO_2} = 1448$ cm$^{-1}$, $v_{SO} = 1419$ cm$^{-1}$. The asymmetric $SO_2$ stretching mode shows only a small variability and appears always between $\approx$1440 and 1460 cm$^{-1}$. The symmetric $SO_2$ stretching mode frequency occurs normally at $\approx$1220 cm$^{-1}$; in the same region (1100-1300
one expects the resonance for the S=C (double) bond. Since there are several bands in this region, an unambiguous assignment is not possible, but the band the band at 1226 cm$^{-1}$ could correspond to $\nu_{\text{sym} S=O}$. In particular the SF (stretching) modes at 875 and 801 cm$^{-1}$ have a rather unusual appearance, as compared to SF$_5$ band shapes. The two bands are of about equal intensity and show no further structure, whereas in SF$_5$-SO$_2$F compounds one finds a multitude of bands of various intensity in the same region.

The mass spectrum shows the molecular ion and appropriate fragment ions, but also one rather intense band which can be accounted for if one assumes rearrangement to

$$F_3OS\quad C=C=O.$$  \hspace{1.5cm} \text{(44a)}

The formation of (F$_2$SO)=C(SO$_2$F)COF and alkyl fluoride can be rationalized by a series of eliminations and rearrangements. In the first step, formation of the HF elimination product is conjectured:

$$F_5S\quad C-H \quad C=O \quad + \quad F_3B\cdot N(C_2\text{H}_5)_3 \quad \rightarrow \quad F_4S\quad C=\quad C=O.$$  \hspace{1.5cm} \text{(21c)}

R = CH(CH$_3$)$_2$ (21a), CH$_3$ (21b)

It has to be assumed that the newly-formed F$_4$S= ester is unstable towards rearrangement. The ester could undergo ring closure, lose RF and undergo transformation by a [1,3]-sigmatropic shift to 44:
Alternatively, attack of the alkoxy group at SF₄ is possible, followed by elimination, ring opening and again [1,3]-sigmatropic shift. Although a number of other mechanisms can be formulated, which involve migration of F⁻, a concerted mechanism seems more likely for the following reasons:

1. F⁻ migration is deemed unlikely in the reaction medium. BF₄⁻ is not a good source of F⁻.
2. In the mass spectrum there is a moderately strong m/e = 105 peak. This could indicate the fragment SOF₃⁺, formed from 44 in the reversal of the last step in the scheme formulated above in the mass spectrometer. For the mass 105 the composition (MH-SOF₄)⁺ is also formulated.
3. Ketene 21 was stable in the presence of BF₄⁻ NH(C₂H₅)₃⁺ and 44a is not expected to be very different.

If the mechanism were ionic, then F⁻ addition to 44a would be a likely step; this would mean that 44a could compete successfully with BF₃ for F⁻. An intermediate (F₃SO⁻)=C(SO₂F)COF would be formed which would give 44 upon loss of F⁻. 1,3 fluorine migrations can apparently take place thermally as found for (CF₃)₂SO₂ (pyrolysis of (CF₃)₂(SO₂)₂(CF₃)₂, leading to F₃CC(SO₂F)=CF₂)₆⁷a, slow conversion at room temperature of (CF₃)₂C=P(O₂(CH₂)₂)OCH₃ to F₂C=C(CF₃)PF(O₂(CH₂)₂)OCH₃)₆⁷b, pyrolysis of (CH₃)₃SnP(CF₃)CH(CF₃)₂, leading to (CF₃)FP=C(CF₃)=CF₂)₆⁷c.
and the spontaneous conversion of \((\text{CF}_3)_2\text{C} = \text{C}=\text{O}\) to \(\text{F}_2\text{C}(\text{CF}_3)\text{CF}0\)\(^{67d}\). Light induced 1,3 F migration was found for tetratrifluoromethyl ethylene\(^e\)-> \(\text{F}_2\text{C} = \text{C}(\text{CF}_3)\text{CF}(\text{CF}_3)_2\).

Further reactions of the trifunctional molecule could not be completed successfully. Reaction with diethylamine led to an inseparable mixture of many compounds, as evidenced by the \(^{19}\text{F}\) n.m.r. spectrum and reaction with CsF in CH\(_3\)CN led to the uptake of one equivalent of CsF, but the fluorine spectrum showed the presence of at least two COF groups. Separation was not attempted.

While the esters 21a and 21b gave 44 when treated with \(\text{F}_3\text{B} \cdot \text{N}(\text{C}_2\text{H}_5)_3\) similar treatment of the ester \(\text{SF}_5\text{CH}_2\text{COOCH}_3\) resulted in decomposition; only residual ester and \(\text{SF}_4\) was observed in the \(^{19}\text{F}\) n.m.r. spectrum of the reaction mixture.

**NUCLEAR MAGNETIC RESONANCE SPECTRA**

Most of the compounds that were synthesized and described have no unusual n.m.r. spectra. Covalent \(\text{SF}_5\) compounds appear in a certain chemical shift range, where \(\phi_A\) ranges from \(\approx 60-85\) ppm and \(\phi_B\) from \(\approx 45\) to 75 ppm (CCl\(_3\)F = 0.0 ppm), and \(J_{AB}\) is usually 150 \(\pm\) 7 Hz. The octahedral environment of the sulfur leads to two different kinds of fluorine, the 4 equatorial fluorines (B) and one axial fluorine (A), and the spin system is classified as \(\text{AB}_4\). Furthermore, when the chemical shift difference \(\phi_A - \phi_B\) becomes small (\(\approx < 3\)ppm), the usual \(\text{AB}_4\) spectrum, which
consists of a doublet of a multiplet (the B-resonance) and a 9-line pattern, which is actually the superimposition of a singlet, a triplet and a quintet, becomes degenerate and appears as a multiplet. It cannot be interpreted by manual means anymore. As long as an $AB_4$ spectrum is seen, however, such a hand-analysis can be made with reasonable accuracy. The values that are obtained are usually acceptable, although in most cases not too accurate, as an exact determination of $\phi_B$ and $J_{AB}$ rests upon an accurate assignement of lines within the B-resonance. This is not usually possible, as either the lines are not well enough resolved or it is not simple to determine the correct lines within the irregular multiplet. A correct assignment is realistically only possible with a spectral simulation program. Relationships between the individual resonance lines and the sought-after quantities of $\phi_A-\phi_B$ and $J_{AB}$ were published by Borden et al.\textsuperscript{68} and by Harris and Packer.\textsuperscript{69} Packer's and Harris's procedure was used here. Only $\phi_A$ can be measured accurately as the center line (the singlet) of the 9-line pattern which represents the resonance of the axial fluorine. Most of the $^{19}$F n.m.r. spectra were normal $AB_4$ spectra, with $\phi_A$ downfield of $\phi_B$ (e.g. 25), then there were intermediate cases (small $\phi_A-\phi_B$, e.g. 25a), the degenerate case (e.g. 21a, 21b, 30), and again the $AB_4$ case, this time with $\phi_A$ upfield of $\phi_B$. Fig.1 exemplifies the change an $AB_4$ spectrum undergoes when $\phi_A-\phi_B$ is successively
altered. In the end, one has the mirror image of the original spectrum if one goes from $\phi_{A}-\phi_{B} = x$ ppm to $\phi_{A}-\phi_{B} = -x$ ppm. The $\phi_{A}-\phi_{B} = 0$ spectrum is a singlet, although the coupling constant is 150 Hz. This is a similar case to e.g. methane or benzene, where coupling cannot be directly observed.
Figure 1. The change of an AB$_4$ spectrum (84.7 MHz) for various $\phi_A - \phi_B$. $J_{AB} = 150$ Hz. 
a.) $\phi_A - \phi_B = 30$ ppm. b.) $\phi_A - \phi_B = 10$ ppm. c.) $\phi_A - \phi_B = 3$
ppm. d.) $\phi_A - \phi_B = 0$ ppm. e.) $\phi_A - \phi_B = -30$ ppm.
In the $^{19}$F n.m.r. spectra of carbanionic species (25, 32a, 32b, 33, 34, 45) it is observed that both the $\text{SO}_2\text{F}$ and the $\text{SF}_5$ resonances appear, contrary to the expectation that increased charge density lead to increased shielding and thus an upfield shift, at a downfield position as compared to neutral compounds. This shift showed only a minor dependence on the cation ($\text{Cs}^+$, $\text{Ag}^+$, $\text{Na}^+$, $\text{HN(C}_2\text{H}_5)_3^+$, $\text{Hg}^2+$). A related observation was made with perfluoroalkyl carbanions, e.g. Dyatkin et al notice this effect for $(\text{CF}_3)_3\text{C}^-\text{Cs}^+$ and attribute it to the "reverse redistribution of electron density" via fluorine atoms to the vacant inner orbitals of the metal atoms. A complexation of the anion with the metal cation is assumed. More recently, the same property was discussed by Bayliff and Chambers who observed this downfield shift for $\text{Cs}^+$ and $(\text{CH}_3)_2\text{N})_3\text{S}^+$ cations and it was noticed that the theory of magnetic shielding was in "some disarray". Apparently, there is no adequate theory that can explain satisfactorily this phenomenon in an anion, as it is also observed with anions that do not have vacant inner orbitals. A related observation was made in neutral compounds, where the fluorine resonance was shifted to a deshielded position by an electron donating substituent. This downfield shift is also observed in a number of compounds that have to be considered as salts or zwitterions (e.g 26). A common feature of the anions that were characterized here is the
presence of an SO$_2$F, SF$_5$ andCOR at a carbon: in all cases they are enolates.

The CFO group in the compounds F$_5$SC(SO$_2$F)=CFO$^-$ [Cs,Ag,Na]$^+$ is in a strongly shielded position (17ppm), as compared to neutral CFO-compounds. This suggests that the charge is concentrated mainly at the CFO-group; the lowering of the C=O stretch frequency from F$_5$SCH(SO$_2$F)CFO is about 90 cm$^{-1}$ for 25 (1766 cm$^{-1}$), underscoring the latter conclusion. Similar low-field resonances were reported earlier for the anions FO$_2$SNCFO$^-$ (Cs,K,Na)$^+$ and F$_5$SC(CF$_3$)$_2^-$ (Cs,Rb)$^+$ although they were not discussed.

In the mercurial (F$_5$SC(SO$_2$F)COF)$_2$Hg the COF resonance appears at a somewhat less shielded position (21.6ppm), as compared to the sodium, silver and cesium salts. The SF$_5$ and SO$_2$F resonance in the mercury compound is in a less deshielded position than in these salts. This observation parallels with the usual property of mercurials as more covalent compounds (they are often soluble in nonpolar solvents and can be distilled), which probably holds here, too. The C=O resonance of the mercurial appears at 1826 cm$^{-1}$, which argues for a more covalent compound, too.

In the $^{13}$C spectra of perfluoroalkyl carbanions$^{68}$ $^6$Rf$^-$ increased shielding is seen at the anionic center, as expected from theory, but deshielding is observed at the carbon atoms next to the anionic center. A similar effect was observed for 25 and 22,
although both carbons showed a downfield shift as compared to 22. While in the perfluorocarbanions the charge is localized (disregarding hyperconjugation), it can be distributed over the enol system in 25. This can account for the simultaneous deshielding, if associated with increased charge density, in 25. The observation of a low field resonance for an SF₅-compound can thus be rather generally associated with an anionic site next to SF₅; one has to be cautious, though, because some SF₅-hydrocarbons (having the grouping F₅SCH₂) show φₐ ≈ 75 ppm, and φₐ ≈ 85 ppm, which is very close to the range that was observed for the anions.

The ¹⁹F n.m.r. spectra of compounds 23, 42 and 43 are complicated. They are classified as AA'BCXX' (23) and as ABCDXY (42, 43) spin systems. For 42 and 43 there are also chiral centers which complicated the analysis even more. The ¹⁹F n.m.r. spectrum of 23 is similar to the spectra of other SF₄= compounds. The spectral parameters were obtained by several steps. 1.: An AA'BC spectrum consists of two (approximately) independent subspectra. This allowed to determine φₐ, φₐ and Jₐ, reducing the number of unknown parameters to 3 (φₐ, Jₐ, Jₐ). 2.: φₐ was guessed from the appearance of the spectrum, and the remaining couplings were assumed to fall into a certain range (145-165 Hz). This latter assumption was warranted by comparison with similar
compounds. Furthermore, each line in the SF₄ spectrum showed further triplet splitting of different magnitude in the A, B and C part of the spectrum. This represented the coupling to the CF₂ group, the chemical shift of which was also measurable and which appears roughly as a quintet. A complete set of spectral parameters could then be calculated with the aid of an iterative n.m.r. simulation program. From these values, and in analogy to the structure of CH₂=SF₄ (X-ray structural determination), it was now possible to assign the following structure to 23:

In this structure, the axial fluorines F_B and F_C (the SF₄S= group is a fragment of the SF₆ octahedron) lie in the plane of the sultone ring. The alternative structure where F_A and F_A' lay in the plane of the sultone ring would give the same n.m.r. spectrum. That 23 has an AA'BC spectrum shows also that the fluorine atoms do not undergo exchange, as was found for similar F₄S=N systems.
Figure 2. Experimental and calculated $^{19}$F n.m.r. spectrum of $\text{F}_4\text{S}=\text{CCF}_2\text{OSO}_2$, 23.

The $^{13}$C n.m.r. spectrum of 23 (table 2) reveals two resonance groups, one at 114.5 ppm and a weaker one at 80.7 ppm. From the intensities and the splitting patterns the resonance at 80.7 ppm is assigned to $\text{C}^\alpha$, as the absence of an attached spin-one half nucleus decreases relaxation, leading to partial saturation. The $\text{C}^\alpha$ resonance appears approximately as a quintet of triplets; it is assumed that the four fluorines have coupling constants of comparable magnitude ($\approx 32$ Hz) to $\text{C}^\alpha$ which would account for the quintet splitting, while the triplet coupling (7.8 Hz) would arise from coupling to the fluorines at $\text{C}^\beta$. The other resonance signal ($\text{C}^\beta$) is a triplet of doublets. The triplet coupling
(282 Hz) is nearly the same as in 1-(293 Hz) and is attributed to the C\textsuperscript{β}-F couplings, and the smaller doublet coupling (23.2 Hz) is assumed to arise from trans-C\textsuperscript{β}-F\textsubscript{B} coupling. This latter coupling was unchanged on raising the sample temperature to 40°C, implying that not multiple ring conformations are resolved.

The \textsuperscript{19}F n.m.r. spectrum of 42 shows three types of fluorine in the ratio of 1:2:1, which identifies it clearly as a cis-SF\textsubscript{4}R compound, because for a trans arrangement a singlet is observed for the fluorine resonances, due to their equivalence, whereas the CF\textsubscript{2} group appears as an AB system with asymmetric coupling to the proton, analogous to sultone 1. The coupling within the SF\textsubscript{4} arrangement is complicated and only partial interpretation of the spectrum is possible. In 42 as well as in 43 one observes actually four fluorine resonances. Two of these lie close together, respectively, and are very similar, except that in one of them additional coupling is seen. This suggests that the rotation of the SF\textsubscript{4}X (X = Cl, OCH\textsubscript{3}) group is actually hindered, perhaps through hydrogen bonding between either a fluorine or X and the hydrogen at C\textsubscript{α} of the sultone ring. The assignment of the different fluorines and coupling constants followed the one given for similar compounds. The CF\textsubscript{2} group appears as an AB system.

The \textsuperscript{13}C n.m.r. spectrum of 42 with proton coupling shows a triplet and two multiplets, which collapse upon \textsuperscript{1H}-
decoupling into one at 109 ppm. The multiplet consists of nine lines, which show additional splitting and may be assigned to Cα (the same numbering is assumed as for 22). The remaining resonances, from Cβ, appear as four lines, the middle two of which overlap to produce a distorted triplet centered at 113.4 ppm.

**TABLE I**

19F N.M.R. AND I.R. TABLE OF SF5-ANIONS

<table>
<thead>
<tr>
<th>Structure</th>
<th>Counterion</th>
<th>φA</th>
<th>φB</th>
<th>φX</th>
<th>JAB</th>
<th>VCO</th>
</tr>
</thead>
<tbody>
<tr>
<td>F5SC(SO2F)COOC3H7</td>
<td>Na⁺</td>
<td>95.3</td>
<td>82.8</td>
<td>67.3</td>
<td>150.4</td>
<td>-</td>
</tr>
<tr>
<td>F5SC(SO2F)COOC3H7</td>
<td>2</td>
<td>93.5</td>
<td>81.7</td>
<td>66.2</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>F5SC(SO2F)COOC3H7</td>
<td>3</td>
<td>97.0</td>
<td>82.1</td>
<td>66.3</td>
<td>154.0</td>
<td>1615</td>
</tr>
<tr>
<td>F5SC(SO2F)COOC3H7</td>
<td>2</td>
<td>96.1</td>
<td>82.0</td>
<td>65.8</td>
<td>151.3</td>
<td>1589</td>
</tr>
<tr>
<td>F5SC(SO2F)COOC3H7</td>
<td>4</td>
<td>95.4</td>
<td>82.1</td>
<td>67.0</td>
<td>154.2</td>
<td>1640</td>
</tr>
<tr>
<td>F5SC(SO2F)COOC3H7</td>
<td>5</td>
<td>89.6</td>
<td>79.7</td>
<td>71.3</td>
<td>154.2</td>
<td>1669</td>
</tr>
<tr>
<td>F5SC(SO2F)COOC3H7</td>
<td>6</td>
<td>95.7</td>
<td>82.0</td>
<td>73.1</td>
<td>150.2</td>
<td>1695</td>
</tr>
<tr>
<td>F5SC(SO2F)COOC3H7</td>
<td>7</td>
<td>95.4</td>
<td>82.1</td>
<td>67.0</td>
<td>154.2</td>
<td>1669</td>
</tr>
<tr>
<td>F5SC(SO2F)COOC3H7</td>
<td>8</td>
<td>95.5</td>
<td>82.6</td>
<td>69.5</td>
<td>151.5</td>
<td>-</td>
</tr>
</tbody>
</table>

1 in isopropanol, 2 in ether, 3 in CD3COCD3, 4 in CDCl3, 5 in CH2Cl2, 6 in acetone, 7 in CH3CN; * neat samples [φ] = ppm; [J] = Hz; [v] = cm⁻¹.

**TABLE II**

13C N.M.R. TABLE OF SF5CαCβ COMPOUNDS

<table>
<thead>
<tr>
<th>Structure</th>
<th>δα</th>
<th>δβ</th>
</tr>
</thead>
<tbody>
<tr>
<td>F5SC(SO2F)COF</td>
<td>25²</td>
<td>104.33</td>
</tr>
<tr>
<td>F5SC(SO2F)COF</td>
<td>22²</td>
<td>88.5</td>
</tr>
<tr>
<td>F5SC(SO2F)COF</td>
<td>31²</td>
<td>86.28</td>
</tr>
<tr>
<td>F4SC1CHCF2OSO₂</td>
<td>42²</td>
<td>108.8</td>
</tr>
<tr>
<td>F5SC(SO2F)=CHO3Si(CH3)₃</td>
<td>39a²</td>
<td>126.95</td>
</tr>
<tr>
<td>F5SCF2CFOJ</td>
<td>2</td>
<td>119.21 (t-t)</td>
</tr>
<tr>
<td>F5SCF2CF2Br</td>
<td>2</td>
<td>108.15</td>
</tr>
</tbody>
</table>

1 in CD3CN, 2 in CDCl3, 3 neat sample referenced to external CDCl3 [δ] = ppm, Si(CH3)₄ = 0.0. J: JCαFα = 314.5 Hz; JCβFβ = 323.1 Hz; JCβFα = 41.0 Hz
The infrared spectra of 23, 42 and 43 and are in accord with the proposed structures, too. Bands for the asymmetric (1400-1450 cm\(^{-1}\)) and symmetric stretching mode (1200-1250 cm\(^{-1}\)) of the SO\(_2\) group are observed: 1415 and 1192-1235 cm\(^{-1}\) for 23, 1430 and 1214 cm\(^{-1}\) for 42 and 1424 and 1216 for 43. To the C=S bond corresponds a strong band at ≈1200 cm\(^{-1}\).\(^{83}\) Several bands in this region in the i.r. of could be assigned to the C=S bond. Additional typical bands correspond to the S-F stretching modes in the SF\(_4\) groups of 23, 42 and 43 at 800-900 cm\(^{-1}\).

**FEATURES OF X-RAY STRUCTURES\(^{84}\)**

It was a good fortune that several of the SF\(_5\) compounds were crystalline. This allowed to determine some structural features. In one instance, the roentgenographic determination of the structure allowed the identification of the compound altogether, as only very small amounts of material were at hand, and other techniques were not completely conclusive. This was 29, formed in a not-understood way from what is probably F\(_5\)SC(SO\(_2\)F)=COCH\(_2\)O (28). The amide F\(_5\)SCH(SO\(_2\)F)CON(C\(_2\)H\(_5\))\(_2\), 27, and the salt F\(_5\)SC(SO\(_2\)F)COOCH\(_3\)\(^-\) NH(C\(_2\)H\(_5\))\(_3\)\(^+\) 32b were stable, crystalline compounds, which allowed the determination of their structural parameters as solids. The obvious close relationship between these compounds validates comparisons respecting bond-angles and -lengths.
<table>
<thead>
<tr>
<th>Bond/Angle</th>
<th>Distance/Angle [pm/°]</th>
</tr>
</thead>
<tbody>
<tr>
<td>S(1)-F(1)</td>
<td>157.1 (3)</td>
</tr>
<tr>
<td>S(1)-F(3)</td>
<td>157.3 (3)</td>
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<tr>
<td>S(1)-F(4)</td>
<td>158.4 (2)</td>
</tr>
<tr>
<td>S(1)-F(5)</td>
<td>159.5 (3)</td>
</tr>
<tr>
<td>S(1)-C(1)</td>
<td>177.0 (4)</td>
</tr>
<tr>
<td>C(1)-C(2)</td>
<td>144.7 (4)</td>
</tr>
<tr>
<td>C(2)-O(1)</td>
<td>120.7 (5)</td>
</tr>
<tr>
<td>O(2)-C(3)</td>
<td>143.6 (4)</td>
</tr>
<tr>
<td>S(2)-O(3)</td>
<td>139.5 (3)</td>
</tr>
<tr>
<td>S(2)-F(6)</td>
<td>141.9 (4)</td>
</tr>
<tr>
<td>S(1)-F(1)-F(2)</td>
<td>89.6 (1)</td>
</tr>
<tr>
<td>S(1)-F(1)-F(3)</td>
<td>90.0 (1)</td>
</tr>
<tr>
<td>S(1)-F(1)-F(4)</td>
<td>172.8 (1)</td>
</tr>
<tr>
<td>S(1)-F(2)-F(3)</td>
<td>86.8 (1)</td>
</tr>
<tr>
<td>S(1)-F(2)-F(4)</td>
<td>86.1 (1)</td>
</tr>
<tr>
<td>S(1)-F(3)-F(4)</td>
<td>94.0 (1)</td>
</tr>
<tr>
<td>S(1)-F(3)-C(1)</td>
<td>93.1 (1)</td>
</tr>
<tr>
<td>S(1)-C(1)-C(2)</td>
<td>123.5 (3)</td>
</tr>
<tr>
<td>S(1)-C(1)-S(2)</td>
<td>115.4 (3)</td>
</tr>
<tr>
<td>C(1)-C(2)-O(1)</td>
<td>115.4 (3)</td>
</tr>
<tr>
<td>C(1)-C(2)-O(2)</td>
<td>116.2 (3)</td>
</tr>
<tr>
<td>C(1)-C(2)-C(3)</td>
<td>112.1 (2)</td>
</tr>
<tr>
<td>C(1)-S(2)-F(6)</td>
<td>104.6 (2)</td>
</tr>
<tr>
<td>C(1)-S(2)-F(6)</td>
<td>101.3 (2)</td>
</tr>
<tr>
<td>C(1)-N-C(12)</td>
<td>113.7 (3)</td>
</tr>
<tr>
<td>C(1)-C(12)</td>
<td>115.7 (3)</td>
</tr>
<tr>
<td>N-C(13)-C(14)</td>
<td>113.9 (4)</td>
</tr>
</tbody>
</table>
### TABLE IV

BOND DISTANCES [pm] AND ANGLES [°] IN $\text{F}_5\text{SCH(SO}_2\text{F)}\text{CON(C}_2\text{H}_5)_2$, 27, STANDARD DEVIATIONS IN PARENTHESES

<table>
<thead>
<tr>
<th>Bond</th>
<th>Distance [pm]</th>
<th>Angle [°]</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{S(1)}-\text{F(5)}$</td>
<td>153.2(4)</td>
<td>95.0</td>
</tr>
<tr>
<td>$\text{S(1)}-\text{F(3)}$</td>
<td>153.7(4)</td>
<td></td>
</tr>
<tr>
<td>$\text{S(1)}-\text{F(4)}$</td>
<td>155.1(4)</td>
<td></td>
</tr>
<tr>
<td>$\text{S(1)}-\text{F(1)}$</td>
<td>155.3(4)</td>
<td></td>
</tr>
<tr>
<td>$\text{S(1)}-\text{F(2)}$</td>
<td>156.8(3)</td>
<td></td>
</tr>
<tr>
<td>$\text{S(1)}-\text{C(1)}$</td>
<td>185.6(5)</td>
<td></td>
</tr>
<tr>
<td>$\text{S(2)}-\text{O(2)}$</td>
<td>139.6(4)</td>
<td></td>
</tr>
<tr>
<td>$\text{S(2)}-\text{O(1)}$</td>
<td>140.2(4)</td>
<td></td>
</tr>
<tr>
<td>$\text{S(2)}-\text{F(6)}$</td>
<td>152.2(4)</td>
<td></td>
</tr>
<tr>
<td>$\text{S(2)}-\text{C(1)}$</td>
<td>180.1(5)</td>
<td></td>
</tr>
<tr>
<td>$\text{O(3)}-\text{C(2)}$</td>
<td>121.3(5)</td>
<td></td>
</tr>
<tr>
<td>$\text{N(1)}-\text{C(2)}$</td>
<td>132.7(6)</td>
<td></td>
</tr>
<tr>
<td>$\text{N(1)}-\text{C(5)}$</td>
<td>147.1(6)</td>
<td></td>
</tr>
<tr>
<td>$\text{N(1)}-\text{C(3)}$</td>
<td>147.7(6)</td>
<td></td>
</tr>
<tr>
<td>$\text{F(5)}-\text{S(1)}-\text{F(3)}$</td>
<td>91.1(3)</td>
<td>107.25</td>
</tr>
<tr>
<td>$\text{F(5)}-\text{S(1)}-\text{F(4)}$</td>
<td>176.2(2)</td>
<td>104.3(3)</td>
</tr>
<tr>
<td>$\text{F(5)}-\text{S(1)}-\text{F(1)}$</td>
<td>90.2(3)</td>
<td>112.6(3)</td>
</tr>
<tr>
<td>$\text{F(5)}-\text{S(1)}-\text{F(2)}$</td>
<td>87.9(2)</td>
<td>117.7(3)</td>
</tr>
<tr>
<td>$\text{F(5)}-\text{S(1)}-\text{C(1)}$</td>
<td>92.2(2)</td>
<td>124.5(4)</td>
</tr>
<tr>
<td>$\text{F(3)}-\text{S(1)}-\text{F(4)}$</td>
<td>89.5(3)</td>
<td>117.4(4)</td>
</tr>
<tr>
<td>$\text{F(3)}-\text{S(1)}-\text{F(1)}$</td>
<td>176.3(2)</td>
<td>118.0(4)</td>
</tr>
<tr>
<td>$\text{F(3)}-\text{S(1)}-\text{C(1)}$</td>
<td>88.9(2)</td>
<td>109.46</td>
</tr>
<tr>
<td>$\text{F(3)}-\text{S(1)}-\text{C(2)}$</td>
<td>93.0(2)</td>
<td>108.54</td>
</tr>
<tr>
<td>$\text{F(4)}-\text{S(1)}-\text{F(1)}$</td>
<td>89.0(3)</td>
<td>108.53</td>
</tr>
<tr>
<td>$\text{F(4)}-\text{S(1)}-\text{F(2)}$</td>
<td>88.4(2)</td>
<td>108.53</td>
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<td>$\text{F(4)}-\text{S(1)}-\text{C(1)}$</td>
<td>91.5(2)</td>
<td>108.53</td>
</tr>
<tr>
<td>$\text{F(1)}-\text{S(1)}-\text{F(2)}$</td>
<td>87.7(2)</td>
<td>113.2(4)</td>
</tr>
<tr>
<td>$\text{F(1)}-\text{S(1)}-\text{C(1)}$</td>
<td>90.4(2)</td>
<td>109.47</td>
</tr>
<tr>
<td>$\text{O(2)}-\text{S(2)}-\text{O(1)}$</td>
<td>120.8(3)</td>
<td>109.46</td>
</tr>
<tr>
<td>$\text{O(2)}-\text{S(2)}-\text{F(6)}$</td>
<td>106.9(2)</td>
<td>109.46</td>
</tr>
<tr>
<td>$\text{O(2)}-\text{S(2)}-\text{C(1)}$</td>
<td>114.5(2)</td>
<td>109.48</td>
</tr>
<tr>
<td>$\text{O(1)}-\text{S(2)}-\text{F(6)}$</td>
<td>107.4(3)</td>
<td>109.48</td>
</tr>
<tr>
<td>$\text{O(1)}-\text{S(2)}-\text{C(1)}$</td>
<td>104.2(2)</td>
<td>109.47</td>
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<td>$\text{C(2)}-\text{N(1)}-\text{C(5)}$</td>
<td>116.7(4)</td>
<td>108.93</td>
</tr>
<tr>
<td>$\text{C(2)}-\text{N(1)}-\text{C(3)}$</td>
<td>127.8(4)</td>
<td>108.93</td>
</tr>
<tr>
<td>$\text{C(5)}-\text{N(1)}-\text{C(3)}$</td>
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<td>108.95</td>
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<td>$\text{H(1)}-\text{C(1)}-\text{C(2)}$</td>
<td>107.25</td>
<td>111.6(5)</td>
</tr>
<tr>
<td>$\text{H(1)}-\text{C(1)}-\text{S(2)}$</td>
<td>107.26</td>
<td>109.51</td>
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<td>$\text{H(10)}-\text{C(6)}-\text{H(11)}$</td>
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<td>$\text{H(9)}-\text{C(6)}-\text{H(11)}$</td>
<td>109.46</td>
<td>109.47</td>
</tr>
<tr>
<td>$\text{H(11)}-\text{C(6)}-\text{C(5)}$</td>
<td>109.43</td>
<td></td>
</tr>
</tbody>
</table>
The crystal structure of the cation, triethylammonium, in 32b has been determined by James for the three anions Cl\textsuperscript{-}, Br\textsuperscript{-} and J\textsuperscript{-}. It was found, that the nitrogen-carbon and the carbon-carbon distances were subject to some change in going from Cl\textsuperscript{-} to J\textsuperscript{-}. The closest resemblance of the cation of 32b is found by comparison to HN(C\textsubscript{2}H\textsubscript{5})\textsuperscript{+} Br\textsuperscript{-}, and from this a significant compression of the ion is concluded. The anion shows some interesting bond-length and -angle changes, as compared to neutral compounds. A comparison will be made between the structures of 32b and 27.

In the anion large angles S\textsubscript{1}-C\textsubscript{1}-S\textsubscript{2}, S\textsubscript{1}-C\textsubscript{1}-C\textsubscript{2} and S\textsubscript{2}-C\textsubscript{1}-C\textsubscript{2} are found. They add up to 359.1\textdegree{} (120.2+123.5+115.4), while the same angle sum in the amide comes to 334.6\textdegree{} (117.7+112.6+104.3). The variation between a tetrahedron and a plane is from 328.4\textdegree{} to 360\textdegree{}. This shows that the anion is almost flat, while the amide shows strong pyramidalization. Usually, anions are pyramidal; this behavior is certainly partially caused by the great steric repulsion of the three large substituents. Because a lone pair can normally be considered as an extra substituent, which would favor a non-planar arrangement, another cause for this near-planarity must be the mesomeric interaction between the anion center at C\textsubscript{1}, the keto group and the SO\textsubscript{2}F
group. This charge dispersal would weaken the effect of the negative charge as a substituent. If one wants to consider hyperconjugation also, then the SF\textsubscript{5} group and additional conjugation in the SO\textsubscript{2}F group has to be included in this scheme, too.

The crystal structures of the anions SF\textsubscript{5}\textsuperscript{-86} and SF\textsubscript{5}O\textsuperscript{-87} were determined recently. While in SF\textsubscript{5}\textsuperscript{-} a significant elongation of the equatorial S-F bonds was found, with d(S-F\textsubscript{E}) > d(S-F\textsubscript{A}), the reverse was seen in SF\textsubscript{5}O\textsuperscript{-}. In 32b a similar effect is seen, with d(S-F\textsubscript{A}) = 159.5 pm and d(S-F\textsubscript{E}) = 157.8 ± 0.6 pm. Corresponding values in 27 are d(S-F\textsubscript{A}) = 156.8 pm and d(S-F\textsubscript{E}) = 154.3 ± 1.0. These values are very similar and do not indicate an important structural effect of the negative charge on the SF\textsubscript{5} moiety. On the other hand the SF\textsubscript{5}-C bond is significantly much shorter in 32b than in 27 (177.0 vs. 185.6 pm). The second C-S bond in 32b is also much shorter than in 27 (167.9 vs. 180.1 pm). Such a bond shrinkage is usually observed when a substituent is attached to an olefinic radical, but this could also indicate conjugation and hyperconjugation of the enolate moiety with both sulfur fragments:

\[
\begin{align*}
\text{F-S=O-CO-X} \quad & \rightarrow \quad \text{F-S=O-CO-X} \\
\text{F-} & \quad \rightarrow \quad \text{F-} \\
\text{F} & \quad \rightarrow \quad \text{F-} \\
\end{align*}
\]

X = CH\textsubscript{3}

The shortening of the S-C bond shifts it towards the bond length in F\textsubscript{4}S=C compounds. Furthermore, this shrinkage is
observed in the C-C bond (32b: 144.7 pm, 27: 154.4 pm). The C=O bond lengths in 27 and 32b are almost identical, which can be accounted for partially by the well-known conjugation in amides: 

\[ \text{O} - \text{C}-\text{NR}_2 \leftrightarrow \text{O}^- - \text{C}=\text{N}^+\text{R}_2, \]

which introduces the same structural feature in 27 as in 32b. The low C=O frequency in amides is supportive of this picture. In formamide \( d(\text{C}=\text{O}) = 124.3 \text{ pm} \), and \( d(\text{C}-\text{N}) = 131.9 \text{ pm} \), while in acetamide \( d(\text{C}=\text{O}) = 124.3 \text{ pm} \) and \( d(\text{C}-\text{N}) = 133.6 \text{ pm} \). One finds \( d(\text{C}-\text{N}) = 132.7 \text{ pm} \) and \( d(\text{C}=\text{O}) = 121.3 \text{ pm} \) in 27. One expects no noteworthy effect at the N-C\(_2\text{H}_5\) bond, and this is supported by the measured bond length of 147.4 pm (average), as compared to \( d(\text{C}'-\text{N}) = 146.3 \text{ pm} \) in methylformamide (this value is calculated). It appears as though the charge in 32b were distributed over the whole molecule. Hyperconjugation seems to play a role in this charge dispersal as concluded from the very substantial bond shortenings. This would be one of the very first direct evidences for this phenomenon. The bond lengths in 29, \((\text{F}_5\text{SCH(SO}_2\text{F})\text{CH}_2)_2\text{O}\), are comparable to 27, which validates the above comparison even further.

![Figure 3. X-ray structure of 32b, F\(_5\)SC(SO\(_2\)F)COOCH\(_3\)\(^-\) NH(C\(_2\)H\(_5\))\(_3\)\(^+\).](image-url)
Figure 4. X-ray structure of 27, F₅SCH(SO₂F)CON(C₂H₅)₂.

27 was obtained by an unexpected transmutation from what was probably F₅SCH=COCH₂O (28) upon storage.

### TABLE V

BOND LENGTHS [pm] AND ANGLES [°] IN (F₅SCH(SO₂F)CH₂)₂O, 29, STANDARD DEVIATIONS IN PARENTHESES

<table>
<thead>
<tr>
<th>Bond</th>
<th>Length [pm]</th>
<th>Bond</th>
<th>Length [pm]</th>
</tr>
</thead>
<tbody>
<tr>
<td>S(1)-F(1)</td>
<td>154.7(4)</td>
<td>S(1)-F(2)</td>
<td>155.9(3)</td>
</tr>
<tr>
<td>S(1)-F(3)</td>
<td>158.2(5)</td>
<td>S(1)-F(4)</td>
<td>156.4(3)</td>
</tr>
<tr>
<td>S(1)-F(5)</td>
<td>156.5(4)</td>
<td>S(1)-C(1)</td>
<td>185.8(5)</td>
</tr>
<tr>
<td>C(1)-C(2)</td>
<td>151.3(7)</td>
<td>C(1)-S(2)</td>
<td>178.9(5)</td>
</tr>
<tr>
<td>C(2)-O(3)</td>
<td>140.5(5)</td>
<td>O(3)-C(2A)</td>
<td>140.5(5)</td>
</tr>
<tr>
<td>S(2)-O(1)</td>
<td>149.3(4)</td>
<td>S(2)-O(2)</td>
<td>139.5(5)</td>
</tr>
<tr>
<td>S(2)-F(6)</td>
<td>141.7(4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F(1)-S(1)-F(2)</td>
<td>89.8(3)</td>
<td>F(1)-S(1)-F(3)</td>
<td>90.1(2)</td>
</tr>
<tr>
<td>F(1)-S(1)-F(4)</td>
<td>176.2(2)</td>
<td>F(1)-S(1)-F(5)</td>
<td>88.5(3)</td>
</tr>
<tr>
<td>F(1)-S(1)-F(5)</td>
<td>87.6(2)</td>
<td>F(1)-S(1)-F(6)</td>
<td>112.2(2)</td>
</tr>
<tr>
<td>F(2)-S(1)-F(3)</td>
<td>95.2(2)</td>
<td>F(2)-S(1)-F(4)</td>
<td>91.0(2)</td>
</tr>
<tr>
<td>F(2)-S(1)-F(4)</td>
<td>88.7(2)</td>
<td>F(2)-S(1)-F(5)</td>
<td>92.6(2)</td>
</tr>
<tr>
<td>F(2)-S(1)-F(6)</td>
<td>116.9(3)</td>
<td>S(1)-C(1)-C(2)</td>
<td>114.8(3)</td>
</tr>
<tr>
<td>C(1)-C(2)-O(3)</td>
<td>110.5(4)</td>
<td>C(2)-O(3)-C(2A)</td>
<td>112.0(5)</td>
</tr>
<tr>
<td>C(1)-S(2)-O(1)</td>
<td>101.8(2)</td>
<td>C(1)-S(2)-O(2)</td>
<td>105.4(3)</td>
</tr>
<tr>
<td>O(1)-S(2)-F(6)</td>
<td>108.4(3)</td>
<td>O(2)-S(2)-F(6)</td>
<td>117.7(3)</td>
</tr>
</tbody>
</table>
It is first noticed that the fragment F₅SCH(SO₂F)CH₂ in 29 has a very similar structure as in 27; the average d(S-Fe) = 156.3 ± 1.5 pm. For d(S-Fa) = 156.5 pm. The bond-length alternation within the group of the equatorial fluorines is considerable. This alternation is found with 27 and 32b as well as other SF₅ compounds. In the SO₂F group a strong bond length difference for the two S-O bonds is found. This is perhaps the result of an error as there is no obvious reason that could account for such a formidable difference. It is, however, noticed that the longer S-O bond is on the opposite side of the SF₅ group at the same carbon, and this oxygen is close to one hydrogen of the CH₂ group. At the same time the angles C-S-O are 101.8 and 105.4°, respectively, with the larger angle pertaining to the C-S-O triple with the longer
S-O bond. This behavior of the angles makes it unlikely that there is an O⋯H bond, because such a bond, although it could elongate the S-O bond, would also lead to a diminishment of the angle C-S-O. The angle at the ether bond is slightly larger than tetrahedral, and it is noticed that the hydrogens of the two CH₂ bridges are nearly eclipsed. The molecule has nearly C₂ symmetry (it is slightly distorted), with the C₂ axis lying in the C-O-C plane. The SF₅ groups are on opposite sides of the plane. The other alternative would have been C₂ symmetry with the axis perpendicular to the C-O-C plane and the SF₅ groups on the same side of it. This arrangement might be the result of the repulsion of the two SF₅ groups, which is smaller if they lie trans to each other. The bond lengths in the rest of the molecule are comparable to similar compounds. For example, d(S(F₅)-C) = 185.8 pm and 185.6 pm in 29 and 27 respectively, and d(S(O₂F)-C) = 178.9 (29) and 180.1 pm (27). The ether bond is almost of the same dimension as found in (F₅CH₂CHO)₃ (which will be described later in this manuscript), i.e. d(C-O) (29) = 140.5 pm, d(C-O) (trimer) = 141.1 pm (average).
CHAPTER V

SYNTHESIS OF PENTAFLUOROSULFUR BROMIDE, SF$_5$Br

Pentafluorosulfur bromide can be obtained by a number of methods. It was first synthesized by heating a mixture of BrF$_5$, Br$_2$ and SF$_4$ to 100°C for several days.$^{89}$ The easiest and fastest procedure consists of heating bromine and disulfur decafluoride to 140-150°C.$^{90}$ The difficulty lies in the accessibility of S$_2$F$_{10}$. Although it is a by-product in the fabrication of SF$_6$, it is usually immediately degraded, in order to obviate its noxiousness, by scrubbing the reaction gases with NaOH. For laboratory purposes it can be made through photochemical reduction of SF$_5$Cl with H$_2$.$^{91}$, where SF$_5$Cl, which was originally obtained by reacting Cs$^+$ SF$_5^-$ (from CsF + SF$_4$ upon heating, 94%) with chlorine$^{92}$ in 75% yield, or by reaction of KF, SF$_4$ and Cl$_2$ (mol ratio 3:1:1) in $\approx$ 80% yield$^{93}$, can now be readily obtained by the cesium fluoride catalyzed reaction of SF$_4$ and ClF in 92-97% yield.$^{94}$

\[
\text{SF}_4 + \text{ClF (CsF)} \quad \longrightarrow \quad \text{SF}_5\text{Cl}
\]

This difficulty in obtaining S$_2$F$_{10}$ easily in sufficient quantity, and the reluctance of some olefins to react with the more easily available SF$_5$Cl at all, made it necessary to seek a new method for preparing SF$_5$Br.
The method is based on a procedure, which employs the reaction of $SF_4$ and BrF. Since pure BrF is not stable above $\approx -30^\circ C$ it is first made by prolonged shaking of Br$_2$ and BrF$_3$, and isolated from the equilibrium mixture by low-temperature vacuum transfer. The equilibrium constant for the following (fast) gas phase reaction has been determined:

$$Br_2 + BrF_3 \rightleftharpoons 3 BrF \quad H = 11.9 \text{ kcal/mol}$$

The BrF is then heated with CsF and $SF_4$ to 100$^\circ$C (24h). The yield seems to be $\approx 90\%$ on a small scale$^{95}$ and 34\% on a larger scale.$^{96}$ Several reactions exist that employ the in situ formation of BrF, e.g. the formation of COBrF from Br$_2$, BrF$_3$ and CO (85\%)$^{97}$ or of SO$_2$BrF from SO$_2$, Br$_2$ and BrF$_3$ ($\approx$quantitative).$^{98}$ It seemed possible, that BrF, generated in situ, should react with $SF_4$ in the presence of CsF when shaken at room temperature. This reaction was indeed observed.

$$BrF_3 + Br_2 \rightleftharpoons 3 BrF$$
$$BrF + SF_4 -CsF-> SF_5Br (+SF_6)$$

Isolated yields of (pure) SF$_5$Br were 25\%, but it was usually used as a crude product, and the secondary products were then purified. In these instances yields of up to 70\% of distilled adducts of SF$_5$Br and various olefins were obtained. This method is thus comparable to the method of McDiarmid and Cohen for the synthesis of SF$_5$Br from S$_2$F$_{10}$ and Br$_2$ with yields to up to $\approx 80\%$. Purification is usually avoided, too, in this case, as bromine transfer at low
enough temperatures is slow, while SF\textsubscript{5}Br transfer is fast and the other possible constituents, SF\textsubscript{6}, SF\textsubscript{4} (decomposition products of S\textsubscript{2}F\textsubscript{10}) and unreacted S\textsubscript{2}F\textsubscript{10} are mostly neutral in the addition reactions that follow.

Sulfur hexafluoride was consistently observed as a by-product of the reaction; it must stem from the oxidation of SF\textsubscript{4} by bromine trifluoride. On occasion, the steel reaction vessel warmed up. Regarding this, and in light of the equilibrium between BrF\textsubscript{3}, Br\textsubscript{2} and BrF, which certainly lies on the BrF\textsubscript{3} side at 100°C, the method of Christe et al., as applied on a large scale by Seppelt et al., appeared unsafe when larger amounts of SF\textsubscript{5}Br were to be synthesized. The disadvantage of allowing the reaction to proceed at room temperature is its relative slowness. Shaking times of up to two weeks were necessary to obtain consistently good results, although on occasion good yields were already achieved after just two days of shaking. In order to preserve SF\textsubscript{4} (which is expensive), only 70-80\% of its theoretical amount was used, since it was observed, that the sulfur tetrafluoride that was added was never completely consumed, and, as the extent of oxidation to SF\textsubscript{6} is not known, this somewhat arbitrary molar ratio was used.

Several adducts were made by using the crude SF\textsubscript{5}Br that was isolated by removing it from the steel reaction vessel at -78°C. Long transfer times were necessary to collect most of the SF\textsubscript{5}Br. By this method, some Br\textsubscript{2} will always transfer,
too, and although no difficulties were encountered that would stem from the presence of bromine, it could be removed by treating the crude \( \text{SF}_5\text{Br} \) at room temperature with mercury\(^9\). It was possible to collect substantial additional amounts of almost pure \( \text{SF}_5\text{Br} \) from vessels that were warmed up intermittently and re-cooled. The chief impurity was then bromine, but \( \text{SF}_6 \) or \( \text{SF}_4 \) were present only in small amounts, as judged from the infrared spectrum. For example, runs with 25-30 g of \( \text{BrF}_3 \) yielded up to 20 g of \( \text{SF}_5\text{Br} \) in this second transfer. Whereas the initial transfer is comparatively fast (transfer times at \(-78^\circ\text{C}\) up to 4 hours), the second collection is very slow and lasted up to 10 hours.
CHAPTER VI

CONCLUSION

In conclusion, starting from sultone 1, intermediates 21, 23 and several salts of the form $Y^+ [F_5SC(SO_2F)COR]^- \text{ were obtained. An outstanding feature was the preferred formation of stable salts which were capable of further reaction. Halogenated compounds of the form } F_5SCX(SO_2F)COZ \text{ and } F_5SCH(Cl,Br)SO_2F \text{ (} X = Br, Cl; Z = F, Cl, OCH(CH_3)_2 \text{) were obtained but it was not possible to introduce other substituents } X, \text{ such as alkyl groups or cyano. Compound 23, a sulfur ylide, could be converted to several other sultones by polar addition of ClF, HF, HCl, CH_3OH and (CH_3)_2CHOH across the F_4S=C bond. These are unprecedented reactions in sultone chemistry, as the } \beta\text{-sultone ring stayed intact; this has not been described before. Cesium fluoride adds quantitatively to form 25. A novel rearrangement of esters of the form } F_5SCH(SO_2F)COOR \text{ to } (O=SF_2)=C(SO_2F)C(O)F \text{ was also found. These reactions provided routes to compounds which were inaccessible so far and are of potential value for the synthesis of sulfonic acids for fuel cells. The crystalline nature of several ester salts allowed to determine their structure by X-ray crystallography. An improved large-scale synthesis of SF_5Br was also found.}
PART II

SYNTHESIS AND REACTIONS OF SF₅-EPOXIDES
Hexafluoropropylene epoxide (47) is the most important of the fluorinated epoxides. The reasons are that it was one of the first perfluoro epoxides to be synthesized, and its reaction chemistry has been most widely investigated. It is comparatively readily available. The methods of its preparation include passing CF₃CF=CF₂ through a solution of KOH and H₂O₂ at low temperature, or a solution of a hypochlorite, treatment with KMnO₄ + anhydrous HF (−70°C, FMnO₃ is proposed as the oxidizing agent)¹ to the treatment with oxygen (or O₂+O₃ or O₂+Br₂ or Cl₂ = light mixtures) or organic peroxides². These syntheses can thus be divided into nucleophilic, electrophilic and radical procedures. A recent review is given in ref. 2, and a somewhat older one in ref. 7. Other fluoro epoxides were made by adding CH₂ or other carbene derivatives³ to ketones or aldehydes or (e.g. CF₂, C(CF₃)₂ to hexafluoroacetone). This can, for example, quite easily provide for octafluoroisobutylene epoxide⁴

\[(\text{CF}_3)_2\text{C}=\text{O} + \text{CF}_2 \longrightarrow (\text{CF}_3)_2\text{C}=\text{O}\text{CF}_2\]\n
The difluorocarbene originates mostly in the following reaction:
(\text{F}_3\text{CCFCF}_2 \text{ (47)}) \rightleftharpoons \text{CF}_3\text{CFO} + \text{CF}_2\text{ (5a,b) (at 150-180°C)}

On occasion, the reaction

\((\text{F}_3\text{C})_3\text{PF}_2 \rightleftharpoons (\text{F}_3\text{C})_2\text{PF}_3 + \text{CF}_2 \text{ F}_3\text{CPF}_4 \text{ etc. (ca. 130°C)}\)

is also used, but \((\text{F}_3\text{C})_3\text{PF}_2\) is difficult to obtain.

Other methods use haloacetates or halohydrines in the synthesis of oxiranes of the type \(\text{R}_f\text{CH}_{2}\text{CHCH}_2\text{O}, \text{R}_f\text{CHCH(R)}\text{O} \) (\(\text{R} = \text{alkyl}\)) and \(\text{R}_f\text{CHCH}_2\text{O}\). The halohydrines or haloacetates are obtained by different routes, i.e. bromination of a suitable ketone, and subsequent reduction,

\[
\begin{align*}
\text{CF}_3\text{CH}_3 & \rightarrow \text{CF}_3\text{CH}_2\text{Br} \rightarrow \text{CF}_3\text{CHCH}_2\text{Br} \rightarrow \text{CF}_3\text{CHCH}_2\text{Br},
\end{align*}
\]

treatment of an olefin with \(\text{Br}_2, \text{Hg(CH}_3\text{COO)}_2 + \text{CH}_3\text{COOH}, \text{and with base,}

\[
\begin{align*}
\text{R}_f\text{CHCH}_2\text{CH}_2\text{O} & \rightarrow \text{R}_f\text{CHCH}_2\text{CH}_2\text{O},
\end{align*}
\]
or addition of \(\text{R}_f\text{J}\) to allyl acetate or allylalcohol and treatment with base,

\[
\begin{align*}
\text{R}_f\text{I} + \text{CH}_3\text{COOCH}_2\text{CH} & \rightarrow \text{CH}_3\text{COOCH}_2\text{CHCH}_2\text{R}_f \rightarrow \text{R}_f\text{CHCH}_2\text{CH}_2\text{O}
\end{align*}
\]

A newer method employs treatment of olefins \(\text{R}_f\text{CF}=\text{CFR}_f, \text{R}_f\text{R}_f'\text{C}=\text{CFR}_f '\) or \(\text{R}_f\text{R}_f'\text{C}=\text{CFR}_f '\) with \(\text{NaOCl}\) in aqueous acetonitrile. This method can be suitably accommodated to phase transfer catalysis conditions and is also important because of the difficulty in epoxidizing internal fluoro olefins with other methods.

Epoxides are valuable intermediates. For unsymmetrical
terminal perfluoro epoxides, rearrangements to either acyl fluorides or fluoro ketones are possible. The formation of a ketone is brought about by acidic reagents (for example, SbF$_5$, $\gamma$-Al$_2$O$_3$, TiO$_2$, Cr$_2$O$_3$), and the formation of acyl fluoride by basic reagents, notably F$^-$. Some cyclic oxides rearrange to ketones with alumina,

\[
\begin{align*}
(CF_2)_n\begin{array}{c} CF \\ CF \end{array} & \quad \rightarrow \quad \begin{array}{c} CF \\ CF \end{array} \quad (C=O) \\
& \quad n = 3, 4 
\end{align*}
\]

Hexafluoropropylene oxide is converted by either alumina or SbF$_5$ to hexafluoroacetone; AlCl$_3$ leads chiefly to the ketone but yields also some pentafluoropropionyl fluoride. Perfluoroisobutylene (a highly electrophilic agent) also causes rearrangement to hexafluoroacetone. On the other hand, octafluoroisobutylene epoxide rearranges with antimony pentafluoride to perfluoroisobutyryl fluoride, as the corresponding ketone does not exist. The formation of a ketone or an acyl fluoride corresponds to two different modes of ring opening and can be related to two different sites of attack (in this case by F$^-$):

\[
\begin{align*}
R_fCF_3 & \quad \rightarrow \quad R_fCaF--CaF_2 \quad \rightarrow \quad R_fCF_2CFO \\
& \quad (\alpha\text{-attack}) \quad \beta\text{-attack}) 
\end{align*}
\]

As for the SF$_5$-epoxides of the form SF$_5$XCF$_2O$ (X=H, halogen), only $\alpha$-attack is of interest, because SF$_5$-ketones, formed in $\beta$-attack mode, are not stable towards nucleophilic attack; loss of SF$_5^-$ occurs. This restricts very severly
their synthetic utility.

Hexafluoropropylene oxide \(47\) can be used for many synthetic purposes, but in the present context its (anionic) oligomerisation behavior is especially important. This can be brought about by different agents, ranging from alkoxides, halide ions, neutral bases, even silver nitrate to activated carbon. The product distribution in the reaction

\[
(n+1) \quad F_3C\overline{C}CF_2O_{47} \quad \longrightarrow \quad CF_3CF_2[CF_2OCF]nCFO_{48}
\]

depends on the catalyst and solvent that is used.\(^{16}\) The value of \(n\) is found between 0 and 100. The process proceeds in the following fashion through the intermediacy of \(F^-\):

\[
\begin{align*}
F_3C\overline{C}CF_2F & \longrightarrow F_3CCF_2CFO \longrightarrow F_3CCF_2CF_2O^-; +
\end{align*}
\]

\[
\begin{align*}
F_3CCF_2[CF_2OCF]CF_2O^- & \longrightarrow F_3CCF_2[CF_2OCF]nCF_2O^- \longrightarrow
\end{align*}
\]

\[
F_3CCF_2[CF_2OCF]nCFO + F^- \quad \nonumber
\]

The polyethers are, after removal of the acyl fluoride group, chemically and thermally extremely inert, high-boiling and non-toxic and have a high dielectric strength.\(^{17}\)

A similar process accounts for the first step in the synthesis of "Nafion", which is a highly conductive perfluorinated polymeric sulfonic acid that enjoys now widespread use ranging from catalysis to fuel-cell electrolytes. The polymerisation is initiated by an alkoxide which is
obtained from tetrafluoroethylene sultone. The reaction then goes through a reductive decarboxylation step and is completed by co-polymerizing the product with tetrafluoroethylene and hydrolyzing the sulfonyl fluoride:

\[
\begin{align*}
0_2S^{-} -> & F_2SCF_2CF_2O^- \quad \text{-----} \\
& \text{CF}_3 \\
& \text{FO}_2\text{SCF}_2[\text{CF}_2\text{OCF}]_n\text{CFO}; + \text{Na}_2\text{CO}_3, \text{heat} \quad \text{-----} \\
& \text{CF}_3 \\
& \text{FO}_2\text{SCF}_2[\text{CF}_2\text{OCF}]_{n-1}\text{CF}_2\text{OCF}=\text{CF}_2 + \text{CF}_2=\text{CF}_2 \quad \text{-----} \\
& \text{---}[\text{CF}-\text{CF}_2]_x\text{---}[\text{CF}_2-\text{CF}_2]_y\text{---} \quad \text{R}_f=\text{SO}_3\text{H} \\
& \text{CP}_2[\text{CFOCF}_2]_{n-1}\text{CF}_2\text{SO}_2\text{F} \quad \text{49}
\end{align*}
\]

The pentafluorothio epoxides were synthesized in order to obtain SF₅ polyethers whence an analogous "SF₅-Nafion" could be synthesized. Possibly improved properties of this material could be higher heat-resistance, greater hydrophobicity, more elasticity (due to a diminished possibility of crystallization). Polymers of SF₅-epoxides are expected to be also highly heat-resistant. The ether linkage in perfluoroethers is thermally extremely stable. For example, the perfluoroether OC₂F₄SF₄C₂F₄ decomposes only above 475°C over NaF under extrusion of SF₄; octafluorotetrahydrofuran is the other product.¹⁸ These high boiling polyethers should also be good dielectric materials, as e.g. F₅SCF=CF₂¹⁹ (b.p. +19°C) and F₅SCF₃²⁰ (b.p. -7°C) are superior to SF₆ (b.p. -63°C).

The attempted synthesis of SF₅-epoxides has once been reported.²¹ The compound that was isolated in the following
reaction, with the oxidant under pressure, an acyl fluoride, cannot have originated in an intermediate epoxide:

\[
\text{F}_5\text{SCF}=\text{CF}_2 + \text{O}_2 \text{ (or O}_3\text{)} \rightarrow \text{F}_5\text{SOCF}_2\text{CFO}
\]

The easy availability of simple fluoroolefins of the form \(\text{F}_5\text{SCX}=\text{CF}_2\) \((X = \text{H} \text{ (52), F} \text{ (50)})\) makes it possible to obtain the epoxides

\[
\text{F}_5\text{S} \quad \bigg| \quad \text{F}.
\]

\((X = \text{H} \text{ (53), X = F (54)})\)

To further this investigation the next logical step would be to insert a \(\text{CF}_2\)-group between the epoxide ring and the substituent to obtain

\[
\text{F}_5\text{SCF}_2 \quad \bigg| \quad \text{CF}.
\]

This requires the availability of olefins of the form \(\text{F}_5\text{SCF}_2\text{CX}=\text{CF}_2\) \((X = \text{H}, \text{halogen, 56})\). To introduce a \(\text{CF}_2\)-group, several possibilities exist. It would also be desirable to have an olefin \(\text{FO}_2\text{SCF}_2\text{CF}=\text{CF}_2\) \((57)\), which upon epoxidation and polymerization should yield a perfluoro polyether sulfonyl fluoride, the direct precursor of a high-density sulfonic acid.
CHAPTER II

RESULTS AND DISCUSSION

An early attempt to obtain a haloacetate by the following reaction failed, although a preliminary reaction was encouraging:

\[ \text{F}_5\text{SCH}_2\text{CH}_2\text{Br} \ (58) + \text{AgOC(O)CH}_3 \rightarrow \text{F}_5\text{SCH}_2\text{CH}_2\text{OC(O)CH}_3 \ (59) + \text{AgBr} \]

but,

\[ \text{F}_5\text{SCHBrCH}_2\text{Br} \ (60) + \text{AgOC(O)CH}_3 \rightarrow \text{F}_5\text{SC} = \text{CH}_2 \ (61) \]

The silver salt reaction was carried out in glacial acetic acid, following a prescription of Park et al.\textsuperscript{23} It was thus decided to pursue direct epoxidation of olefins. The olefins of the form \( \text{F}_5\text{SCX} = \text{CF}_2 \) (X=H,F) are easily prepared. Therefore, the epoxides

\[
\begin{align*}
\text{F}_5\text{S} && \quad \text{X} && \quad \text{O} && \quad \text{F} \\
\end{align*}
\]

\( (X = H \ (53), F \ (54)) \)

should be accessible with a suitable epoxidation procedure. For \( X = F \) the epoxide is the exact analogue of hexafluoropropylene epoxide (47). Earlier attempts to epoxidize \( \text{F}_5\text{SCH} = \text{CH}_2 \) with m-chloroperbenzoic acid in \( \text{CH}_2\text{Cl}_2 \) in a pressure vessel at 90 °C were unsuccessful, as was the ozonisation of \( \text{F}_5\text{SCF} = \text{CF}_2 \), which lead to unidentified
pyrophoric products. Epoxidation was effected by taking advantage of the proclivity of fluorinated olefins towards nucleophilic addition. For example, F\textsubscript{5}SCF=CF\textsubscript{2}, when treated with CO\textsubscript{2}F in CH\textsubscript{3}CN and in the presence of CsF forms F\textsubscript{5}SCF(CF\textsubscript{3})COF\textsuperscript{24}. As the epoxidation of fluorocarbons with hypochlorite passes through an anion intermediate, and the SF\textsubscript{5} group can stabilize negative charges better than a CF\textsubscript{3} group\textsuperscript{25}, it seemed a promising pathway for the preparation of pentafluorothio epoxides. In a two-phase system this route was found effective with a phase-transfer catalyst:

\[
\text{F}_{5}\text{SCX}=\text{CF}_{2} + \text{NaOCl} \rightarrow \text{F}_{5}\text{SCX-CF}_{2} \rightarrow \text{F}_{5}\text{SCX-CF}_{2}
\]

\(\text{X} = \text{H (52), F (50)}\)

1,1,2-trifluorotrichloroethane ("Freon-113") was chosen as solvent for the reaction with F\textsubscript{5}SCF=CF\textsubscript{2} and CCl\textsubscript{4} for F\textsubscript{5}SCH=CF\textsubscript{2}. In the latter case it was easier to isolate the product by distillation, because CCl\textsubscript{2}FCCl\textsubscript{2} boils in the same region as the resulting epoxide. The phase transfer catalyst \(N(n-\text{C}_{8}\text{H}_{17})_{3}\text{CH}_{3}^{+}\text{Cl}^{-}\) was used in a concentration of 10-20 mol %. Temperature control (\(-6^\circ\text{C}\)) and slow addition of the olefin to the cold two-phase system consisting of the phase-transfer catalyst, solvent and basic NaOCl solution, were especially important. Too fast addition led to a temperature rise and the solution became acidic. In these cases no fluorinated products were found. The reactions were monitored by \(\text{F}^{19}\) n.m.r. and terminated with the disappearance of the olefin. In the case of the
perfluoroolefin a by-product was observed which according to its band shape and chemical shift was identified as \(S_2F_{10}\).

It was not possible to obtain until now an olefin of the form \(F_5S(CF_2)_nCX=CF_2\) \((n=1,2, X = H, \text{ halogen, } 56a)\). The following reaction proved unsuccessful:

\[
F_5SCF_2CF_2I + CHF=CF_2 + (\phi COO)_2 \rightarrow F_5SCF_2CF_2CHFCF_2I
\]

Subsequent dehydroiodination should have provided for a terminal olefin. What was observed was at temperatures up to \(170^\circ C\) loss of the \(SF_5\) group and formation of \(ICF_2CF_2I\), but none of the desired adduct.

A report by Witucki shows the low yield addition of \(SF_5Cl\) to tetrafluoroallene: 26

\[
F_5SCl + F_2C=C=CF_2 \rightarrow F_5SCF_2C=CF_2
\]

The synthesis of tetrafluoroallene turns out to be a comparatively complicated endeavour and it was decided that this route, after several costly failures, was not suitable to provide for sizable amounts of epoxides. Carbene addition to the acyl fluoride \(F_5SCF_2CFO\) (65) was also considered. \(F_5SCF_2CFO\) was first described by Haszeldine and Nyman27 as one of the products of the electrochemical fluorination of thioglycollic acid, but it was obtained in a mixture and could only partially characterized. An attempt to obtain this material by sulfur trioxidation of \(F_5SCF_2CF_2J\) was made:

\[
F_5SCF_2CF_2J \text{ (63)} + \text{ oleum + heat } \rightarrow F_5SCF_2CFO \text{ (65)}
\]
Although the reaction worked rather well, the product that was obtained was not pure. Upon cooling the colorless liquid separated into two layers. Despite lengthy trials to separate these materials through flash-evaporation, fractional condensation or preparative gas chromatography, it was not possible to effect separation. Little can be said about the nature of this second constituent, except that it has a singlet in the $^{19}$F n.m.r. spectrum at $\approx -80\text{ppm}$. It is not an SF$_5$ nor a COF compound.

It was then tried to use this crude ($\approx 70\%$) material for further difluorocarbene addition. The CF$_2$ was either generated by heating hexafluoropropylene oxide or by the method of Burton (treatment of the salt $\phi_3\text{PCF}_2\text{Br}^+ \text{Br}^-$ with KF in diglyme at $20^\circ\text{C}$)\textsuperscript{28}. In no case was any product attributable to the reaction

$$\begin{align*}
\text{F}_5\text{SCF}_2\text{CFO}^\phi_{65} + \text{CF}_2 & \rightarrow \text{F}_5\text{SCF}_2\text{CFCF}_2^\phi_{66}
\end{align*}$$

identified. Further attempts, like the addition of SF$_5$Br to CF$_2$=CFCF$_2$OSO$_2$F\textsuperscript{29}, with the possible subsequent treatment of the adduct with fluoride ion failed because of the inertness of the olefin towards SF$_5$Br.

Another attempt to have a CF$_2$ group inserted between the SF$_5$ group and the epoxide was undertaken by first obtaining a hydrogenated epoxide, which then could be perhaps fluorinated. Such an attempt had apparently been undertaken\textsuperscript{30} with ClCH$_2$CHCH$_2$Cl which should have yielded hexafluoro propylene epoxide upon fluorination, but was not
successful. The electrochemical fluorination procedure that was used might not have been appropriate, however, as the product was most likely an oxetane\(^3\). For the present purpose \(SF_5Cl\) was added to allyl acetate. This was a surprisingly slow reaction, as after two weeks of heating the two reactants in \(CCl_3F\) at 100°C there was still unreacted olefin present and only a yield of \(\approx 60\%\) could be realized:

\[
F_5SCl + CH_2=CHCH_2OC(O)CH_3 \rightarrow F_5SCH_2ClCH_2OC(O)CH_3 \quad (67)
\]

Base treatment of the analogous \(R_fI\) addition product gave the epoxide \(R_fCH_2CHCH_2O\) (see above), but in the present case the product was an alcohol, apparently formed by the simultaneous elimination of \(HCl + CH_3COOH\):

\[
F_5SCH_2ClCH_2OC(O)CH_3 + \text{base} \rightarrow F_5SCH=CHCH_2OH \quad (\approx 90\%), \quad (68)
\]

The elimination was originally conducted with aqueous \(KOH\) (in a two-phase system), but it appeared as though water were retained tenaciously by the product. This was chiefly concluded from the i.r. spectrum which shows a broad feature coincident with the water spectrum. Running the reaction under anhydrous conditions (with sodium methoxide in methanol) yielded the same product, however.
CHAPTER III

REACTIONS OF F$_5$SCFCF$_2$O

Most of the reactions of the epoxide F$_5$SCFCF$_2$O (54) were not quantitatively analyzed because of the impossibility of separating the products that were formed. Nevertheless, one can draw one major inference from these experiments. In all reactions loss of SF$_5$ was observed and SF$_4$ and CF$_3$CFO could be identified as major products by their characteristic infrared frequencies. Similar observations have been made before by DesMarteau and Sekiya$^{32}$ and by Berry and Fox. In one mode of reaction (attack at CF$_2$), the compound F$_5$SNCF$_2$O was found to lose SF$_5^-$ upon base-treatment,

\[ F_5SN-CF_2 + F^- \rightarrow SF_4 + CF_3NO + COF_2 + \text{others} \]

COF$_2$ would originate in the attack at nitrogen. The loss of SF$_5^-$ was inferred from the presence of SOF$_2 + SiF_4$; SF$_4$ was not observed, but SOF$_2$ is the usual hydrolysis product. In the other case, F$_5$SCCF$_3$ (62) was found to lose SF$_5^-$ when bases were present. There is a parallelism between the case of F$_5$SCFCF$_2$O (54) and the isomer F$_5$S(CF$_3$)C=O. While all anions seem to attack the epoxide at the CF$_2$ group, leading to an intermediate anion, the attack in the case of the
ketone is at the CO group, leading to the same intermediate:

\[
\text{F}_5\text{S}^\ominus\text{C-}\text{CF}_3^\ominus\text{+F}^- \rightarrow \quad \text{F}_5\text{S}^\ominus\text{C-}\text{CF}_3^\ominus \leftarrow \text{F}^- \quad \text{F}_5\text{S}\text{CFCF}_2\text{O}^\ominus
\]

\[
\text{CF}_3\text{CFO} \quad \text{SF}_5^- \rightarrow \text{SF}_4^- \quad \text{+F}^- \quad \text{SF}_5^- \quad \text{F}_5\text{CFCF}_2\text{O}^\ominus
\]

In the above case, the expected product would be CF\text{3CFO} + SF\text{4}. F\text{5SCFCF}_2\text{O}\text{ as an analogue to hexafluoro propylene epoxide and should exhibit similar reaction behavior.}\text{33}

While CF\text{3CFCF}_2\text{O} reacts with methanol to give methyl-2-methoxytetrafluoro propionate, its SF\text{5 analogue does not lead to a corresponding product:}

\[
\text{F}_3\text{CCFCF}_2\text{O} + \text{CH}_3\text{OH} \rightarrow \text{F}_3\text{CF(OCH}_3\text{)}\text{COOCH}_3 \quad (96\%)
\]

\[
\text{F}_5\text{SCFCF}_2\text{O} + \text{CH}_3\text{OH} \rightarrow \text{F}_5\text{SCF(OCH}_3\text{)}\text{COOCH}_3
\]

The product that is obtained is a mixture of several compounds, of which one constitutes 58% (g.c.). These products could not be identified due to the small amounts of the epoxide that were available, but the SF\text{5 group was destroyed, as seen in the 19F n.m.r. spectrum.}

Water, which leads with hexafluoropropylene oxide to 2,2-dihydroxytrifluoropropionic acid, does not seem to react appreciatively with F\text{5SCFCF}_2\text{O}.

\[
\text{F}_3\text{CCFCF}_2\text{O} + \text{H}_2\text{O} \rightarrow \text{F}_3\text{CC(OH)}_2\text{COOH} \quad (68\%)
\]

\[
\text{F}_5\text{SCFCF}_2\text{O} + \text{H}_2\text{O} \rightarrow \text{no appreciable reaction}
\]

The reaction was carried out at slightly raised temperature in a Pyrex-glass vessel with a large excess of water, and the progress of the reaction was monitored by measuring the gas pressure occasionally and checking the i.r. spectrum of
the gaseous material. There was a slight initial pressure drop (≈ 13 %), which might have its origin in the solubility of the epoxide in water. The change in the i.r. spectrum was confined to the appearance of one weak band at 1374 cm⁻¹, and after 1 week circa 80 % of the epoxide could be recovered. The reaction vessel was severely etched, and a white material had been dislodged from it. This residue, although not identified, originated certainly in the reaction of the glass vessel with hydrogen fluoride. The epoxide does seem to react initially with water, releasing HF, which turns the reaction mixture acidic, halting further reaction, but allowing the HF to etch the vessel in a continuous fashion.

The epoxide is completely inert to boron trifluoride; ring opening does not occur.

The reaction with cesium fluoride in acetonitrile showed the formation of mainly SF₄ and CF₃CFO.

\[ \text{F}_5\text{SCF}_2 \text{CF}_2 + \text{CsF} + \text{CH}_3\text{CN} \rightarrow \text{SF}_4 + \text{CF}_3\text{CFO} + \text{other compound(s)} \]

Preparative gas chromatography showed also the formation of another (minor) product, but it could not be identified. The reaction was conducted with heating and it was first tried to separate the products by fractional condensation. This separated the solvent from the mixture, but the SF₄ and CF₃CFO were collected together in the -196°C trap (76 % of theory). It was also not possible to separate the products by g.c., but the eluted band was clearly asymmetrical and
when an i.r. spectrum at different stages of the elution was measured, the changing composition of the eluent was clearly seen. SiF₄, SOF₂ and CF₃CFO could be identified, but no information concerning their amounts was obtained. The fourth product (≈ 6% of total area) has strong bands at 816 and 909 cm⁻¹, and also in the C-F stretch region (1047-1213 cm⁻¹). This could indicate a compound that has S-F and C-F bonds, but the appearance and position makes it unlikely that an SF₅ group is present. The few bands that are present indicate that it is a simple compound. Whether it originates in F⁻ attack at CF instead of CF₂ (the origin of SF₄ and CF₃CFO) must remain an unanswered question.

Further attempts to detect CsSF₅ (which should be a primary product of decomposition by CsF) were unsuccessful. When the reaction was conducted at room-temperature in the course of several weeks, blackening was observed, but the residue, when treated with Br₂ did not yield SF₅Br. DesMarteau and Sekiya showed the presence of CsSF₅ by treating the solid residue with fluorine:

\[ \text{CsSF}_5 + F_2 \rightarrow \text{CsF} + \text{SF}_6, \]

and Seppelt, Fuchs and Bittner³⁴ could isolate CsSF₅ and RbSF₅ from the reaction of the respective fluorides and F₅SC(CF₃)≡CF₂:

\[ \text{F}_5\text{S} \equiv \text{CF}_2 + \text{CsF} \rightarrow \text{F}_5\text{S} \equiv \text{CF}_2 \text{C}^+ \rightarrow \text{CsSF}_5 + \text{F}_3\text{C} \equiv \text{CF}_3 \]

These two reactions show that loss of SF₅⁻ is possible when
other stable product can be formed at the same time. The weight increase of the cesium fluoride in the reaction with $\text{F}_5\text{SCFCF}_2\text{O}$ was only 24% of the expected value.

The reaction of $\text{F}_5\text{SCFCF}_2\text{O}$ with $\text{SbF}_5$ did not lead to the formation of $\text{F}_5\text{SC(0)CF}_3$. Formation of C=O compounds (carbonyl fluorides or ketones) is observed other with perfluoroolefin epoxides$^{35}$. Instead, the i.r. spectrum of the product shows the presence of $\text{CF}_3\text{CFO}$, $\text{SiF}_4$ + perhaps CO ($2185 \text{ cm}^{-1}$). The presence of other products is also possible. The products were not further analyzed except by qualitative view. That $\text{SF}_4$ was not observed comes from the following reaction:

\[ \text{SbF}_5 + \text{SF}_4 \longrightarrow \text{SF}_3^+ + \text{SbF}_6^- \]

A conversion into the isomers $\text{F}_5\text{SCF}_2\text{CFO}^{36}$ or $\text{F}_5\text{SC(0)CF}_3$ is thus not possible by this route. When $\text{F}_5\text{SCFCF}_2\text{O}$ was heated in the presence of oxygen, no reaction was observed at 50 °C, but two days of heating at 130-140°C produced a strong band at 1898 cm$^{-1}$ and there were further bands attributable to $\text{SF}_4$, $\text{SOF}_2$ and $\text{SiF}_4$. No $\text{F}_5\text{SOCF}_2\text{CFO}$ was detected; this product could have been formed in a similar fashion as was found in the reaction of $\text{F}_5\text{SCF}=\text{CF}_2 + \text{O}_2$. Although the epoxide was not subjected to heat alone, it is quite possible that this is the effect of thermal decomposition rather of the involvement of oxygen.

Bases as weak as diethylether will cause this reaction: When $\text{F}_5\text{SCFCF}_2\text{O}$ is dissolved in $(\text{C}_2\text{H}_5)_2\text{O}$ slow
formation of SF₄ is observed. In the case of tetrafluoroethylene epoxide this reaction results in the formation of a diether ROCF₂CF₂OR.³⁷

These reactions show that a preferred mode of action does involve attack at the CF₂ group with the following loss of SF₅⁻ (→ SF₄ + F⁻), always leading to SF₄ and CF₃CFO.

This loss of SF₅ was also observed with F₅SC(O)CF₃³⁸ and bases, and is also similar to the observation made by DesMarteau and Sekiya with F₅SNCF₂O, and as described above, by Seppelt, Bittner and Fuchs with F₅SC(CF₃)₂⁻ Cs⁺.

Although F₅SCHCF₂O was also synthesized, its reactivity was not investigated. It is expected that it will lose SF₅⁻ even more readily than F₅SCFCF₂O in anionic reactions, because it lacks the α-fluorine which might stabilize a negative charge at Cₐ.

SYNTHESIS OF THE STARTING OLEFINS

The two olefins that were used as starting materials were synthesized by modified published versions. F₅SCH=CF₂ (52) was obtained as described in conjunction with the synthesis of F₅SCHCF₂OSO₂, while F₅SCF=CF₂ (50) was obtained by dehydrobromination of F₅SCHCF₂Br with aqueous KOH, also in excellent yield. The more common method of dehydrohalogenation with solid KOH or with KOH in petroleum ether is less advantageous, because it is more time consuming and yields are lower. With the analogous
F₅SCHFCF₂Cl, it was reported that decomposition occurred with aqueous KOH.³⁹
CHAPTER IV

INFRARED, $^{19}$F AND $^{13}$C N.M.R. SPECTRA

In the infrared spectra the epoxide ring band is seen at 1530 (SF$_5$CFCF$_2$O) and $\approx$1500 cm$^{-1}$ (SF$_5$CHCF$_2$O), respectively, at the upper limit of the values observed for other fluorinated epoxides. The $^{19}$F n.m.r spectra of the two epoxides show the typical AB$_4$ spectra of the SF$_5$ group. It is conspicuous that the SF$_5$ resonances in the epoxides have moved upfield, as compared to the olefins, by 6-10 ppm. The CF$_2$ groups in F$_5$SCFCF$_2$O and in F$_5$SCHCF$_2$O appeared as AB and MX system, respectively; in the case of the perfluoro compound this system was rather complicated due to extensive overlap. There was also an explicit dependence on the solvent. For example, the two fluorines in the CF$_2$ group of SF$_5$CFCF$_2$O had a very small chemical shift difference in the pure sample and appeared as an incomprehensible multiplet; in 1,1,2-trifluorotrichloroethane a complicated, partially overlapped, but interpretable, AB system was observed, while in CDCl$_3$ the two resonances were clearly separated. It was possible to reproduce the CF$_2$ resonance with an n.m.r. simulation program.$^{40}$ The coupling within the CF$_2$ group in hexafluoropropylene oxide is 43.5 Hz$^{41}$, considerably much larger than in SF$_5$CFCF$_2$O, but comparable to the value in
The $\text{-}\alpha,\beta$-trans coupling in fluoro olefins and in other fluorinated epoxides is found to be negative$^{14}$.

Most likely, $J_{XF}$ is also negative in the present case, but it is given here only as the absolute value.

![Figure 6. Recorded (a, in CDCl$_3$) and simulated (b) AB part of the $^{19}$F n.m.r. spectrum of SF$_5$CFCF$_2$O. Linewidth = 2.6 Hz. $\phi_F^1-\phi_F^2 = 1.02$ppm.](image)

In the $^{13}$C n.m.r. spectra of both epoxides, no unusual shifts are observed; the resonances and coupling constants for both carbons appear in the expected regions, and are comparable with the shifts and couplings in the $\beta$-sultone F$_5$SCHCF$_2$OSO$_2$ and other (partially) fluorinated oxiranes, with the exception of $J_{CaFa}$ in SF$_5$CFCF$_2$O.
TABLE VI

$^{19}$F AND $^{13}$C N.M.R. DATA† OF SF$_5$-EPOXIDES

<table>
<thead>
<tr>
<th></th>
<th>$F_5SCF$-CF$_2$</th>
<th>$F_5SCH$-CF$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\phi_A$</td>
<td>60.3 (0.95)</td>
<td>68.1 (1.00)</td>
</tr>
<tr>
<td>$\phi_B$</td>
<td>46.2 (4.33)</td>
<td>53.0 (3.90)</td>
</tr>
<tr>
<td>$\phi_F^1$</td>
<td>110.5 (m) (2.06)</td>
<td>-119.3 (d-d) (0.90)</td>
</tr>
<tr>
<td>$\phi_F^2$</td>
<td>&quot; (m)</td>
<td>-105.2 (d-d-p) (1.00)</td>
</tr>
<tr>
<td>$(\phi, \delta)_X$</td>
<td>-129.0 (d-septet) (1.00)</td>
<td>5.30 (m)</td>
</tr>
<tr>
<td>$J_{AB}$</td>
<td>152.3</td>
<td>153.3</td>
</tr>
<tr>
<td>$J_{AX}$</td>
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<td>-</td>
</tr>
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</tr>
<tr>
<td>$J_{BF^2}$</td>
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</tr>
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</tr>
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<td>4.1</td>
</tr>
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<td>44.0</td>
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<td>$\delta_{C\alpha}$</td>
<td>$\approx$107.4 (d-m)</td>
<td>81.45 $\approx$(d-p) *</td>
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<tr>
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<td>108.40 (t)</td>
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<td>$\approx$337 Hz</td>
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</tr>
<tr>
<td>$J_{CF^2C\beta}$</td>
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<td>295.6</td>
</tr>
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</table>

$[^{\phi}] = \text{ppm}; [J] = \text{Hz};$ band areas are in parentheses. †Neat sample. ‡$^{13}$C spectra are run in CDCl$_3$ solution. *The individual bands show further structure, but they are only ill-resolved. **The coupling constants for the CF$_2$ fluorine atoms were determined from the simulated spectrum in CDCl$_3$ solution (Figure 6).
CHAPTER V

CONCLUSION

Two SF$_5$-epoxides were synthesized and the reaction chemistry of the perfluoro compound was investigated in more detail. It was found that loss of the SF$_5$ group was the dominant reaction with nucleophiles. It was not possible to polymerize this epoxide. Attempts to synthesize an olefin with a CF$_2$ group between the double bond and the SF$_5$ group failed; this should have diminished the tendency for losing the SF$_5$ moiety.
PART III

TOWARDS THE SYNTHESIS OF PENTAFLUOROTHIO MALONIC ACID
CHAPTER I

INTRODUCTION

Pentafluorosulfur compounds have not yet gained much use; nevertheless, the \( \text{SF}_5 \) group does exhibit some rather intriguing properties, which make it comparable, and in some respects superior to a \( \text{CF}_3 \) group. Comparing the n-octanol-water partition coefficients of a compound bearing different substituents, one obtains the hydrophobicity coefficient, which indicates the affinity of a substituent towards a hydrophobic medium such that a larger number corresponds also to a larger affinity towards the hydrophobic phase. In this series, \( \text{SF}_5 \) is only superseded by trans-\( \text{CF}_3 \text{SF}_4 \). Such a property is desirable in several respects. The incorporation of fluorine into biologically active molecules has several motives. A fluorine atom is only slightly larger than a hydrogen atom (van der Waals radii, \( \text{F} = 1.35 \text{Å}, \text{H} = 1.2 \text{Å} \)), but has rather different properties. If the fluorine atom is in a position of the molecule that is at some point subject to chemical change in a biochemical transformation, then the normal reaction sequence will be interrupted. Another reason for introducing fluorine is the interruption of a biochemical cycle by the formation of poisonous by products in a chemical reaction; this can be the case in the
hydrolysis of a CF₃ group. These rationales have brought about the synthesis of certain nucleoside analogues, as 5-fluorouracil, 5-fluoro-2'-deoxyuridine, 5-fluorocytosine and 5-trifluoromethyl-2'-deoxyuridine², which are used as anti-viral and cancerostatic drugs. The activity of 5-trifluoromethyl-2'-deoxyuridine seems to be related to its incorporation into the cell DNA. Other derivatives of deoxyuridine were also investigated, as the 5-pentafluoroethyl and 5-enneafluoro-n-butyl-2'-deoxyuridine, but were found to have only lessened activity against herpes simplex virus (type 1 and 2)³. 5'-trifluoromethylthioribose⁴, a sugar analogue, shows very high (in-vitro) efficacy against the malaria parasite. A third reason is to increase the lipid solubility of certain drugs, which may be, according to Filler, the most significant factor in improving pharmacological activity.⁵ Discounting for particular properties of the SF₅ substituent, which are not investigated and are possibly absent, due to its usual chemical inertness, it is thus expected that drugs, in particular central nervous system agents, for which the increase of lipophilicity is considered specially important, due to the increase of absorption and transport across the brain-blood barrier, containing this substituent would be more efficacious than their CF₃ (or F) analogues. In the case of 5-trifluoromethyl-2'-deoxyuridine rapid hydrolysis of the
trifluoromethyl group upon administration is observed, with the formation of inorganic fluoride and 5-carboxyuracil. Under certain conditions, an SF$_5$-group is, however, also subject to chemical degradation, either under reducing or basic conditions.

Several SF$_5$-compounds have been patented as fumigants, SF$_5$CF=CF$_2$, SF$_5$CFICF$_3$, SF$_5$N=CF$_2$ and as a soil fumigant, SF$_5$CH$_2$COOH. Insecticidal properties are described for compounds derived from SF$_5$Cl and ethylene or propylene.
CHAPTER II

A GENERAL OUTLINE FOR THE SYNTHESIS OF SF₅-COMPOUNDS

Most SF₅ compounds are obtained by addition of SF₅Br or SF₅Cl to olefins. This method allows one to obtain SF₅-compounds with the SF₅-group at a secondary carbon only in exceptional cases. On the other hand it is not usually possible to manipulate simple SF₅-compounds chemically towards the introduction of a carbon substituent at the SF₅-carbon. Examples for this failure have been given in the first part of this manuscript.

Whereas in CF₃ chemistry a number of reactions already exist, that allow CF₃-substitution in suitable substrates, in the SF₅-field only few such methods are available:

1. Addition of SF₅Cl or SF₅Br to multiple bonds.¹¹
2. Addition of S₂F₁₀ to multiple bonds.¹²
3. Oxidation of disulfides with AgF₂ or CoF₃.¹³
4. Anodic fluorination of thiols or disulfides.¹⁴
5. Direct fluorination.¹⁵

Of these methods the most suitable ones are the addition of SF₅X to multiple bonds, while oxidations of -SH or -SS- are used for special classes of compounds.

The addition of SF₅X to a multiple bond, which can be induced by light, heat or free-radical initiators, is a
radical reaction and involves in the first stage cleavage to 
$\text{SF}_5\cdot$ and $X\cdot$, followed by addition to the multiple bond.$^{16}$ 
The chain reaction is maintained by reaction of the primary 
product with $\text{SF}_5X$:

$$\begin{align*}
\text{SF}_5X & \quad\longrightarrow\quad \text{SF}_5\cdot + X\cdot \quad (X=\text{Br,Cl}) \\
X\cdot + \text{C}=\text{C} & \quad\longrightarrow\quad \text{X-C-C}\cdot \quad (a) \\
\text{SF}_5\cdot + \text{C}=\text{C} & \quad\longrightarrow\quad \text{F}_5\text{S-C-C}\cdot \quad (b) \\
\text{F}_5\text{S-C-C}\cdot +\text{SF}_5X & \quad\longrightarrow\quad \text{F}_5\text{S-C-C-X} + \text{SF}_5\cdot
\end{align*}$$

Only reaction $(b)$ will lead to propagation, as in the next 
step both $(a)$ and $(b)$ will react with $\text{SF}_5X$, abstracting $X\cdot$
and leaving $\text{SF}_5\cdot$ behind. The reaction step $(a)$ continues 
with $(b)$ in this second step, explaining why neither $\text{X-C-C-X}$
nor $\text{F}_5\text{S-C-C-SF}_5$ are usually observed as products in these 
reactions.$^{17}$ It was shown later, that the addition of $\text{SF}_5\cdot$
to an olefin is reversible$^{18}$, an observation that was also 
made upon occasion in the course of this work.

The addition of $\text{SF}_5X$ $(X=\text{Br,Cl})$ to olefins or acetylene 
leads only to compounds which contain the $\text{SF}_5$-group at the 
carbon not bearing a substituent:

$$\begin{align*}
\text{CH}_2=\text{CHCH}_3 + \text{SF}_5X & \quad\longrightarrow\quad \text{F}_5\text{SCH}_2\text{CHXCH}_3 \\
\text{CH}_2=\text{CF}_2 + \text{SF}_5X & \quad\longrightarrow\quad \text{F}_5\text{SCH}_2\text{CF}_2\text{X} \\
\text{CH}_2=\text{CHCOOC}_2\text{H}_5 + \text{SF}_5X & \quad\longrightarrow\quad \text{F}_5\text{SCH}_2\text{CHBrCOOC}_2\text{H}_5 \\
\text{CH}_2=\text{CHOAc} + \text{SF}_5X & \quad\longrightarrow\quad \text{F}_5\text{SCH}_2\text{CHXOAc}
\end{align*}$$

$\text{SF}_5\cdot$ is an electrophilic radical and will thus add 
preferentially to positions with higher charge density. This 
is a chief obstacle in making a whole class of $\text{SF}_5$
compounds, that are additionally substituted at the carbon bearing the SF$_5$-group. Although it is possible to obtain compounds of the form SF$_5$CHX- (X=H,Br,Cl,F), this must be considered an exception, as with an asymmetrically substituted ethylene (CF$_2$=CHCl e.g.) the addition will not be uni-directional.

Only in special cases can C$_\alpha$ be substituted with something other than hydrogen or halogen:

\[ \text{F}_3\text{CCH}=\text{CF}_2 + \text{SF}_5\text{Cl} \text{(hv)} \longrightarrow \text{(F}_3\text{C)}\text{F}_5\text{SCHCF}_2\text{Cl}^{19} \]

\[ (\text{F}_3\text{C})_2\text{C}=\text{CH}_2 + \text{SF}_5\text{Cl} \longrightarrow (\text{F}_3\text{C})_2\text{F}_5\text{SCCH}_2\text{Cl}^{20} \]

\[ \text{F}_5\text{SCH}=\text{CF}_2 + " ~" \longrightarrow (\text{F}_5\text{S})_2\text{CHCF}_2\text{Cl}^{21} \]

\[ \text{H}_3\text{C}=\text{COR} + " ~" \longrightarrow \text{F}_5\text{S}(\text{CH}_3)\text{C}=\text{C}(\text{Cl})\text{OR}^{22} \]

\[ \text{F}_5\text{SC}=\text{CH} + \text{CH}_2\text{N}_2 \longrightarrow 3\text{-SF}_5\text{-pyrazol} (60\%) + \text{4-SF}_5\text{-pyrazol} (40\%)^{23} \]

\[ \text{F}_5\text{SC}=\text{CF}_2 + \text{CH}_2=\text{CHCH}=\text{CH}_2 \longrightarrow \text{F}_5\text{SCF}=\text{CF}_2 \text{H}_2\text{C}--\text{CHCH}=\text{CH}_2^{24} \]

\[ \text{F}_5\text{SC}=\text{CF}_2 + \text{COF}_2 \text{(CsF)} \longrightarrow \text{F}_5\text{SCFCF}_3^{25} \]

\[ \text{F}_5\text{SC}=\text{CF}_2 + \text{SO}_3 \longrightarrow \text{F}_5\text{SCXCF}_2 \text{X=H,F}^{26a,b} \]

\[ \text{F}_5\text{SCF}(\text{CF}_3)\text{I} + \text{C}_2\text{H}_4 \text{(hv)} \longrightarrow \text{F}_5\text{SCF}(\text{CF}_3)\text{CH}_2\text{CH}_2\text{I}^{27} \]

It is obvious, that steric requirements in the addition reactions of SF$_5$Cl are of minor importance, for otherwise the direction of addition should have been the opposite.

It was intended to synthesize the compound 5-SF$_5$-uracil, as the analogue of 5-trifluoromethyl-uracil. These compounds are simply related to the respective 2'-uridines.

An important intermediate for syntheses would be
pentafluorothio malonic acid, $F_5SCH(COOH)_2$ or its esters. As with other malonic acid derivatives, this should allow the synthesis of pyrimidine derivatives, in particular barbituric acids, and further manipulation of $F_5SCH(COOR)_2$ ($R = H$, alkyl) should lead to $\alpha$-$SF_5$ acrylic acid. The CF$_3$-analogue, CF$_3$CH(COOC$_2$H$_5$)$_2$, was synthesized some years ago by Knunyanz$^{28}$, its di-acyl fluoride was described by Krespan$^{29}$, and CH$_2$=C(CF$_3$)COOH by Japanese researchers. Trifluoromethyl malonic ester was obtained by successive methanolysis of (CF$_3$)$_2$CHCOOCH$_3$; trifluoromethyl malonyl fluoride was obtained by reaction of acetic anhydride with $\alpha$-CF$_3$ perfluoroacryloyl fluoride, and $\alpha$-CF$_3$ acrylic acid by a catalytic carbonylation procedure of 2-bromo-3,3,3-trifluoroproylene$^{30}$ 1,2-dibromo-3,3,3,-trifluoropropane$^{31}$ in good yield. 5-trifluoromethyluracil or 5-trifluoromethyluridine are accessible by several different routes: by ring-closure$^{32}$, replacement of halogen by reaction with copper perfluoroalkyl compounds$^{33}$ or by a radical substitution of the hydrogen at position 5 in pyrimidines$^3$. In order to achieve this goal for the SF$_5$ analogue, a suitable reagent for cyclization - most advantageously an acrylic or malonic acid - is necessary; an analogous reaction to the copper catalyzed R$_f$-substitutions for SF$_5$ groups has not been devised.
CHAPTER III

RESULTS AND DISCUSSION AND THE SYNTHESIS OF
SF$_5$CH(COOC$_2$H$_5$)CHBROCOCH$_3$

The introduction of an $\alpha$-carbon substituent in SF$_5$-compounds should be possible by several routes. A particularly easy reaction would have been the addition of SF$_5$Br to maleic anhydride. This compound was completely inert towards SF$_5$Br, probably due to the electrophilic double bond. Although the equilibrium $\text{OClCCCH}_2\text{CClO} \rightleftharpoons \text{OClCCCH=Cl} + \text{HCl}$ was described$^{34}$, no product corresponding to SF$_5$Cl addition to malonylchloride (more exactly, to chlorocarbonyl ketene) was observed in the respective reactions at $\approx 90^\circ\text{C}$ in a steel bomb; only decomposition was seen.

Nucleophilic displacement of halogen by cyanide was another possibility that was considered:

$$\text{F}_5\text{SCHBrCH}_2\text{OCH}_3 + \text{CN}^- \rightarrow \text{F}_5\text{SCH(C=N)CH}_2\text{OCH}_3$$

In this reaction, loss of the SF$_5$ group was observed and a tarry residue was obtained. The $\beta$-bromo-$\beta$-SF$_5$-ethylmethyl ether was obtained by nucleophilic addition of CH$_3$O$^-$ to F$_5$SCBr=CH$_2$.

$$\text{F}_5\text{SCBr=CH}_2 + \text{CH}_3\text{O}^- \rightleftharpoons \text{CH}_3\text{OH} \rightarrow \text{F}_5\text{SCHBrCH}_2\text{OCH}_3$$

(61%)
It was initially hoped that the bromine in the olefin might be replaceable by the methoxy group, which then could have been hydrolyzed to a ketone:

\[
\text{F}_5\text{SC}=\text{CH}_2 + \text{CH}_3\text{O}^- \rightarrow \text{F}_5\text{SC}=\text{CH}_2 \rightarrow \text{F}_5\text{S}\_\text{C}=\text{C-CH}_3 \rightarrow
\]

It was later learnt that its CF$_3$ analogue loses SF$_5^-$ upon attack of anions at the carbonyl group; likely, the above ketone would have behaved similarly and this would narrow the synthetic utility of such SF$_5$-ketones severely. An example of this behavior is described in the chapter dealing with the epoxide F$_5$SCFCF$_2$O.

Carbonylation of SF$_5$CBr=CH$_2$, by analogy to CF$_3$CBr=CH$_2$, was also not possible. The reaction led to the loss of the SF$_5$-group.

\[
\text{SF}_5\text{CBr}=\text{CH}_2 + \text{CO} + \text{N(C}_2\text{H}_5\text{)}_3 + \text{PdCl}_2(\text{P}_3\text{)}_2 \ (	ext{tetrahydrofuran})
\]

\[\text{--X--->} \quad \text{F}_5\text{S}
\]

\[\quad \downarrow
\]

\[\quad \text{C}=\text{CH}_2
\]

\[\quad / \]

\[\quad \text{HOOC}
\]

This reaction leads with F$_3$CC(Br)=CH$_2$ to F$_3$CC(COOH)=CH$_2$ in good yield$^{26}$. The Tollens reaction of SF$_5$CH$_2$COOCH$_3$ was not successful,

\[
\text{F}_5\text{SCH}_2\text{COOCH}_3 + (\text{CH}_2\text{O})_n + \text{HN(C}_2\text{H}_5\text{)}_2 \rightarrow \text{F}_5\text{SC(COOCH}_3\text{)}\text{CH}_2\text{OH},
\]

neither did it lead to carbon coupling with SF$_5$CH$_2$CHO:
Although some product was formed, it could not be identified, due to its miniscule mass.

The main difficulty with the aldehyde SF₅CH₂CHO was that it was laborious to obtain it pure in sufficient quantities. Its synthesis involves dehydration (with phosphorous pentoxide) of its hydrate, F₅SCH₂CH(OH)₂, which in turn is obtained by hydrolysis of the vinyl ether F₅SCH=CHOCH₃ in aqueous hydrochloric acid³⁵ᵃ,ᵇ, or by treatment of the adducts F₅SCH₂CHClOAc with methanolic potassium hydroxide³⁶, respectively, followed by precipitation of the aldehyde with water. The intermediate vinyl ether is obtained by several procedures, either by treatment of F₅SCH₂CHCl₂ with excess sodium methoxide³⁵ᵃ, by reaction of F₅SCH=CHCl with methanolic KOH³⁷ or a suspension of sodium methoxide in xylene³⁵ᵇ, or by reaction of methanolic CH₃ONa with F₅SCH=CHBr (this work). The aldehyde reacts with water, forming the hydrate, although usually mixtures of the aldehyde and the hydrate are obtained in the above procedures:

\[
\begin{align*}
F₅SCH₂CHCl₂ + 2CH₃ONa & \rightarrow F₅SCH=CHOCH₃ \\
F₅SCH=CHCl + CH₃ONa & \rightarrow F₅CCH=CHOCH₃ + NaCl \\
F₅SCH=CHBr + CH₃ONa & \rightarrow F₅CCH=CHOCH₃ + NaBr \\
F₅CCH=CHOCH₃ + HCl (reflux) & \rightarrow F₅SCH₂CHO + \\
& F₅SCH₂CH(OH)₂ + CH₃Cl \uparrow
\end{align*}
\]
All the published procedures were found unsatisfactory. For this reason a new method of synthesis was sought.

The new synthesis is simply the acidic hydrolysis of the adduct $\text{F}_5\text{SCH}_2\text{CHClOAc}$, which is easily obtained from $\text{SF}_5\text{Cl}$ and vinyl acetate as described. The corresponding basic hydrolysis with two equivalents of $\text{KOH}$ was published earlier, but proved to be difficult to control, with ensuing low yields of the aldehyde.

Also tried was whether the basic hydrolysis would lead to the aldehyde with only one equivalent of $\text{KOH}$ in methanol, because, depending on the substituent that would be first displaced, two products were possible, which should have been hydrolyzable:

$$\text{F}_5\text{SCH}_2\text{CHClOAc} + \text{OH}^- \rightarrow \text{F}_5\text{SCH}_2\text{CHOAc} + \text{Cl}^-$$

What was found was a mixture of products, $\text{F}_5\text{SCH}_2\text{CH(OCH}_3\text{)CC(O)CH}_3$, $\text{F}_5\text{SCH}_2\text{CH(OH)}_2$ and $\text{F}_5\text{SCH}_2\text{CHO}$ (the latter two products in $\approx 45\%$ yield):

$$\text{F}_5\text{SCH}_2\text{CHClOAc} + \text{OH}^- + \text{CH}_3\text{OH} \rightarrow \text{F}_5\text{SCH}_2\text{CH(OCH}_3\text{)(OAc)}_2 \quad (24.8\%)$$

From other reactions, which showed that nucleophilic
displacement in SF₅ compounds is usually not possible, because it leads to the loss of SF₅⁻, it is inferred that this reaction is rather an elimination-addition reaction:

\[
\text{F}_5\text{SCH}_2\text{CHClOAc} \rightarrow \text{F}_5\text{S} \begin{array}{c} \text{C=O} \\ \text{H} \end{array} \text{CH}_2\text{H} \quad + \quad \text{CH}_3\text{OH} \quad \rightarrow \quad \text{F}_5\text{S} \begin{array}{c} \text{C=O} \\ \text{H} \end{array} \text{CH}_2\text{H} \begin{array}{c} \text{CH(OH)} \text{OC(O)} \text{CH}_3 \\ \text{OAc} \end{array}
\]

The formation of \(\text{F}_5\text{SCH}_2\text{CH(OH)}_2\) and \(\text{F}_5\text{SCH}_2\text{CHO}\) is then explainable on the same grounds as either the hydrolysis of \(\Delta\) (\(\rightarrow\) \(\text{F}_5\text{SCH}_2\text{CHO}\)) or the nucleophilic addition of water to \(\Delta\) followed by hydrolysis of the primary \(\text{F}_5\text{SCH}_2\text{CH(OH)}\text{OC(O)}\text{CH}_3\) (\(\rightarrow\) \(\text{F}_5\text{SCH}_2\text{CH(OH)}_2\)). A parallel reaction is observed with the compound \(\text{F}_5\text{SCH=CHBr}\) when treated with sodium methoxide in methanol. The major product originates in vinyl substitution, but a minor product is an acetal, which is formed by a nucleophilic addition reaction from the primary product:

\[
\text{F}_5\text{SCH=CHBr} + \text{CH}_3\text{O}^- \rightarrow \text{F}_5\text{SCH=CHOCH}_3 \quad \rightarrow \quad \text{F}_5\text{SCH}_2\text{CH(OCH}_3\text{)}_2
\]

The chloro vinyl derivative reacts analogously. In fact, the dimethyl acetal was first obtained by this route.²²

It was difficult to separate the vinyl ether and the acetal by distillation and the hydrolysis of these acetals (the diethyl acetal was also prepared in the assumption that hydrolysis would be faster²⁸) was extremely difficult. In an experiment the diethyl acetal (which was prepared by a method described below) was heated for about two weeks with
concentrated HCl to 100°C. The mixture turned slowly dark and a tarry precipitate appeared, but the acetal could be detected (besides the aldehyde) to the very end. When it was heated with gaseous HCl a chloro acetal was formed:

\[ \text{F}_5\text{SCH}_2\text{CH(OCH}_3\text{)}_2 + \text{exc. HCl(g)} \rightarrow \text{F}_5\text{SCH}_2\text{CHCl(OCH}_3\text{)} \]

The acidic hydrolysis of \( \text{F}_5\text{SCH}_2\text{CHClOAc} \) did not pose any problems, although the yield is only \( \approx 50\% \). This is still higher than in most published procedures. In only one case a yield of almost 90% was claimed.

\[ \text{F}_5\text{SCH}_2\text{CHCl}(\text{OCH}_3) + \text{HCl(aq)} \rightarrow \text{heat} \rightarrow \text{F}_5\text{SCH}_2\text{CHO} \]

In contrast to the other hydrolytic methods, the aldehyde was obtained and not the aldehyde hydrate, so that the dehydration step could be omitted. The only successful reaction of this aldehyde was its trimerization by acid catalysis to yield a trioxane. At 80°C, 85% phosphoric acid did not produce any change, concentrated sulfuric acid led to charring at elevated temperatures (90-105°C), but converted the aldehyde at room temperature quickly to the solid trimer, by the usual, acid-catalyzed polymerization of aldehydes:

\[ 3 \text{F}_5\text{SCH}_2\text{CHO (H}^+\text{)} \rightarrow \text{CHCH}_2\text{SF}_5 \quad 86\% \]
Initially, the trimer was found as large, very appealing, lustrous, uncolored crystals as a product of partial conversion when the aldehyde was stored in the refrigerator for several weeks at \( \approx -5^\circ C \). Conversion seems to hinge upon unknown variables, because it was usually very slow, never complete and did not proceed to the same degree for different preparations of the aldehyde. It is possible that residual acid or perhaps small amounts of the aldehyde hydrate caused the conversion in this case. The trimer was initially found when the phosphorous pentoxide that was used to dehydrate the aldehyde hydrate was dissolved in water and extracted with chloroform. This compound was then recrystallized from hexane and exhibited a sharp melting point (110.5-111.5°C), and an \(^1H\) n.m.r. spectrum consistent with a 1,3,5-trioxane structure, although the mass spectrum indicated a dimer with ionization voltages as low as 15 eV. The structure of this material was determined crystallographically, and will be discussed subsequently.

As described earlier (Part I), it was not possible to substitute the anions \( F_5SC^-\left(SO_2F\right)COR \) \((R = F, OCH\left(CH_3\right)_2)\) with carbon substituents, nor was it possible to react the ketene \( F_5SC\left(SO_2F\right) = C = O \) with \( O = C = NSO_2Cl \). This latter reaction should have led to a derivative of a malonic acid\(^{39}\):

\[
\begin{align*}
F_5S\text{C}=\text{C}=O + O=\text{C}=\text{NSO}_2\text{Cl} \rightarrow & \quad F_5S\text{C}=\text{C}=\text{O}\cdot\text{NSO}_2\text{Cl} \\
\text{(+ROH } & \rightarrow \text{F}_5\text{S}\text{C}=\text{OOR})
\end{align*}
\]
In these cases, the SO₂F group in the primary products should have been removable by heating with chloro or fluorosulfonic acid⁴⁰.

It is quite clear that a functionality in position α to the SF₅ group has to be introduced from the very beginning. To achieve this goal, several observations from different reactions were used. The adduct F₅SCH₂CHClO(0)CH₃ is formed readily, but relatively slowly at elevated temperatures from vinyl acetate and pentafluorothiochloride⁴¹:

\[
\text{F}_{5}\text{SCl} + \text{CH}_2=\text{CHO}(0)\text{CH}_3 \rightarrow \text{F}_{5}\text{SCH}_2\text{CHClO}(0)\text{CH}_3 \ (>80\%) 
\]

The analogous reaction with SF₅Br is very difficult to control; only at low temperature (<=-110°C) in an inert solvent (CCl₃F) is the reaction controllable.

\[
\text{F}_{5}\text{SBr} + \text{CH}_2=\text{CHO}(0)\text{CH}_3 \rightarrow \text{F}_{5}\text{SCH}_2\text{CHBrO}(0)\text{CH}_3 \ (\approx50\%)
\]

When the precautions of low temperature and high dilution are not observed, another product is formed which originates probably in BrF addition to the olefin; it was also not possible to run the reaction on a large scale. This product is of a very decomposable nature and could never be satisfactorily characterized, but it did not contain an SF₅ group.

On the other hand, SF₅Br adds in a controlled fashion to ethyl acrylate:
The product cannot be dehydrohalogenated, it loses instead "HSF₅" as a pseudo halogen hydride, both with N,N-dimethyl aniline⁴² and diazabicyclononane⁴³:

\[ \text{F}_5\text{SBr} + \text{CH}_2=\text{CHCOOC}_2\text{H}_5 \rightarrow \text{F}_5\text{SCH}_2\text{CHBrCOOC}_2\text{H}_5 \]

The SF₄ can be detected in the n.m.r. spectrum. The result is not surprising, although unfortunate, as the hydrogen in 1-position should be the most acidic one in the molecule. The synthesis of β-SF₅ acrylates has thus to be conducted by a more elaborate course.

A further important observation is that the adduct F₅SCH₂CHClOC(O)CH₃ reacts with alcohols, completely analogously to the compounds JCH₂CHClOC(O)CH₃ and BrCH₂CHClOC(O)CH₃, to form acetals by setting the mixture aside⁴⁵:

\[ \text{F}_5\text{SCH}_2\text{CHClOC(O)CH}_3 + \text{ROH (exc.)} \rightarrow \text{F}_5\text{SCH}_2\text{CH(OR)}_2 \ (70-80\%) \]

\[ \text{R = CH}_3, \text{C}_2\text{H}_5, \text{but not CH(CF}_3)_2 \quad + \text{ROAc} + \text{HCl} + \text{H}_2\text{O} \]

These acetals can be oxidized by percarboxylic acid to a carboxylic acid⁴⁶:

\[ \text{F}_5\text{SCH}_2\text{CH(OCH}_3)_2 + \text{m-Cl-φCOOH} \rightarrow \text{F}_5\text{SCH}_2\text{COOCH}_3 \ (60\%) \]

Esters of pentafluorosulfur acetic acid are mostly made by converting the acyl chloride, which is easily accessible
from ketene (H₂C=CH₂) and alcohols. For small scale preparations the abovementioned alternative synthesis can be used which avoids the use of the noxious ketene.

The final oxidation of the acetal with m-chloro perbenzoic acid was done without addition of sulfuric acid, as recommended by Heywood and Phillips. During the reaction a precipitate was observed, that dissolved later again. This could have been the intermediate as suggested by Heywood and Phillips, a hemiacetal m-chloro-perbenzoate, but this point was not further followed.

If one combines all the above observations one recognizes the following pattern:

1. SF₅X (X = Br,Cl) adds to vinyl acetate as well as to ethyl acrylate such that the SF₅ group is attached at the terminal carbon. In the case of ethyl acrylate this is a site of reduced charge density, and in the case of vinyl acetate of increased charge density:

\[
\begin{align*}
\text{H}_2\text{C}=\text{C}-\text{O}-\text{C}(\text{O})\text{CH}_3 & \quad \leftrightarrow \quad \text{H}_2\text{C}=\text{C}^{2+}-\text{O}^{2-}\text{C}(\text{O})\text{CH}_3, \\
\text{H}_2\text{C}=\text{C}-\text{C}-\text{OC}_2\text{H}_5 & \quad \leftrightarrow \quad \text{H}_2\text{C}=\text{C}^{+}-\text{COCH}_3
\end{align*}
\]

The addition to vinyl acetate is much more efficient than the addition to ethyl acrylate as inferred from the almost uncontrollable reaction of the former with SF₅Br and the comparatively mild reaction with the latter.

2. If the charge density is the dominating criterion for the addition then the addition of an electrophilic
radical trans to an acetyl group should preponderate in the addition trans to a carboxylic ester group. This is also consistent with the transfer coefficients of bromo and chloro alkanes with different substrates. For example, vinyl acetate reacts almost 10000 times faster with CCl₄ than methyl acrylate, which is attributed to the greater reactivity of the intermediate chain propagating radical. Bromo alkanes react even faster than chloro alkanes because of the weaker C-Hal bond. As the S-Br bond is weaker than the S-Cl bond, this should be true for SF₅Br additions, as compared to SF₅Cl additions, too. This is observed experimentally.

3. Such an addition direction should be possible with β-acetyl acrylic ester.

4. The adduct thus obtained should be convertible to an acetal and thence to an ester, or to an aldehyde-ester, as with the adduct F₅SCH₂CHClOC(O)CH₃:

\[ \text{F}_5\text{SX} + \text{H}_3\text{CC(O)OCH=CHCOOC}_2\text{H}_5^{49} \rightarrow \text{F}_5\text{SCH} \rightarrow \text{F}_5\text{SCH} \]

\[ \rightarrow \text{F}_5\text{SCH} \rightarrow \text{F}_5\text{SCH} \]

\[ \rightarrow \text{F}_5\text{SCH} \rightarrow \text{F}_5\text{SCH} \]

The aldehyde-ester should be reducible to an alcohol and thence convertible to an acrylic acid:
This acid would be equivalent to the compound $\text{H}_2\text{C}=-\text{COOH}$ that was used to synthesize a number of 5-$\text{CF}_3$-uracils.

It was found that $\text{SF}_5\text{Cl}$ was completely unreactive towards $\beta$-acetylaclrylate in $\text{CCl}_3\text{F}$; without the solvent decomposition occurred of both the olefin and the $\text{SF}_5\text{Cl}$. $\text{SF}_5\text{Br}$ decomposed in a thermal reaction in the same fashion with and without solvent, but when the reaction was done in a glass bomb in the laboratory light, or better, with the irradiation of a sunlamp, $\text{SF}_5\text{Br}$ addition could be indeed observed. No addition was observed when the glass bomb was kept in the dark. By-product formation could be decreased when the irradiation was done in an ice-bath, such that only the upper part of the bomb was exposed to light, while the lower part was in the cold-bath and the solution was stirred. The product that was obtained, a colorless high-boiling oil, was very unstable and decomposed within days at room temperature but was stable in the refrigerator. That the product is a secondary $\text{SF}_5$ compound was deduced from its $^{19}\text{F}$ n.m.r spectrum, which showed the $\text{SF}_5$ resonance at $\approx 70$ ppm ($\phi_B$) and $80$ ppm ($\phi_A$). There was a second (minor) product with $\phi_B$ at $52$ ppm. The observation is made that electronegative substituents at the $\text{SF}_5$ carbon lead to an upfield shift of the $\text{SF}_5$ resonance (e.g. with the epoxides
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F₅SCFCF₂O and F₅SCHCF₂O, as compared to their starting olefines), so that the product with φB = 52 ppm is probably F₅SCH(OAc)CHBrCOOC₂H₅. The product could be purified by distillation. The infrared spectrum is also supportive of the assumed structure of the chief product F₅SCHCOOC₂H₅(CHBrOAc). Two C=O bands were present, one of them (1783 cm⁻¹) similar to the compounds F₅SCH₂CH(Cl,Br)OAc (1778 cm⁻¹), and the other at 1754 cm⁻¹, similar to F₅SCH₂COOCH₃ (1757 cm⁻¹). For subsequent reactions the impure material was used. It was found that the material reacted in a very intricate manner with alcohols. When ethanol was added to it in excess the reaction, as followed by ¹⁹F n.m.r. spectroscopy, displayed the disappearance and appearance of several SF₅ compounds, with the formation of two major product after several days. The infrared spectrum showed the disappearance of the band at 1783 cm⁻¹, corresponding to the loss of the acetyl group. Distillation of the material resulted in decomposition and the formation of at least four new SF₅ compounds. Attempts to separate them by preparative gas chromatography were unsuccessful.

Attempts to induce reduction of the SF₅Br adduct with NaBH₄ were unsuccessful, treatment with aqueous HCl led to decomposition (this should have resulted in the formation of an aldehyde, analogously to F₅SCH₂CHClOAc), but treatment with an excess of gaseous HCl led to mainly one product, but it could again not be separated.
Although with this reaction one important problem was solved, namely the synthesis of a secondary SF$_5$-compound, the subsequent screening reactions gave a disappointing result. This behavior is not understood at this point. It cannot be ruled out that the probably severe sterical crowding prevents effective acetal formation.

This work was stopped at this point.
CHAPTER IV

THE X-RAY STRUCTURE OF $(\text{SF}_5\text{CH}_2\text{CHO})_3$

The mass spectrum of this aldehyde oligomer did not show a molecular ion signal, even at ionization potentials as low as 15 eV. The highest mass was seen at $m/e = 341$ (($\text{SF}_5\text{CH}_2\text{CHO})_2\text{H}^+$), which could have originated in a dimer. It was thus of interest to determine the actual structure. To determine the molecular weight would have answered the question of the structure (dimer or trimer) also, but it seemed important to have more structural information in order to build up a reference file for SF$_5$ compounds, as this is a developing field of chemistry, with little information available from the literature.

The crystal structure of $(\text{SF}_5\text{CH}_2\text{CHO})_3$ consists of trimeric units with threefold symmetry in which the trioxane ring is situated in a chair-type conformation. The SF$_5$CH$_2$ groups lie in an equatorial position. A similar chair-type conformation has been found for $\alpha$- and $\beta$-parabutylchloral, $(\text{CH}_3\text{CHClCCl}_2\text{CHO})_3$, in which the 1,1,2-trichloropropyl groups are located in a cis,cis arrangement; all CH$_3$CHClCCl$_2$-groups are in the equatorial positions$^{50}$.

In $\alpha$-(CH$_3$CHClCCl$_2$CHO)$_3$, the C-O bond lengths vary from 1.396(7) Å to 1.425(8) Å and the C-O-C and O-C-O bond
angles for the ring are 109.9(5)° to 110.1(5)° and 109.6(5)° to 110.5(5)°, respectively. In (SF₅CH₂CHO)₃ a larger alternation of C-O bond-lengths of 1.435 Å and 1.387 Å is found with C-O-C and O-C-O bond angles of 109.6° and 110.4°, respectively. This alternation must be due to the steric influence of the CH₂SF₅ groups which are rotated about the C-C bonds so as to lie closer to the oxygen atom involved in the longer C-O bond.

There is no evidence for isomerism as is reported for other trioxane compounds, e.g., in parachloral an α-form and a β-form exist⁵¹.

The α-form is derived from the cis-trans isomerism of the CCl₃ substituents. The α-trans-isomer assumes a tub conformation, apparently stabilized by an intramolecular hydrogen bond. The β-form, or all-cis isomer, has a twisted chair conformation.

**Figure 7.** X-ray stereographic view of (F₅SCH₂CHO)₃
TABLE VII

NON-HYDROGEN BOND LENGTHS [pm] AND ANGLES [*] FOR
(SF$_5$CH$_2$CHO)$_3$. STANDARD DEVIATIONS IN PARENTHESES

<table>
<thead>
<tr>
<th>Bond</th>
<th>Length (pm)</th>
<th>Bond</th>
<th>Length (pm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S-C(1)</td>
<td>179.8(6)</td>
<td>S-F5)</td>
<td>157.4(6)</td>
</tr>
<tr>
<td>S-F(1)</td>
<td>158.1(3)</td>
<td>O-C(2)</td>
<td>143.5(4)</td>
</tr>
<tr>
<td>S-F(2)</td>
<td>156.2(4)</td>
<td>C(1)-C(2)</td>
<td>151.0(6)</td>
</tr>
<tr>
<td>S-F(3)</td>
<td>155.2(4)</td>
<td>C(2)-O(A)</td>
<td>138.7(6)</td>
</tr>
<tr>
<td>S-F(4)</td>
<td>157.3(3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C(1)-S-F(1)</td>
<td>89.6(2)</td>
<td>C(1)-S-F(5)</td>
<td>177.3(2)</td>
</tr>
<tr>
<td>C(1)-S-F(2)</td>
<td>92.0(3)</td>
<td>F(1)-S-F(5)</td>
<td>87.8(2)</td>
</tr>
<tr>
<td>F(1)-S-F(2)</td>
<td>90.4(2)</td>
<td>F(2)-S-F(5)</td>
<td>88.5(3)</td>
</tr>
<tr>
<td>C(1)-S-F(3)</td>
<td>92.3(3)</td>
<td>F(3)-S-F(5)</td>
<td>87.2(3)</td>
</tr>
<tr>
<td>F(1)-S-F(3)</td>
<td>90.3(2)</td>
<td>F(4)-S-F(5)</td>
<td>87.4(2)</td>
</tr>
<tr>
<td>F(2)-S-F(3)</td>
<td>175.7(3)</td>
<td>C(2)-O-C(2A)</td>
<td>109.6(4)</td>
</tr>
<tr>
<td>C(1)-S-F(4)</td>
<td>95.2(2)</td>
<td>S-C(1)-C(2)</td>
<td>118.1(3)</td>
</tr>
<tr>
<td>F(1)-S-F(4)</td>
<td>175.3(3)</td>
<td>O-C(2)-C(1)</td>
<td>108.8(4)</td>
</tr>
<tr>
<td>F(2)-S-F(4)</td>
<td>89.4(2)</td>
<td>O-C(2)-O(A)</td>
<td>110.4(4)</td>
</tr>
<tr>
<td>F(3)-S-F(4)</td>
<td>89.6(2)</td>
<td>C(1)-C(2)-O(A)</td>
<td>105.8(3)</td>
</tr>
</tbody>
</table>
Neither pentafluorosulfur malonic ester nor \( \alpha \)- or \( \beta \)-SF\(_5\) acrylic ester were obtained. It was, however, possible to devise a method for the synthesis of SF\(_5\)-acetaldehyde acetals (from 1-chloro-1-acetoxy-2-pentafluorothio ethane) and their conversion to esters, and the synthesis of a secondary SF\(_5\) compound, F\(_5\)SCH(COOC\(_2\)H\(_5\))CHBrOCOCH\(_3\), which should have been convertible, in analogy to the abovementioned procedure, to an acetal, F\(_5\)SCH(COOC\(_2\)H\(_5\))CH(O\(_2\)H\(_5\))\(_2\) and thence to an ester, F\(_5\)SCH(COOC\(_2\)H\(_5\))\(_2\). The chemical behavior of this system is not understood. A new method for the synthesis of pentafluorothio acetaldehyde was also found.
PART IV

EXPERIMENTAL PART
CHAPTER I

GENERAL METHODS

Volatile materials were handled in conventional Pyrex-glass vacuum lines, equipped with either mercury manometers or Heise-Bourdon tube gauges and a Televac vacuum gauge. $^1$H and $^{19}$F n.m.r. spectra were usually run at 90 ($^1$H) and 84.67 ($^{19}$F) MHz on a Varian EM-390 spectrometer, otherwise they were run on a General Electric QE-300 spectrometer at 300 ($^1$H) and 75 ($^{13}$C) MHz, or on a Bruker AMX-400 spectrometer ($^{19}$F at 376 MHz, $^{13}$C at 100.6 MHz, $^1$H at 400.6 MHz). The following abbreviations are used to indicate splitting patterns: s = singlet; d = doublet; t = triplet; q = quartet; p = pentet, quintet; m = multiplet. Infrared spectra were obtained from neat samples between KBr or NaCl plates on a Nicolet DX-20 spectrometer. Band intensities are indicated by the following: vs = very strong; s = strong; m = medium; w = weak; vw = very weak, br = broad, sh = shoulder. The band positions are always given in 1/cm. The mass spectra were obtained on a VG 7070 mass spectrometer under the conditions indicated with each compound. In the mass spectra of bromine or chlorine compounds the masses of fragments of only one isotope are listed, i.e., for $^{35}$Cl and $^{79}$Br. The mode of ionization is
indicated by the following short-hands: e.i. = electron impact; c.i. = chemical ionization; n.i. = negative ion. Acetonitrile was distilled from phosphorous pentoxide; other solvents were used without further treatment. Chemicals that were used in standard reactions were obtained from Ozark-Mahoning-Pennwalt (BrF₃, SF₄), Aldrich or Peninsula Chemical Research Corporation (PCR) (CsF, C₂HF₃, CH₂CF₂), NaF (Baker), Mallinckrodt (CH₃OH, C₂H₅OH). The sources of other chemicals will be named in the respective preparations where they were used. F₅SCHCF₂O₃SO₂ was prepared as described.

Elemental analyses were carried out by Mikroanalytisches Laboratorium Beller, Göttingen, Federal Republic of Germany.
SF₅Br from BrF₃ + Br₂ + SF₄:

In a typical experiment, a 150 ml stainless steel bomb was charged with CsF (7.53 g, 49.2 mmol) and dried (15 hours, 100-110°C, high vacuum). Bromine trifluoride (29.72 g, 217 mmol) was added by vacuum transfer, then bromine (34.4 g, 215 mmol), and lastly SF₄ (48.71 g, 451 mmol equals 70% of theory, in two portions, 1. 32.42 g, then 16.28g); the bomb was allowed to attain room temperature in an automatic shaker between the two additions. Shaking was continued for 289 h at room temperature, and 74.5 g of crude SF₅Br were transferred out from the bomb (at -78°C) during 4 hours (56.8%). Major by-products were SF₆ and unreacted SF₄. This crude material was used without further purification and reacted with acetylene to the adduct F₅SCH=CHBr (57% yield after distillation). Similar reactions were conducted with other olefins, and yields ranging from 39 - 69% were realized:

<table>
<thead>
<tr>
<th>Olefin</th>
<th>Adduct</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHF=CF₂</td>
<td>F₅SCHFCF₂FBr</td>
<td>38.6-57.8</td>
</tr>
<tr>
<td>CH₂=CF₂</td>
<td>F₅SCH₂CF₂Br</td>
<td>69.5</td>
</tr>
<tr>
<td>HC=CH</td>
<td>F₅SCH=CHBr</td>
<td>45.5-61.6</td>
</tr>
</tbody>
</table>
Although only small amounts of bromine transfer at -78°C, its olefin adducts might interfere later. However, it was easily removed by shaking the crude SF₅Br with mercury (10-15 g for 70 g of crude SF₅Br; the mercury may be used several times). It was also found that using more CsF (up to 50 mol% with respect to BrF₃) did not raise the yield of SF₅Br. Shaking was generally not extended beyond 2 weeks. When a vessel had been used several times, yields dropped. It was also found that there were still substantial quantities of SF₅Br left, even when a transfer had lasted for several hours at -78°C. Up to 20 g of SF₅Br that contained almost no SF₄ and SF₆ could be collected from bombs with a similar charge as described above, by prolonged transfer (up to 10 h) at -78°C, after thawing and recooling. It was later found that quantities of up to 56 g of SF₄ could be transferred at once into the 150 ml bombs containing the BrF₃+Br₂ mixture, without any incident. Larger quantities were not tried.

**¹³C n.m.r. spectrum of F₅SCH₂CH₂Br (100.6 MHz, Si(CH₃)₄):**

A CDCl₃ solution, ≈ 50% in the SF₅ compound was used. The spectrum is ¹H decoupled. \( F^A F_4^B S C_a H F_a C_\beta F_\beta^1 F_\beta^2 B r \): \( \delta_\alpha = 108.15 \) ppm, \( \delta_\beta = 114.87 \) ppm, \( \delta_{\beta^1} = 23.3 \) Hz; \( J_{CaF_\alpha} = 249.0 \) Hz; \( J_{CaF^B} = 23.3 \) Hz; \( J_{CaF^1_{\beta^2}} = 36.0 \) Hz; \( J_{CaF^2_{\beta^1}} = 27.0 \) Hz; \( J_{C_\beta F_\beta^1_{\beta^2}} = 15.4 \) Hz; \( J_{C_\beta F_\beta^2_{\beta^1}} = 307.8 \) Hz;
\[ J_{C\beta \alpha} = 30.2 \text{ Hz.} \]

**F_{5}\text{SCH} = \text{CF}_2(52):**

To an aqueous potassium hydroxide solution (81.5 g, 42.5% by weight KOH) in a 100 mL Pyrex-glass round bottomed flask, equipped with a magnetic stirring bar, F_{5}\text{SCH}2\text{CF}_2\text{Br} (33.70 g, 124.4 mmol) was added at room temperature. A 20-cm reflux condenser that was connected to a cold trap (-78°C) was then attached to the reaction vessel. The mixture was slowly heated with vigorous stirring to a temperature of 100°C, and warm water (40°C) was passed through the condenser; after the disappearance of the bottom layer in the reaction flask (30 min), heating was maintained for another 3 hours. The product was collected in the cold trap (22.90 g, 120.5 mmol, 96.9% yield). A gas chromatogram (Carbowax, 20% on Chromosorb W, 100°C) showed only one product to be present; the i.r. spectrum agreed with the literature values, except that for the latter several other lines (weak) are reported, indicating that the present product is purer.

**F_{5}\text{SCH}(\text{SO}_2\text{F})\text{COOCH(CH}_3\text{)}_2(21a):**

To a 100 ml Pyrex-glass reaction vessel, equipped with a Teflon stir bar and a Kontes Teflon valve, and containing dry sodium fluoride (30.08 g, 716.2 mmols), isopropanol (10.13 g, 168.8 mmol) was added by vacuum condensation. The
sultone $\text{F}_5\text{SCHCF}_2\text{OSO}_2$ was then added in three portions (12.37g, 45.8 mmol; 8.70g, 32.20 mmol; 15.91g, 58.9 mmol) at -196 °C, with the vessel being warmed to room temperature after each addition and stirred for a few minutes. For each addition slight warming was noticed when room temperature was reached. After addition was complete, the mixture was stirred at 52°C for 15 h and at 68°C for 0.5 h. The crude product was condensed (41.29g) and distillation gave 31.12g (100.5 mmol) of a colorless oily liquid with an unpleasant odor; yield = 73.3%; b.p. 92-94°C (26-28) torr.

Infrared spectrum (neat sample, KBr, cm$^{-1}$): 2995, m; 2952, w,sh; 1761, s; 1443, s; 1395, vw; 1382, w; 1352, vw; 1296, m; 1276, m; 1221, w; 1168, vw; 1102, m-s; 981, vw; 875, vs,sh at 900; 837, w-m; 819, w-m; 801, w-m; 781, w; 769, w; 733, vw; 688, vw; 650, vw; 612, w; 572, s; 491, vw; 458, vw.

$^{19}$F n.m.r. spectrum (neat sample, external CCl$_3$F): multiplet, centered at $\delta = 70.9$ ppm, area = 5.0 ($\text{SF}_5$); multiplet at $\delta = 58.8$ ppm, area = 1.0 ($\text{SO}_2\text{F}$).

$^1$H n.m.r. spectrum (neat sample, Si(CH$_3$)$_2$ ext.): multiplet, $\delta = 6.4$ ppm, area = 1.0 ($\text{F}_5\text{SC-H}$); septet, $\delta = 5.4$ ppm, area = 1.1, (O-CH(CH$_3$)$_2$); doublet, $\delta = 1.5$ ppm, area = 6.2, (O-CH(CH$_3$)$_2$), $J_{\text{H-H}} = 6.22$ Hz.

Mass spectrum (c.i., m/e, species, %): 311, (M+H)$^+$, 0.32; 269, (M-3H-2F)$^+$, 4; 205, (M+H-(CH$_3$)$_2$CHCOO-F)$^+$, 7.5; 143, $\text{C}_2\text{HF}_2\text{OS}_2^+$, 14.8; 127, $\text{SF}_5^+$, $\text{FSCSO}_2^+$, 20.5; 125, $\text{C}_2\text{H}_2\text{FO}_3\text{S}^+$, 8.4; 99, $\text{C}_4\text{H}_3\text{SO}^+$, 9.2; 91, $\text{C}_2\text{FOS}^+$, 4.2; 89, $\text{SF}_3^+$,
100; 79, FSCO+, 4.1; 70, SF2+, 10.8; 67, FSO+, 24.6; 65, 
CH$_2$FS+, 19; 59, C$_3$H$_7$O+, 55.8; 58, C$_3$H$_6$O+, 5.5; 55, C$_3$H$_3$O+, 
4.8.

**Anal.** Calcd for C$_5$H$_8$F$_6$O$_4$S$_2$: C, 19.36; H, 2.60; F, 36.8; S, 20.67. Found: C, 19.27; H, 2.55; F, 37.1; S, 20.82.

$\text{F}_5\text{SC(SO}_2\text{F)=C=O (21), Method 1:}$

In a representative experiment, $\text{F}_5\text{SCH(SO}_2\text{F)COOCH(CH}_3\text{)}_2$
(2.65 g, 8.55 mmol) was mixed with powdered $\text{P}_4\text{O}_{10}$ (Baker, 8.85 g, 31.1 mmol) in a 100 mL Pyrex-glass round bottomed flask. The flask was connected to a short vacuum-transfer line, with a 10 mL receiving flask that was cooled to -196°C. The pressure was lowered to 500 torr and the mixture was heated to 100°C (10 min). The reaction flask was cooled to -196°C, evacuated, and reheated (100°C). In 30 min 1.60 g (6.4 mmol) of crude ketene was collected. Prolonged heating led to transfer of unreacted ester (0.34 g, 1.1 mmol). Yield: 85.9% (with respect to recovered ester). In similar experiments, the yields varied from 49 to 89 percent. In this fashion, 4.92 g of crude ketene 21 was obtained from 9.35 g (30.16 mmol) of ester. Distillation (125-128 torr, (54-56)°C) gave 4.09 g (16.36 mmol) of pure ketene 21 in 54.2% overall yield.

$^{19}\text{F n.m.r. spectrum (CDCl}_3, \text{CCl}_3\text{F ext., AB}_4\text{X spectrum): }$$\varphi_A = 73.1$ ppm, (nine lines), area = 1.0; $\varphi_B = 86.7$ ppm, (d-m), area = 3.9; $\varphi_X = 76.3$ ppm, (SO$_2$F), pentet,
area = 1.0; J_{AB} = 155 \text{ Hz}, J_{BX} = 6.8 \text{ Hz}.

^{13}\text{C n.m.r. spectrum (100.6 MHz, CDCl}_3, \text{ Si(CH}_3)_4\text{): } \delta_1 = 86.28 \text{ ppm, d-p, (F}_5\text{SC)}; \delta_2 = 174.81 \text{ ppm, } \approx \text{ pentet (ill-resolved, CO)}; J_{1X} = 39.70 \text{ Hz (X = SO}_2\text{F)}; J_{2B} = 32.46 \text{ Hz}; J_{2B} \approx 3.3 \text{ Hz.}

Infrared spectrum (neat sample, KBr, cm}^{-1}: 2193(vs), 1445(vs), 1305(m), 1214(vs), 1025(vs), 875(vs), 845(s,sh), 801(s), 689(m), 635(m), 618(m), 590(m), 520(w), 470(w), 432(vw).

Mass spectrum (electron impact, m/e, species, %): 250, M\textsuperscript+ (32S), 16.4; 231, (M-F)\textsuperscript+, 3.2; 129, (M-SO}_2\text{F}_3\text{)}\textsuperscript+, 5.3; 127, SF\textsubscript{5}\textsuperscript{+}, FSCSO\textsubscript{2}\textsuperscript{+}, 94.7; 91, C\textsubscript{2}FOS\textsuperscript{+}, 6.4; 89, SF\textsubscript{3}\textsuperscript{+}, 100; 72, C\textsubscript{2}OS\textsuperscript{+}, 1.9; 70, SF\textsubscript{2}\textsuperscript{+}, 11.6; 67, FSO\textsuperscript{+}, 25.2; 63, CFS\textsuperscript{+}, 7.1.

Anal. calcd: for C\textsubscript{2}F\textsubscript{6}O\textsubscript{3}S\textsubscript{2}: C, 9.60; F, 45.6; S, 25.63. Found: C, 9.86; F, 45.4; S, 25.82.

**NaF catalyzed rearrangement of F\textsubscript{5}SC\textsubscript{2}CF\textsubscript{2}OSO\textsubscript{2}:**

In a dry 100mL Pyrex-glass flask, equipped with a Kontes Teflon valve and a stirrrring bar, NaF (5.30 g, 126.2 mmol) was dried and the sultone F\textsubscript{4}SCHCF\textsubscript{2}OSO\textsubscript{2} (14.99g, 44.4 mmol) was added. The mixture was stirred at (83-85)°C for 4 days. Vacuum transfer afforded 14.15 g of a mixture which consisted (^{19}\text{F n.m.r.) solely of a 14:86 (}%) mixture of ketene 21, F\textsubscript{5}SC(SO\textsubscript{2}F)CO, and the acyl fluoride F\textsubscript{5}SCH(SO\textsubscript{2}F)COF 22 (yield \approx 95%). It was later found that using an excess of finely ground NaF speeds up the
rearrangement considerably. In this case much of the contents of the flask solidify, and it is possible to transfer out from the heated reaction vessel mixtures with much higher ketene contents (>80%), but 21 could never be obtained pure by this method.

\[
F_5SC(SO_2CO) (21), \text{Method 2:}
\]

To 5.00 g of the 14:86 (%) mixture of ketene 21 and acyl fluoride 22, as obtained from the NaF-catalyzed rearrangement of the sultone \( F_5SC(SO_2CF_2O) \) (1), in a 25 ml Pyrex-glass round bottomed flask, equipped with a stirring bar, the base \( BF_3\cdot N(C_2H_5)_3 \) (18, 3.17 g, 18.8 mmol) was added. The flask was attached to a vacuum-transfer line, with a 10 mL receiving flask that was cooled to -196°C. The pressure in the line was lowered to 500 torr, and the mixture was heated to 100°C over a 25-min period. At \( \approx 85°C \) a crystalline precipitate was observed in the homogeneous melt, and which disappeared at \( \approx 90°C \), leaving an upper brown and a lower colorless liquid layer. Heating was continued at 100°C (25 min). The product (4.45 g) was transferred under high vacuum (30 min) to the 10 mL receiving flask cooled to -196°C. The \(^{19}F\) n.m.r. and i.r. spectra showed that only ketene 21 was present, although the product was slightly fuming. Yield (based on the 14:86 mixture): 95%.
**F₅SC(SO₂F)CO in a sealed tube:**

Ketene **21** (0.49g, 1.96 mmols) was heated in an evacuated dry 50 mL Carius tube to 220°C for 4h. The liquid inside was slightly discolored but 0.48 g of unchanged ketene was recovered.

**F₅SC(SO₂F)CO and iso-propanol:**

To 118 mg of iso-propanol (1.94 mmol) in a 1 ml dry reaction vessel ketene **21** (497 mg, 1.99 mmol) was added dropwise. Initially, the reaction was vigorous, but subsided after addition of approximately 1/2 of the ketene. After addition was complete, the reaction vessel was kept at room temperature (1/2 h) and 568 mg of product was collected (94.2%). The infrared spectrum and ^19^F n.m.r. spectra agreed with the spectra of the ester F₅SCH(SO₂F)COOCH(CH₃)₂, as described above. Gas chromatography (Carbowax 20%, 6ft, on Chromosorb W, 110°C) of the product showed the product to be essentially pure (>95%).

**F₅SC(SO₂F)CO and 2,2,2-Trifluoroethanol:**

To 0.50 g (2.0 mmol) of ketene **21** in a dry 30ml Pyrex-glass reaction vessel, equipped with a Kontes Teflon valve, CF₃CH₂OH (0.24g, 2.4 mmol) was condensed at -196°C. The mixture was allowed to attain room temperature slowly and heated to 83°C for 1 hour. The liquid product was transferred out, washed with water (2x0.5 ml), the lower
layer was taken up with a pipet, transferred to a 5ml flask and dried (Na$_2$SO$_4$). The infrared spectrum of the product (0.59 g, 1.7 mmol) was essentially identical with the published spectrum of F$_5$SCH(SO$_2$F)COOCH$_2$CF$_3$. The $^{19}$F n.m.r. spectrum revealed that the product was chiefly the trifluoroethyl ester with some F$_5$SCH$_2$SO$_2$F as a minor impurity. Yield 82% (n.m.r. spectrum).

E$_5$SC(SO$_2$F)CO and H$_2$O:

Into a 100 ml reaction vessel, equipped with a Kontes Teflon valve and containing water (40.5 mg, 2.25 mmol) ketene 21 (485.3 mg 1.94 mmol) was condensed. The mixture was allowed to attain room temperature and to remain there for 2 h. It was then cooled to -78°C and the infrared spectrum of the off-gas showed CO$_2$ to be present with trace impurities. The amount of CO$_2$ formed was 85.4 mg (1.94 mmol), theor. (1.94 mmol, 85.4 mg). The residue was transferred at elevated temperature and dried (Na$_2$SO$_4$). A total of 369.0 mg of product was collected. A gas chromatogram (Carbowax, 20%, 6 ft, on Chromosorb W, 100 °C) showed only one product; its infrared and $^{19}$F n.m.r. spectra were identical with the published spectra of F$_5$SCH$_2$SO$_2$F. Yield: 84.9%.

E$_5$SC(SO$_2$F)CO and (C$_2$H$_5$)$_2$NH (27):

Ketene 21 (0.49g, 1.96 mmol) was condensed into a 50
ml Pyrex-glass reaction vessel, equipped with a Kontes Teflon valve and containing 2 ml of dry CH₂Cl₂ at -196°C. The vessel was warmed above the melting point of the CH₂Cl₂, shaken, and cooled again to -196°C, and diethylamine (0.142 g, 1.94 mmol) was added by vacuum-transfet. Upon being warmed to the melting temperature the solution was shaken, frozen (-196°C) and allowed to attain room temperature slowly. During warming to room temperature the solution turned brownish. At room temperature it was decanted, concentrated and recrystallized from CH₂Cl₂, leaving 302 mg (0.93 mmol of F₅SCH(SO₂F(C)N(C₂H₅)₂, m.p. 108-110°C. Yield: 47.7%.

¹⁹F n.m.r. spectrum (CDCl₃, CCl₃F, ext.): multiplet, δ = 72.2 ppm, area = 4.8 (SF₅); multiplet, δ = 62.3 ppm, area = 1.0 (SO₂F).

¹H n.m.r. (300 MHz, CDCl₃, Si(CH₃)₄): triplet, δ = 1.19 ppm, J_H-H = 7.1 Hz, area = 3.0 (CH₃); triplet, δ = 1.32, J_H-H = 7.1 Hz, area = 3.0 (CH₃); multiplet, centered at 3.5 ppm, area = 4.3 (CH₂); pentet, δ = 6.00 ppm, J_H-F = 4.0 Hz, area = 0.9 (CH).

I.r. spectrum (neat sample, squeezed on KBr plate), cm⁻¹: 2990(w), 1661(vs), 1460(vw), 1432(s), 1390(w), 1363(w), 1320(w), 1278(w), 1222(m), 1212(m), 1129(w), 1096(w), 1083(vw), 953(w), 899(m), 860(vs), 820(m), 799(m), 740(w), 647(w), 562(m), 488(w), 458(w).

Mass spectrum (70 eV, mass, species, %): 323, M⁺, 1.6;
308, (M-CH$_3$)$^+$, 12.1; 280, C$_6$H$_4$F$_6$NO$_3$S$_2$$^+$, 2.0; 231, C$_2$F$_5$O$_3$S$_2$$^+$, 4.6; 196, (M-SF$_5$)$^+$, 14.9; 176, (M-HF-SF$_5$)$^+$, 5.8; 127, (SF$_5$, CFO$_2$S$_2$)$^+$, 7.4; 125, C$_2$H$_2$FO$_3$S$^+$, 2.9; 112, (C$_2$H$_5$)$_2$NCOC$^+$, 2.9; 105, C$_2$HO$_3$S$^+$, 2.7; 100, C$_5$H$_{10}$NO$^+$, 5.8; 89, SF$_3$$^+$, 14.8; 73, C$_2$HOS$^+$, 2.0; 72, (C$_2$H$_5$)$_2$N$^+$, C$_2$OS$^+$, 42.4; 71, C$_4$H$_9$N$^+$, 3.4; 70, C$_4$H$_8$N$^+$, SF$_2$$^+$, 4.4; 67, FSO$^+$, 5.5; 58, C$_2$H$_4$NO$^+$, SCN$^+$ (rearr.), 100; 56, C$_3$H$_6$N$^+$, C$_2$H$_2$NO$^+$,C$_2$S$^+$, 21.5; 54, C$_2$NO$^+$, 2.0.

**Anal. calcd** for C$_6$H$_{11}$F$_6$NO$_3$S$_2$: C, 22.29; H, 3.43; F, 35.3; N, 4.33; S, 19.83. Found: C, 22.37; H, 3.54; F, 35.2; N, 4.26; S, 19.91.

F$_5$SC(SO$_2$F)=C=O and HSi(CH$_3$)$_3$ (39a):

The reaction was carried out in dry glassware under strict exclusion of moisture.

Into a 50 ml pear shaped Pyrex-glass vessel, equipped with a Kontes Teflon valve, 1.1536 g of 21 were added by vacuum-transfer. The stoichiometric amount of SiH(CH$_3$)$_3$ (Petrarch) was added by vacuum transfer (190 torr, 0.455 l, 295 °C), and the vessel was allowed to attain 0°C slowly by allowing it to warm up in a pre-cooled Dewar container. A white, somewhat moist-looking solid was obtained. Another $\frac{1}{2}$ equivalent of SiH(CH$_3$)$_3$ was added (total = 1.5 equiv.). No more reaction was observed. The vessel was immersed in an ice-bath and volatile materials were pumped off through a -196°C cold-trap (45 minutes). Residue = 1.3445 g (theor. =
1.4951 g). The material was transferred to a sublimation apparatus in a dry-box and sublimed in an oil bath (48°C). A white crystalline solid collected quickly on the cold-finger and the sublimation was over in ¼ hour. Yield = 0.9290 g (62.1 %), m.p. (sealed tube) 44-48°C. The compound seems to be perfectly stable when kept in a vacuum-sealed glass tube, but turns dark slowly in a screwcap vial in the dry-box.

$^1$H n.m.r. spectrum (CDCl$_3$, Si(CH$_3$)$_4$ ext.): $\delta_1$ = 0.43 ppm, s (evidence of shoulders), area = 9.00 (Si(CH$_3$)$_3$); $\delta_2$ = 8.10 ppm, d-p (poorly resolved), area = 0.95; coupling to AB$_4$X ($^1$F): $J_{2B}$ $\approx$ 4.2 Hz; $J_{2X}$ $\approx$ 1.7 Hz.

$^{19}$F n.m.r. spectrum (CDCl$_3$, CCl$_3$F ext.): $\phi_{SF_5}$ $\approx$ 75 ppm, m, int. = 5.00 (the appearance of the spectrum is still $\approx$ AB$_4$, but $J_{AB}$ = 160.7 Hz, well above the usual range, indicating that the spectrum cannot be interpreted by simple means); $\phi_{SO_2F}$ = 68.6 ppm, partially resolved pentet, intensity = 1.09, $J_{SF_4-SO_2F}$ = 8.3 Hz.

$^{13}$C n.m.r. spectrum (100.6 MHz, CDCl$_3$, Si(CH$_3$)$_3$): $\delta_1$ = 1.83 ppm, quartet (CH$_3$), $J_{1H}$ = 116.4 Hz; $\delta_2$ = 128.95 ppm, m; $\delta_3$ = 159.24 ppm, d, $J_{3H}$ = 179.0 Hz.

I.r. spectrum (neat sample, KBr, cm$^{-1}$): 3028, w; 2976, w; 2919, vw; 1600, m-s,br; 1415, s; 1363, m; 1310, m; 1257, m-s; 1209, s; 1044, m-s; 980, w; 874, vs,sh; 861, vs; 822, vs; 772, vs; 721, s; 682, s; 634, m-s; 609, s; 579, vs; 494, w-m; 460, m.
Mass spectrum (e.i., 70 e.V., mass, species, > 1 %, only $^{28}\text{Si}$ listed): 309, (M-CH$_3$)$^+$, 21.5; 261, (M+H-3CH$_3$)$^+$, 7.0; 213, (M-Si(CH$_3$)$_3$-2F)$^+$, 2.2; 187, (F$_5$SCSO)$^+$, 1.3; 185, CHF$_4$O$_2$S$_2$$^+$, 11.9; 165, CF$_3$O$_2$S$_2$$^+$, 4.4; 126, C$_4$H$_2$OSSi$^+$, 1.1; 125, C$_5$H$_5$SSi$^+$, 13.3; 124, C$_5$H$_3$SSi$^+$, C$_2$HFO$_3$S$^+$, 3.7; 105, C$_2$HO$_3$S$^+$, 4.6; 103, CHSF$_3$$^+$, 1.0; 97, C$_2$Si(CH$_3$)$_3$$^+$, CH$_2$SO$_2$F$^+$, 9.6; 92, C$_4$OSi$^+$, COS$_2$$^+$, 2.1, 91, FSCCO$^+$, 1.8; 89, SF$_3$$^+$, 24.2; 81, C$_3$HOSi$^+$, 2.6; 79,(3.6), 78,(6.5), 77,(100.0), C$_4$HSi$^+$, 76, C$_4$Si$^+$, 1.6; 75,(2.9), 74,(4.8), 73,(56.3), Si(CH$_3$)$_3$$^+$; 72, C$_3$H$_8$Si$^+$, 2.2; 70, SF$_2$$^+$, 2.4; 67, SOF$^+$, 3.4; 64, SO$_2$$^+$, 4.3; 63, CF$_2$$^+$, 7.1; 62, CH$_2$OS$^+$, 1.5; 57, C$_2$H$_5$Si$^+$, 1.4.

Anal. calcd for C$_5$H$_{10}$F$_6$O$_3$S$_2$Si: C, 18.52; H, 3.11; F, 35.2; S, 19.77. Found: C, 18.42; H, 3.00; F, 35.5; S, 19.92.

F$_5$SC(SO$_2$F)=C=O and Acetone (30):

Dry acetone (1.05 g, 18.10 mmol) and 0.67 g of 21 (2.68 mmol) were mixed at room temperature in a 5 ml flask; the flask was stoppered and kept at room temperature overnight (17 h). The solution was yellow the next morning and all ketene was consumed. Water was added (3 ml), the mixture was swirled and the water pipetted off. This was repeated twice with 2 ml H$_2$O each. After drying (Na$_2$SO$_4$) 0.73 g of a light yellow product was obtained. This material was subjected to preparative gas chromatography (20 % SE-30, 45/60, 3m, 95 °C). Three bands were eluted of which the
first one ($R_f = 13$ minutes) was acetone, the second one ($R_f = 17$ minutes) evaporated rapidly upon warming up, leaving a very small amount of an oily, colorless residue, and the third band ($R_f = 23$ minutes) was almost pure $\text{F}_5\text{SC}(\text{SO}_2\text{F})$ (this was confirmed with the infrared spectrum). The residue from fraction 2 was just enough to obtain the $^1\text{H}$ and $^{19}\text{F}$ n.m.r. and infrared spectra.

$^1\text{H}$ n.m.r. spectrum (400 MHz, CDCl$_3$, internal Si(CH$_3$)$_4$): $\delta_1 = 2.00$ ppm, s, (3 H); $\delta_2 = 4.89$ ppm, s, (1.86 H); $\delta_3 = 5.77$ ppm, m, (0.85 H).

$^{19}\text{F}$ n.m.r. (CDCl$_3$, external CC$_3$F): $\phi = 73.0$ ppm, m, area = 5.00 (SF$_5$); $\phi = 60.9$ ppm, m, area = 0.94 (SO$_2$F).

I.r. spectrum (neat sample, on KBr, cm$^{-1}$): 2979, w; 1776, s; 1683 (C=O), w-m; 1446, s; 1382, vw; 1301, m; 1267, m; 1226, m-s; 1212, m-s; 1193, m-2; 1148, m-s; 981, vw; 874, vs; 833, m; 816, m,sh; 796, m; 773, w-m; 747, w; 688, w; 647, w; 610, w-m; 569, s; 487, w-m.

$\text{F}_5\text{SC}(\text{SO}_2\text{F})=\text{C}=\text{O}$ and paraaldehyde (28):

In a 50 ml Carius tube 0.66 g of 21 (2.64 mmol) and 0.08 g of trioxane (TCI, 0.89 mmol) were heated to 155 °C after evacuuation at -196°C for 2 hours. The mixture had darkened and 0.60 g of colorless liquid was vacuum-transferred; a dark residue was left behind. Of the three components (g.c., 20 % SE-30, 45/60, 3m, 95 °C) the major constituent ($R_f = 15.0$ min) was collected via preparative
gas chromatography (the same column was used), yielding 0.25 g of product, probably $F_5SC(SO_2F)=COCH_2O$, 28, (34 %), as a colorless, volatile liquid of a peculiar odor. The elemental analysis could not be obtained correctly, as the liquid deposited some solid after some time, and underwent an apparent chemical transformation.

$^1H$ n.m.r. spectrum (300 MHz, internal Si(CH$_3$)$_4$, CDCl$_3$): $\delta_1 = 7.10$ ppm, t, area = 1.0, $J = 4.65$ Hz ($= J_{12} = J_{1X}$); $\delta_2 = 7.26$ ppm, $\approx$ sextet, area = 1.0, $J_{\text{average}} = 3.48$ Hz;

$^{19}F$ n.m.r. spectrum: (CDCl$_3$, external CCl$_3$F): AB$_4X$ $\phi_A = 74.4$ ppm, 9 lines, area = 1.00; $\phi_B = 68.3$ ppm, d-m-d, area = 4.87; $\phi_X = 60.6$ ppm, d-p, area = 1.05 SO$_2$F); $J_{AB} = 155.3$ Hz, $J_{BH}^1$ or $J_{BH}^2 = 3.44$ Hz (average); $J_{BX} = 9.67$ Hz; $J_{1X} = 4.65$ Hz.

I.r. spectrum (neat sample, capillary film on KBr, cm$^{-1}$): 3142, w-m; 3051, w; 2981, vw; 1614, w; 1440, s-vs; 1379, s; 1227, s-vs; 1135, w; 1060, vw; 1005, m; 867, vs; 820, s-vs; 775, s; 710, vw; 687, m-s; 631, m-s; 582, s-vs; 535, m; 481, w-m; 443, m.

G.c.mass spectrum (e.i., 70 eV, mass, fragment, % > 1): 236, (M-CH$_2$O)$_2^+$, 6.8; 217, (M-CH$_2$O$_2$-F)$^+$, 4.5; 133, C$_3$HS$_2$O$_2^+$, 2.6; 129, $^{34}$SF$_5^+$, SF$_3$C$_2$O$^+$, 3.3; 128, $^{33}$SF$_5^+$, CHFO$_2$S$_2^+$, 8.5; 127, SF$_5^+$, 75.5; 109, CHO$_2$S$_2^+$, 2.6; 108, SF$_4^+$, 4.3; 91, $^{34}$SF$_3^+$, 3.8; 90, $^{33}$SF$_3^+$, 1.1; 89, $^{32}$SF$_3^+$, 100.0; 83, SO$_2$F$^+$, 11.5; 77, CHO$_2$S$^+$, 2.3; 76, CO$_2$S$^+$, 1.7; 70,
Product of the autoconversion of 28 (29):

A sample of 28 was kept for several weeks sealed in a glass tube in the freezer (-12°C). It was found that the originally clear liquid was partially converted to a white solid. The tube was opened and the contents were dissolved in acetone and the solvent was allowed to evaporate slowly at room temperature; within several days, small needle-shaped crystals were obtained. This material was apparently free of impurities. The conversion seems to be more complete in capillary tubes where after several weeks of storage seemingly all the liquid (28) was converted to a solid.

$^1$H n.m.r. spectrum (400 MHz, CDCl$_3$, internal Si(CH$_3$)$_4$): ABX $\delta_A = 4.448$ ppm, d-m, multiplet is probably d-p (partially superposed with second line of $\delta_A$); $J_{AB} = 12.9$ Hz; $J_{ASF_4F} = 2.2$ Hz; $J_{ASO_2F}$ or $J_{AX} = 4.4$ Hz. $\delta_B = 4.512$ ppm, d (each line has shoulders), area A+B = 2.00 (CH$_{AHB}$). $\delta_X = 5.98$ ppm, m, area = 0.85. F$_5$SCH.

$^{19}$F n.m.r. spectrum (375 MHz, CDCl$_3$, internal CCl$_3$F): AB$_4$X, $\phi_A = 76.5$ ppm, 9 lines; $\phi_B = 68.3$ ppm, asymmetric doublet; $\phi_X = 62.3$ ppm, ill-resolved pentet (some lines have shoulders). $J_{AB} = 146.6$ Hz; $J_{BX} \approx 11.7$ Hz.
I.r. spectrum (neat sample, squeezed on KBr): 2972, m;
2909, vw; 1431, vs; 1377, m; 1257, m; 1237, m, sh; 1222, s;
1202, m-s; 1143, s; 1080, m; 1060, m; 1045, m; 988, m; 940,
vw; 877, s-vs, sh; 857, vs; 803, s; 743, m-s; 726, m-s; 706,
m; 677, m-s; 649, m; 634, m-s; 620, m-s; 600, m; 580, s-vs;
554, s; 534, s; 483, s.


F₅SC(SO₂F)=C=O • O=C(N(CH₃)₂)₂ (26):

To 0.68 g of ((CH₃)₂N)₂C=O (5.9 mmol) in a 10 ml pear-shaped flask, equipped with a Kontes Teflon valve connector, 1.18 g of F₅SC(SO₂F)=C=O (4.7 mmol) were added through vacuum-transfer. The mixture was allowed to attain room-temperature slowly. It bubbled slightly and became viscous. After warming cautiously with a heat-gun, and keeping the sample at room-temperature for 45 min, volatile materials were removed by pumping them off on the vacuum-line overnight, leading to a loss of 0.24 g. The sample was still oily and was re-crystallized from CH₂Cl₂ (-12°C), where large colorless, extremely moisture-sensitive crystals were formed. Yield: 0.94 g = 54.5 %. M.p. = 100-102°C (sealed tube).

1H n.m.r. (CDCl₃, Si(CH₃)₄ ext.): δ = 3.42 ppm, s.

19F n.m.r. (CH₂Cl₂, CCl₃F ext.): (AB₄X) φ = 89.6 ppm, 9 lines, int. 1.0 (F_A); φ = 79.7 ppm, d-m, int. = 3.88 (F_B);
$\phi = 71.3$ ppm, $p$, int. $= 1.0$ (F$\text{X,}$SO$\text{2F}$); $J_{AB} = 154.2$ Hz; $J_{BX} = 9.1$ Hz.

I.r. spectrum (cm$^{-1}$, neat sample): 2910, w, broad; 1726, s; 1669,s; 1653,s,sh; 1531, m; 1442, m-w; 1414, m-s; 1390, s; 1248, m-s; 1203, s; 1179, s; 1057, m-s; 1049, m-s; 895, m; 858, vs; 842, vs; 777, vs; 729, vs; 713, s; 660, vs; 615, m; 587,m.

Anal. calcd for $\text{C}_7\text{H}_12\text{F}_6\text{N}_2\text{O}_4\text{S}_2$: C, 22.96; H, 3.03; F, 31.1; N, 7.65; S, 17.50. Found: C, 22.10; H, 3.46; F, 31.3; N, 8.27; S, 16.70.

$\text{F}_5\text{SC(SO}_2\text{F)}=\text{C}=0$ and $\text{SF}_5\text{Br}$:

Pentafluorosulfur bromide (0.55 g) and 21 were heated in a 50 ml Carius tube for 2 days to 110°C. After $\approx 1$ h, large crystals were observed at the colder parts of the tube, but disappeared later. After 2 days, 0.87 g of a brownish liquid was transferred out; its i.r. spectrum was almost identical with the i.r. spectrum of crude $\text{F}_5\text{SCBr(SO}_2\text{F)}\text{COF}$ (25b). Repeated fractional distillation resulted in 0.12 g of a slightly brown solid. Both the i.r. and the $^{19}$F n.m.r. spectrum were consistent with the composition 25b.

$\text{F}_5\text{SC(SO}_2\text{F)}=\text{C}=0$ and $\text{SO}_3$:

Ketene 21 (0.96 g, 3.8 mmol) was heated with (SO$_3$)$_3$ (0.31 g, 3.9 mmol) under anaerobic conditions in a 50 ml
Carius tube (the lower two thirds were in the upright oven) to 128°C (20 hours) then to 150°C (50 hours). The mixture was slightly darkened. Infrared spectroscopy (gas phase) showed only a weak CO$_2$ band; the liquid sample showed a very strong keten band, and $^{19}$F n.m.r. indicated chiefly unchanged keten, and bands at $\approx 73$, 61 and 50 ppm (weak). There was no sign of an A$_2$BC spectrum as would be expected for the compound F$_4$S=C(SO$_2$F)$_2$. The product was condensed back into the Carius tube and heated to 235°C (20 hours). I.r spectroscopy indicated an increase in CO$_2$, and the presence of $^{21}$, SiF$_4$ and perhaps COF$_2$ (weak). The mixture was condensed back into the reaction vessel and heated for another 24 h to 270°C. A comparatively strong CO$_2$ band was present in the i.r spectrum, but there was also still $^{21}$ and SF$_4$ (very strong). The product was condensed out (1.08 g before, 0.36 g after degassing) and its $^{19}$F n.m.r spectrum showed $^{21}$, + bands at $\approx 69$, $\approx 56$ (5:1, probably F$_5$SCH$_2$SO$_2$F) and 45 ppm.

**Preparation of Cs$^+$[F$_5$SC(SO$_2$F)COF]⁻ (25):**

Into a 30 ml Pyrex-glass reaction vessel, with a ground glass cap and a side-arm with a ground glass stopcock containing 3.0555 g of dried CsF (20.1 mmol, PCR) 2.62 g of acetonitrile was distilled from P$_4$O$_{10}$. Then 6.6535 g l (26.6 mmol) was vacuum-transferred (-196°C) into the vessel. Upon reaching room temperature most of the CsF dissolved in a
vigorously reaction causing the flask to warm. Dissolution slowed down towards the end and was complete 3 1/2 h of shaking. A slightly turbid and thick yellowish solution was obtained. Pumping away excess ketene and solvent at room temperature through a cold trap (-196°C) to constant weight left 8.0795 g of an off-white solid. This corresponds to a ketene: CsF ratio of 0.9985 and a yield of 99.91%, m.p. = 148-151 °C

$^{19}$F n.m.r. spectrum, AB$_4$MX (acetone, CCl$_3$F ext.): $\phi_A = 92.3$ ppm (9 lines), $\phi_B = 79.5$ ppm (d-m); $\phi_M = 68.5$ ppm, d-p, SO$_2$F; $\phi_X = 16.4$ ppm, d-p, COF; intensity A:B:M:X = 1.0: 4.4: 1.1: 1.0. $J_{AB} = 148.0$ Hz; $J_{BM} = 9.7$ Hz; $J_{BX} = 21.5$ Hz; $J_{MX} = 7.1$ Hz.

$^{13}$C n.m.r. spectrum (100.6 MHz, CD$_3$CN, Si(CH$_3$)$_4$): $\delta_1 = 104.33$ ppm, either d-quartet or d-quintet, but a quintet is assumed, F$_5$SC; $\delta_2 = 174.81$ ppm, d, COF; $J_{1B} = 20.7$ Hz; $J_{1COF} = 70.60$ Hz; $J_{2COF} = 297.5$ Hz.

Infrared spectrum: (neat sample, NaCl, cm$^{-1}$): 1766 (vs); 1395 (m); 1380 (m); 1257 (m); 1203 (S); 1077 (m); 948 (w); 864 (s); 843 (vs); 789 (vs); 747 (m); 726 (m); 705 (m); 666 (m); 616 (s); 584 (s); 569 (s).

Anal. calc.: for C$_2$CsF$_7$O$_3$S$_2$: C, 5.97; Cs, 33.06; F, 33.1; S, 15.95. Found: C, 6.00; Cs, 33.13; F, 33.6; S, 15.95

HgF$_2$ + 21 (45):

Fresh mercury(II)-fluoride (0.4089 g, 1.71 mmol,
Aldrich) was kept overnight in a 50 ml pear-shaped flask with a Kontes Teflon valve connector under high vacuum (0.003 torr) at room temperature. A solution of 0.91 g of 21 (3.64 mmol) in 1.0 ml of dry acetonitrile was added via vacuum transfer, and the flask was allowed to warm up slowly, while being swirled, once the contents started melting. The mercury fluoride dissolved largely in a vigorous reaction, and the process was moderated by dipping the flask occasionally into liquid nitrogen. A small residue was left behind, but was dissolved after keeping the flask at room temperature for 66 hours. Excess ketene and solvent were removed at room temperature by pumping on the flask through a -196°C trap for 8 hours. The rate of weight loss slowed down constantly, but never reached zero. After 8 hours, 1.2559 g of a white slightly sticky mass was left behind. Although this corresponds to a ratio 21:HgF_2 = 1.976:1.000, the product was apparently a non-stoichiometric compound (F_5SC(SO_2F)CFO)_2Hg·nCH_3CN, as inferred from the melting behavior and the i.r. spectrum, and therefore the complex must be volatile. Melting range 40-≈80°C, bubbling starts at 104°C.

^19F n.m.r. spectrum (CH_3CN, external CCl_3F): \( \phi_A = 90.6 \) ppm, 9 lines , area = 1.00; \( \phi_B = 79.9 \) ppm, d-m, area = 4.13; \( \phi_{SO_2F} = 66.1 \) ppm, broad singlet, area = 0.97; \( \phi_{CFO} = 21.2 \) ppm, broad singlet, area = 0.97; \( J_{AB} = 154.3 \) Hz. All lines are broadened.
I.r spectrum (neat sample on KBr, cm\(^{-1}\)): 3016, w; 2966, w; 2952, w-m; 2629, w; 2423, w; 2385, w; 2347, w-m (CO\(_2\)?): 2305, m; 2276, m; 2053, w, vbr; 1826, vs, br; 1414, vs, br; 1288, s; 1233, s-vs; 1195, s-vs; 1126, s-vs; 1078, s, sh; 1040, s; 964, m-s, br; 855, vs, br; 792, sh, s-vs, br; 682, m-s, br; 663, m-s; 618, s; 583, s-vs, br; 493, m-s; 465, s.

AgF + 21 (25d):

The sample was prepared as described with 25. Initially, strong reaction was observed which slowed down eventually. A colorless liquid was obtained, which left a glassy mass behind, when the solvent was pumped off at room temperature, but no weight corresponding to a stoichiometric composition was obtained ([F\(_5\)SC(SO\(_2\)F)CFO]-Ag\(^+\cdot n\)CH\(_3\)CN). Upon warming, a weight corresponding to less than a 1:1 composition of AgF and 21 was obtained. The compound deposited slowly a dark precipitate when exposed to light, but this seems to suppress further decomposition, as the complex could still be detected by \(^{19}\)F n.m.r. spectroscopy when a CH\(_3\)CN solution was exposed to the irradiation of a sun lamp overnight.

\(^{19}\)F n.m.r. spectrum (CH\(_3\)CN, external CCl\(_3\)F): AB\(_4\)MX \(\phi_A\) = 94.0 ppm, 9 lines, area = 1.09; \(\phi_B\) = 79.9 ppm, d-m, area = 4.0; \(\phi_M\) = 69.7 ppm, d-p (not well resolved), area = 0.89 (SO\(_2\)F); \(\phi_X\) = 17.0 ppm, d-p, area = 0.78 (CFO). \(J_{AB}\) = 155.3 Hz; \(J_{BM}\) = 10.2 Hz; \(J_{BX}\) = 19.7 Hz; \(J_{MX}\) = 6.3 Hz.
**NaN₃ + 21 (46):**

When a solution of 21 in dry acetonitrile was shaken with a few crystals of NaN₃ at 0°C in an n.m.r. tube, the salt dissolved rapidly, and a pale yellow solution was formed. ¹⁹F n.m.r. spectroscopy indicated the complete consumption of 21, with the formation of a single anionic species. When the solution was allowed to warm, it started to bubble, and turned dark yellow. From ¹⁹F n.m.r. measurements it is inferred that the compound [F₅SC(SO₂F)CF₀]⁻ Na⁺ is formed in this decomposition process of the primary azide anion, and the integrated band intensity is less than 1/2 of the initial intensity of 21.

In a preparative scale experiment, HCl (aqueous) was added after dissolution of most of the NaN₃, and n.m.r. measurements and g.c. indicated the formation of F₅SCH₂SO₂F, rather than F₅SCH(SO₂F)NH₂, which should have a substantially different chemical shift and retention time.

¹⁹F n.m.r. spectrum of [F₅SC(SO₂F)CON₃]⁻ Na⁺ (CH₃CN, external CCl₃F): AB₄X \( \phi_A = 95.9 \) ppm; \( \phi_B = 82.6 \) ppm; \( \phi_{SO₂F} = 69.5 \) ppm; \( J_{AB} = 151.5 \) Hz; \( J_{BSO₂F} = 10.3 \) Hz.

**Pentafluorothiofluorosulfonylchloroacetylfuoride (25a):**

To 2.0 g of Cs⁺ [F₅SC(SO₂F)COF]⁻ (5.0 mmol) in a 75 ml steel reaction vessel, chlorine (0.551, 210 torr, room temperature, 6.3 mmol) was added via vacuum transfer. The vessel was heated to 85°C for 2h, then kept at room
temperature for 18 h. The volatile products (1.25 g), as collected by vacuum transfer, were obtained as a clear liquid. Excess chlorine was removed by shaking with ≈1g of mercury and the product thus obtained by vacuum transfer, a clear and colorless liquid, was analytically pure (1.14 g, 75.3 %).

$^{19}$F n.m.r. spectrum (pentane, CCl$_3$F ext.): $\phi_1 = 63.6$ ppm, m, int. = 5.1, (SF$_5$); $\phi_2 = 49.3$ ppm, m, int. = 1.0 (SO$_2$F); $\phi_3 = 31.3$ ppm, m, int. = 1.0 (COF).

Infrared spectrum (neat liquid, KBr, cm$^{-1}$): 1873 (s); 1853 (s); 1458 (vs); 1223 (vs); 1175 (s); 955 (m); 890 (vs); 834 (s); 808 (m); 790 (m); 739 (vw); 700 (m); 694 (m); 669 (vw); 635 (m-w); 623 (m-w); 612 (m-s); 579 (vs); 569 (vs,sh); 504 (w); 496 (w); 472 (w); 417 (w).

Mass spectrum (electrom impact, 70 eV, mass, species, >1%): 158, C$_2$ClFOS$^+$, 23.4; 155, C$_3$ClF$_4$S$^+$, C$_2$FO$_3$S$_2^+$, 5.5; 129, C$_2$ClF$_2$S$^+$, 2.2; 127, SF$_5^+$, CFO$_2$S$^+$; 125, CHO$_3$S$_2^+$, 1.1; 124, C$_2$HFOS$_2^+$, 2.9; 113, C$_2$ClF$_2$O$^+$, 5.2; 108, SF$_4^+$, 1.5; 103, CH$_2$SF$_3^+$, 2.0; 101, CF$_3$S$^+$, 1.7; 95, C$_3$ClOS$^+$, CFS$_2^+$, 7.0; 94, C$_2$ClFO$^+$, 8.7; 91, C$_2$FOS$^+$, 3$^4$SF$_3^+$, 4.8; 89, SF$_3^+$, 100.0; 86, $^{35}$ClFS$^+$, 6.6; 85, C$_3$ClF$_2^+$, 8.5; 83, SO$_2$F$^+$, 3.4; 81, C$_2$F$_3^+$ (rearr.), 11.2; 79, CSOF$^+$, 1.3; 78, C$_2$FO$_2^+$ (rearr.), 1.1; 77, CHS$_2$O$^+$, C$_2$H$_2$FS$^+$ (two fragments), 1.4, 2.5; 76, CSO$_2^+$, C$_2$HFS$^+$, 1.2; 75, C$_2$FS$^+$, 8.0; 73, C$_2$HOS$^+$, 1.1; 72, C$_2$OS$^+$, 1.2; 70, SF$_2^+$, 44.8; 69, CF$_3^+$ (rearr.), 8.6; 67, SOF$^+$, 58.9; 66, C$_3$ClF$^+$, 19.8; 65, SO$_2$H$^+$, 5.6; 64, SO$_2^+$,
54.3; 63, CFS⁺, 17.3; 60, COS⁺, 1.8; 51, SF⁺, 12.0.

Anal. calcd for C₂ClF₇0₃S₂: C, 7.89; Cl, 11.64; F, 43.7; S, 21.05. Found: C, 7.85; Cl, 11.9; F, 44.0; S, 21.23.

Pentafluorosulfurfluorosulfonylchloromethane (25h):

In a 5.0 ml Pyrex flask 0.74 g of F₅SCCl(SO₂F)(COF) (2.4 mmol) were mixed with 0.65 g water (distilled). The flask was stoppered lightly and kept at room temperature for 21 h. After pipetting off the water layer, adding ≈ 1g of powdered P₄O₁₀ in the cold under swirling in small portions, and vacuum transferring away the volatile product, and repeating the drying procedure, 0.61 g (97.1%) of a clear liquid was obtained once. The material was analytically pure as seen by gas chromatography (30% SE-30, 2m, 120°C, Rᵢ = 15.3 min.), ¹H and ¹⁹F n.m.r.

Infrared spectrum (neat liquid, KBr, cm⁻¹): 2992 (m); 1448 (vs); 1262 (m-w); 1232 (s-vs); 1205 (m-s); 1192 (m-w,sh); 874 (vs); 851 (vs); 811 (s); 774 (m-s); 757 (w,sh); 688 (vw); 669 (m); 612 (m); 581 (s); 571 (s); 566 (s); 499 (vw); 490 (w-m); 463 (m); 453 (vw); 423 (vw); 405 (m).

¹H n.m.r. (neat liquid, CH₂Cl₂ ext. standard): δ = 6.14 ppm, pentet. Jₓᵧ = 4.20 Hz.

¹⁹F n.m.r. (neat liquid, ext. CCl₃F): φ₁ = 61.5 ppm, m, int. = 5.0, (degenerate AB₄ spectrum, SF₅); φ₂ = 49.8 ppm, p, int. = 1.0 (SO₂F). Jₓᵧ₂ = 9.5 Hz.
Mass spectrum (electron impact, 70eV, mass, species, >1%): 175, (M-SO₂F)+, 1.8; 131, (M-SF₅)+, 2.7; 129, (M+H-5F-Cl)⁺, 2.5; 127, SF₅⁺, 53.1; 108, SF₄⁺, 2.2; 91, C₂FOS⁺, 2.1; 89, SF₃⁺, 45.7; 83, SO₂F⁺, 2.6; 79, CFOS⁺, 1.8; 70, SF₂⁺, 7.9; 69, H₂SOF⁺, CF₃⁺, 16.7; 68, HSOF⁺, 1.6; 67, SOF⁺, 100.0; 64, SO₂⁺, 8.1; 63, CFS⁺, 1.2; 51, SF⁺, 4.8

Anal.calcd. for CHClF₆O₂S₂: C, 4.64; H, 0.39; Cl, 13.71; F, 44.1, S, 24.80. Found: C, 4.72; H, 0.34; Cl, 13.74; F, 44.3; S, 24.84.

F₅SCBr(SO₂F)COF (25b):

Bromine (0.37 g, 2.31 mmol) and 21 (0.95 g, 2.36 mmol) were gently heated in a 30 ml Carius tube under anaerobic conditions with a heat gun, when most of the bromine was consumed rapidly; heating was continued at 82°C (1 h), but a brownish color persisted. The product was vacuum transferred out (0.67 g), treated with mercury (~ 2 g) to remove excess bromine, after which 0.42 g of crude F₅SCBr(SO₂F)COF was collected. This material could not be purified by fractional condensation or distillation and was used as such in subsequent reactions.

I.r. spectrum of the crude material (neat sample, KBr, cm⁻¹): 2985, vw; 2674, vw; 1868, s,sh; 1846, vs; 1455, s-vs; 1290, w; 1230, s; 1171, s; 1103, w; 946, m-s; 881, vs; 835, s,sh; 814, s; 791, s; 692, s; 650, m; 610, s; 579, s; 568, s,sh; 497, m; 492, m; 466, m; 458, m.
$^{19}$F n.m.r. (CCl$_3$, external CCl$_3$F): $\delta = 74$ ppm, int. = 4 (impurity); $\delta = 67.6$ ppm, m, int. = 30 (SF$_5$); $\delta = 60-62$, several lines, int. = 4 (impurity); $\delta = 50.6$ ppm, m; int. = 6 (SO$_2$F); $\delta = 35.3$ ppm, m, int. = 6 (COF).

F$_5$SCHBrSO$_2$F (25j):

To 1.31 g of crude F$_5$SCBr(SO$_2$F)COF in a 4.0 ml conical vial 5 drops of water were added, and the flask was lightly stoppered; no visible reaction occurred. The mixture was kept at room temperature for 18 hours and occasionally swirled. Phosphorous pentoxide was added with cooling and the volatile materials were transferred out (0.84 g); a light, slightly yellow liquid was obtained. Gas chromatography (20% SE-30, on Chromosorb W, 3m, 120°C): 2 bands, $R_f = 5.6$ minutes, area = 14.6 % (same $R_f$ as F$_5$SCH$_2$SO$_2$F), $R_f = 12$ minutes, area = 85.6 % (F$_5$SCHBr(SO$_2$F)). Preparative g.c. (20% SE-30, 3m, 115°C) afforded 0.24 g (23.5 %) of a colorless liquid.

$^1$H n.m.r. spectrum (CDCl$_3$, Si(CH$_3$)$_4$, 300 MHz): $\delta = 6.40$ ppm, p, 4.58 Hz.

$^{19}$F n.m.r. spectrum (CDCl$_3$, external CCl$_3$F): $\delta = 64-74$ ppm, distorted AB$_4$ spectrum, maximum at 66.7 ppm, int. = 5.2 (SF$_5$); $\delta = 53.3$ ppm, m, int. = 1.0 (SO$_2$F).

Infrared spectrum (neat sample, KBr, cm$^{-1}$): 2997, m; 2922, vw; 1445, s; 1256, w-m; 1225, m-s; 1166, w; 1072, w, broad; 871, vs; 820, s; 785, m; 761, m-s; 690, w; 663, m;
Mass spectrum (e.i., m/e, fragment, % > 1, only $^{79}$Br listed): 219, (M-SO$_2$F)$^+$, 1.6; 175, (M-SF$_5$)$^+$, 20.5; 129, $^{34}$SF$_5$$^+$, 1.9; 127, $^{32}$SF$_5$$^+$, 52; 111, CHBrF$^+$, (rearr.), 30.0; 108, SF$_4$$^+$, 2.2; 97, CH$_2$FS$_2$$^+$, 4.0; 95, CSO$_2$F$^+$, CFS$_2$$^+$, 1.6; 93, CH$_2$Br$^+$, 2.4; 92, CHBr$^+$, 11.8; 91, two fragments, CBr$^+$, ?, 2.4, 2.5; 89, SF$_3$$^+$, 61.5; 83, SO$_2$F$^+$, CHSF$_2$$^+$, 1.3; 80, HBr$^+$, 3.6; 79, Br$^+$, 4.4; 70, SF$_2$$^+$, 10.2; 69, CF$_3$$^+$, 5.0; 68, FSOH$^+$(?), 1.1; 67, FSO$^+$, 100.0; 64, CHSF$^+$, SO$_2$$^+$, 7.5; 63, CSF$^+$, 1.2; 51, SF$^+$, 6.3.

Anal. calcd for CHBrF$_6$O$_2$S$_2$: C, 3.96; H, 0.33; F, 37.6; S, 21.16; Br, 26.37. Found: C, 4.02; H, 0.37; F, 38.0; S, 21.26; Br, 26.28.

F$_5$SCH(SO$_2$F)CH$_2$CH$_2$Br (25i):

A 75 ml stainless steel bomb was charged with 0.24 g of benzoyl peroxide (1.0 mmol, Aldrich), 1.73 g of crude F$_5$SCBr(SO$_2$F)COF ($\approx$ 5 mmol) and 0.57 g of ethylene (21.9 mmol) and heated at 95°C for 24 hours. At room temperature, a colorless liquid (1.01 g after degassing) was transferred out; its i.r. and $^{19}$F n.m.r. spectra was virtually identical with the spectra of authentic F$_5$SCH$_2$SO$_2$F. The residue in the bomb (0.78 g), a brown, partially crystalline mass, was dissolved in ether, and both the i.r. and $^{19}$F spectra indicated the absence of a COF group in the crude product,
but there was a C=O band at 1690 cm\(^{-1}\). The material that caused this band could not be removed by vacuum distillation, but stirring a CH\(_2\)Cl\(_2\) solution of the collected impure fraction, the pot residue and the rinse of the distillation head with dilute NaHCO\(_3\) solution for circa 15 minutes caused its disappearance. The CH\(_2\)Cl\(_2\) solution was pipetted off, dried (Na\(_2\)SO\(_4\)) and transferred to a 10 ml flask. Distillation (1 mm Hg) afforded a small amount of colorless liquid (45-46°C), but lowering the pressure (≈ 0.5 mm Hg), and heating the distillation head with a heat gun yielded some more product (0.23 g, 14.0%).

\(^1\)H n.m.r. spectrum (neat sample, external Si(CH\(_3\))\(_4\)): 
\(\delta_1 = 5.79 \text{ ppm}, \text{ m, partially resolved, probably a septet, } J \approx 5.4 \text{ Hz, intensity } = 0.97 \text{ (F}_5\text{SCH)}; \delta_2 = 3.23 \text{ ppm}, \text{ m, intensity } = 2.03 \text{ (CH}\_2\text{CH}_2\text{Br)}; \delta_3 = 3.94 \text{ ppm, t, } J_{23} = 6.0 \text{ Hz, intensity } = 2.00 \text{ (CH}_2\text{Br). All lines are broadened.}

\(^19\)F n.m.r. spectrum (neat liquid, external CCl\(_3\)F): 
AB\(_4\)X \(\phi_A = 75.3 \text{ ppm, d of nine line pattern (the doublet is poorly resolved), intensity } = 1.00; \phi_B = 66.0 \text{ ppm, d-m, intensity } = 4.00; \phi_X = 54.8 \text{ ppm, } \approx \text{ pentet, intensity } = 0.94 \text{ (SO}_2\text{F). } J_{AB} = 149.6 \text{ Hz; } J_{BX} = 11.3 \text{ Hz; } J_{AH}^1 \approx 4.2 \text{ Hz.}

I.r. spectrum (neat liquid on KBr, cm\(^{-1}\)): 3062, vw; 2960, w; 2940, w,sh; 1428, s-vs; 1364, vw; 1318, w; 1268, m,sh; 1252, m; 1216, s; 1179, w; 1158, vw,sh; 989, w,br; 970, vw,sh; 941, w; 871, vs; 820, m-s; 809, m-s,sh; 775, m-s; 756, m,sh; 672, m; 645, vw; 613, m; 589, s; 579, s,sh;
Mass spectrum (e.i., 70 eV, m/e, fragment, % > 1, only

\[ {^79}\text{Br} \]): 330, M\(^+\), 0.5; 253, \((\text{C}_3\text{H}_2)^{81}\text{BrF}_2\text{O}_2\text{S}_2 = \text{M}-3\text{H}-4\text{F})^+\), 6.6;
252, (\text{M}+\text{H}-\text{Br})^+, 3.4; 251, (\text{M}-\text{Br})^+, \text{C}_3\text{H}_2^{79}\text{BrF}_2\text{O}_2\text{S}_2^+, 71.0;
246, (\text{M}-\text{SO}_2\text{F}-\text{H})^+, 6.2; 202, (\text{M}-\text{SF}_5-2\text{H})^+, 7.4; 167, (\text{M}-\text{Br}-\text{SO}_2\text{F}-\text{H})^+, 4.2; 135, (\text{M}-\text{CH}_2\text{Br}-\text{SO}_2\text{F}-\text{F})^+, 1.7; 127, \text{SF}_5^+, 20.3;
123, \text{C}_2\text{FOS}_2^+, 5.2; 122, \text{C}_2\text{H}_2\text{O}_2\text{S}_2^+, 1.9; 119, \text{C}_3\text{H}_3\text{Br}^+, 46.1;
113, \text{C}_2\text{F}_3\text{S}^+, 1.4; 111, \text{CFOS}_2^+, 1.9; \text{CHO}_2\text{S}_2^+, 2.3; 108, 
\text{CO}_2\text{S}_2^+, \text{SF}_4^+, 1.1; 107, \text{C}_2\text{FS}_2^+, 1.8; 97, \text{CH}_2\text{SO}_2\text{F}^+, 4.7; 93,
\text{CH}_2\text{Br}^+, 16.7; 91, \text{C}_3\text{H}_4\text{FS}^+, 3.4; 89, \text{SF}_3^+, 47.4; 87, \text{C}_3\text{FS}^+, 1.2; 83, \text{SO}_2\text{F}^+, 2.9; 81, \text{CH}_2\text{SOF}^+, 1.5; 79, \text{CSOF}^+, 2.7; 71,
\text{C}_3\text{H}_3\text{S}^+, 1.4; 70, \text{SF}_2^+, 6.5; 69, \text{C}_3\text{HS}^+, \text{CF}_3^+, 4.6; 68, \text{C}_3\text{S}^+, 1.3; 67, \text{FOS}^+, 25.5; 65, \text{CH}_2\text{SF}^+, 3.1; 64, \text{SO}_2^+, \text{CHSF}^+, 2.0;
60, \text{C}_2\text{H}_4\text{S}^+, 7.9; 59, \text{C}_2\text{H}_3\text{S}^+, 100.0; 57, \text{C}_2\text{HS}^+, 7.9; 55, 
\text{C}_3\text{F}^+, 8.7; 51, \text{SF}^+, 5.9.

\textbf{Anal. calcd} for \text{C}_3\text{H}_5\text{BrF}_6\text{O}_2\text{S}_2: \text{C}, 10.88; \text{H}, 1.52; \text{Br},
24.13; \text{F}, 34.43; \text{S}, 19.37. \text{Found: C}, 10.85; \text{H}, 1.50; \text{Br},
24.17; \text{F}, 34.5; \text{S}, 19.33.

\textbf{F}_5\text{SCBr}_2\text{SO}_2\text{F} (25f): 

Bromine was added dropwise to a stirred solution of 25 (2.00 g, 5.0 mmol) in a 25 ml flask in 0.5 ml of acetonitrile at room temperature. A white precipitate was formed instantly, and after some time, bubbling and foaming ensued, with the paling of the bromine color. More bromine was added occasionally, until a faint red color persisted
and no more bubbles were observed (~ 15 minutes). Stirring was continued for 5 min. $^{19}$F n.m.r. indicated only one product, devoid of a COF group, to be present. Because of the failure to separate the product from the solvent by distillation, the reaction mixture was subjected to fractional condensation (-20, -78, -196°C), where the material in the -20°C trap was used in three successive trap-to-trap distillations respectively. This resulted in 0.73 g (38.4 %) of an almost colorless, glassy, partially crystalline material, m.p. 39-43°C. The material of the -78 °C trap was subjected to repeated fractional condensations, but it was not possible to obtain a solid material. The product (-10 °C trap, 0.56 g) showed an almost similar i.r. spectrum to the solid, while the -78 °C trap (0.59 g) contained sizable amounts of CH$_3$CN. The -196 °C trap (0.64 g) was almost free of any SF$_5$ product.

$^{19}$ F n.m.r. spectrum (CH$_3$CN, CCl$_3$F ext.): AB$_4$X spectrum, $\phi_A = 67.0$ ppm, 9 lines, int. = 1.00; $\phi_B = 58.0$ ppm, d-m, int. = 4.11; $\phi_X$ (SO$_2$F) = 39.9 ppm, ~ s (not resolved); $J_{AB} = 149.0$ Hz.

I.r. spectrum (neat samples, on KBr, cm$^{-1}$): 1441 (vs); 1268 (vw,sh); 1222 (vs); 1215 (m); 1100 (m,sh); 1087 (w,sh); 1049 (vw); 972 (w); 882 (vs); 857 (vs); 831 (vs); 787 (m-s); 767 (m-s); 749 (m,sh); 715 (m-s); 703 (m-s); 685 (s); 654 (vw); 618 (m-s); 605 (s); 566 (vs); 498 (m); 489 (m); 471 (m); 463 (m).
Mass spectrum (e.i., 70 eV, mass, fragment, % > 10): 299, F$_5$S$^{79}$Br$_{81}$Br$^+$, 53.4 (triplet, 1:2:1); 281,; 255, C$^{81}$Br$_{79}$BrSO$_2$F$^+$, 54.3 (triplet); 191, CBr$_2$F$^+$ (rearr.), 24.8 (triplet); 172, C$^{81}$Br$_{79}$Br$^+$, 19.4 (triplet; )159, CH$^{79}$BrSOF$^+$, 11.0; 127, SF$_5^+$, 33.9; 92, C$^{79}$Br$^+$, 26.0; 89, SF$_3^+$, 92.6; 80, H$^{79}$Br, 15.7; 79, 79Br, CFSO$^+$, 14.2; 70, SF$_2^+$, 20.7; 67, FSO$^+$, 100.0; 64, SO$_2^+$, 64.8; 48, SO$^+$, 30.1; 44, CS$^+$, 22.1.

Anal. calcd for CBr$_2$F$_6$O$_2$S$_2$: C, 3.14; Br, 41.84; F, 29.8; S, 16.79. Found: C, 3.32; Br, 40.95; F, 29.4; S, 16.65.

F$_5$SCCl(SO$_2$F)COCl (25g):

2.12 g of Cs$^+$ [F$_5$SC(SO$_2$F)COF]$^-$ (5.3 mmol) and 1.31 g Cl$_2$ (18.5 mmol) in a 300 ml Pyrex-quartz vessel were irradiated with a 250 W GE sunlamp from a distance of 5 cm (4h) and 10 cm (12h). The pressure in the reaction vessel had dropped by 1/3, indicating the consumption of 1 equivalent of chlorine; an i.r. spectrum (100 torr, 10 cm path length, KBr windows) showed a COF group, but neither CO nor COClF bands. Irradiation was continued for 24 h with no change in the i.r. spectrum. Dry acetonitrile (1 ml) was then added by vacuum transfer, and irradiation (12h, 10cm) was continued. After 12h, the chlorine color had faded and another 0.84 g of Cl$_2$ (11.8 mmol) were added. After another 17 h of irradiation (20 cm), the chlorine color had faded again. The product was subjected to fractional condensation
(-30, -78, -196°C). The material in the -30°C trap (0.31 g) was (g.c. 5 % QF-1 on Chromosorb W, 1m, 70 °C) 78.6 % F$_5$SCCl(SO$_2$F)COCl, 6.5 % F$_5$SCCl(SO$_2$F)COF, 7.8 % CH$_3$CN and 7.5 % of an unknown material. The contents of the -78°C trap were again subjected to fractional condensation (-45, -78,-196°C). The -45°C trap contained 0.77 g of a product that contained (g.c.) 45.4 % F$_5$SCCl(SO$_2$F)COCl and 14.3 % F$_5$SCCl(SO$_2$F)COF. Almost no product was in the -78°C trap and the material in the -19°C (0.39 g) contained no F$_5$SCCl(SO$_2$F)COCl. The two products from the trap-to-trap distillations were combined and subjected to preparative gas chromatography (QF-1, 1m, 70 °C). A total of 0.20 g of a clear colorless liquid were collected. Yield: 11.8 %. When the above procedure was carried out with CH$_3$CN from the very beginning, the initial formation of F$_5$SCCl(SO$_2$F)COF was observed ($^{19}$F n.m.r.), which upon successive irradiation and chlorine addition decreased, and F$_5$SCCl(SO$_2$F)COCl became the major product. There were other minor SF$_5$ products, which could not be collected in sufficient quantity so as to allow their identification. From comparisons of chemical shift values, one product could have been F$_5$SCHClSO$_2$F and another F$_5$SCCl$_2$SO$_2$F, as conjectured from the similar values of F$_5$SCBr$_2$SO$_2$F ($\phi_A \approx 66$ ppm; $\phi_B \approx 54$ ppm; $\phi_X = 39.5$ ppm, X = SO$_2$F). The product in this case was washed with water, dried and also subjected to preparative gas chromatography, which allowed the collection of 0.16 g of F$_5$SCHClSO$_2$F (12.9 %),
apparently formed by the hydrolysis of $\text{F}_5\text{SCl(SO}_2\text{F)}\text{COCI}$.

$^{19}\text{F}$ n.n.r. spectrum (neat sample, $\text{CCl}_3\text{F}$ ext.): $\phi = 64.5$ ppm, m, int. = 5.3 (SF$_5$); $\phi = 50.6$ ppm, m, int. = 1.0 (SO$_2$F).

Mass spectrum: (negative c.i., isobutane, mass, species > 5%): 257, $\text{C}_3\text{F}_6\text{O}_2\text{S}_2^-$, 10.9; 223, $\text{F}_5\text{SCHSO}_2\text{F}^-$, 46.9; 167, $\text{C}_2\text{F}_5\text{OS}^-$, 6.6; 165, $\text{CF}_3\text{O}_2\text{S}_2^-$, 9.6; 146, $\text{CF}_2\text{O}_2\text{S}_2^-$, 32.0; 130, $\text{CF}_2\text{OS}_2^-$, 9.5; 127, SF$_5^-$, 100.0; 83, SO$_2$F$^-$, 37.5.

Infrared spectrum (neat, KBr, cm$^{-1}$): 1793(s), 1769 (m-s,sh); 1454(vs); 1229(vs); 1022(s); 919(vssh); 890(vs); 878(vssh); 836(s); 798(s); 754(s); 737(s-m); 691(m); 665(vw); 615(s); 599(s); 577(s); 563(s); 515(w); 501(w); 491(w-m); 464(m).

Anal. calcd.: for $\text{C}_2\text{Cl}_2\text{F}_6\text{O}_3\text{S}_2^-$: C, 7.48; Cl, 22.08; F, 35.1. Found: C, 7.59; Cl, 22.30; F, 35.2.

Reaction of 25 with $\text{NO}^+\text{BF}_4^-$ in $\text{CH}_3\text{CN}$ (40):

To a solution of 0.69 g of nitrosonium terafluoroborate (5.9 mmol) in 3.0 ml of dry acetonitrile in a 25 ml round bottomed flask, equipped with magnetic stirring bar, a Claisen head, dropping funnel and a Drierite tube, 1.72 g of 25 (4.3 mmol) in 2.0 ml of dry acetonitrile was added dropwise. A yellowish color was noticed. After 5 minutes, the Claisen head was replaced with a distillation head and the pressure was lowered to 30 mm Hg. The mixture was stirred for 2 hours at room temperature when almost
complete transfer of the contents of the flask into the-196°C cold trap occurred. The cold-trap contents were partially yellow and had a bluish hue in the cold. Upon heating very little of a yellow oil came over; this material showed bands at 2256 and 2340 cm⁻¹. This oil was combined with the cold trap contents, which were now blue in the cold and yellowish-green at room temperature.

¹⁹F n.m.r. of this material (CCl₃F external): φ = 72.1 ppm, m, area = 5.9 (SF₅); φ = 58.9 ppm, m, area = 1.0 (SO₂F); φ = 49.7, m, area = 0.7 (COF). Small signals were also observed at 79.5, 70.0, 78.6, 78.0 76 (broad), 56.0, and 48.9 ppm. The band at 49.7 ppm (COF) had disappeared after keeping the solution at room temperature for 18 hours and several new sharp signals (low intensity) were observed at 80.3, 64.0, 57.0 and 33.6 ppm. The main bands were at φ = 72.9 ppm (m, SF₅), φ = 67.0 (weak) and 59.3 ppm (weak).

Because of this obvious instability of the alleged primary product F₅SC(NO)(SO₂F)COF, it was not tried to isolate it. Instead, water (0.2 ml) was added to the CH₃CN solution, which caused instant discoloration, and some gas bubbles were formed. After 17 hours at room temperature, a single product was observed in the ¹⁹F n.m.r spectrum, which was SF₅CH₂SO₂F. This was shown by adding a small amount of authentical SF₅CH₂SO₂F to the solution in the n.m.r tube, causing only an increase of all resonances that were originally present.
$E_4S=\overline{CSO_2OCF_2}$ (23):

In a vacuum compartment consisting of two 50 ml round bottomed flasks, connected by a transfer line with an outlet to a vacuum line 9.63 g (35.7 mmol) of sultone 1 and 6.68 g of $F_3B\cdot N(C_2H_5)_3$ (39.5 mmol) were heated slowly with stirring in one of the flasks at atmospheric pressure; the other flask was held at -196°C. When the oil bath temperature reached 76°C, the solution bubbled slightly and separated suddenly into two layers. Heating was continued for 2 1/2 minutes, with a final temperature of 80°C. The reaction flask was cooled (-196°C), the vacuum compartment was evacuated and then warmed to 60°C for 45 minutes. In the receiving flask, cooled to -196°C, 8.45 g of product was collected, which consisted ($^{19}$F n.m.r.) of a mixture of 2 parts of 21 and 1 part of 23. Yield of 21 + 23 = 94.8%.

Separation of 21 and 23:

The crude material from above was transferred to a 30 ml reaction flask with a Kontes Teflon valve containing 10.90 g (71.7 mmol) of comminuted cesium fluoride (PCR), shaken and kept at room temperature overnight. A second treatment with 8.58 g of CsF (56.4 mmol) for 3h removed the remaining keten completely, leaving 4.02g of almost pure 23. Yield =43.9 %.(with respect to 1).

To obtain a sample for analysis, part of the above material was subjected to trap-to-trap distillation , where
the traps were held at -10, -63, -110 and -196°C. The material that was collected in the -63°C trap (almost quantitative) was free of one impurity ($\phi = 73$ ppm in the $^{19}\text{F}$ n.m.r. spectrum). A second impurity ($\phi = 68$ ppm in $^{19}\text{F}$ n.m.r., supposedly 1) could be removed by heating the material from the trap-to-trap distillation shortly (1/2 min) with a small amount of $\text{F}_3\text{B} \cdot \text{N}(\text{C}_2\text{H}_5)_3$. The third impurity ($\phi = 70$ ppm, probably $\text{F}_5\text{SCH}_2\text{SO}_2\text{F}$) could not be removed, but accounted for only 1%. The cesium fluoride was rinsed with several small portions of acetone after the reaction and a total of 4.97g of $\text{F}_5\text{SC(SO}_2\text{F})\text{COF}^-\text{Cs}^+$ was obtained; no other compound could be detected.

$^{19}\text{F}$ n.m.r. spectrum (neat sample, $\text{CCl}_3\text{F}$ as external standard): AA'$\text{BCXX}'$ spectrum: $\phi_A = 51.87$ ppm, $\phi_B = 47.16$ ppm, $\phi_C = 40.90$ ppm, $\phi_X = -76.3$ ppm. Integral A:B:C:X = 2.0:1.0:1.0:1.7. $J_{A'B} = J_{AB} = 161.8$ Hz; $J_{AC} = J_{A'C} = 164.9$ Hz; $J_{BC} = 20.4$ Hz; $J_{AX} = 4.1$ Hz; $J_{BX} = 6.0$ Hz; $J_{CX} = 5.4$ Hz.

$^{13}\text{C}$ n.m.r. spectrum: (neat sample, $\text{CDCl}_3$ external): $\delta_\alpha = 80.7$ ppm, (approximately p-t), $J_{\text{SF}_4\text{C}^1} = 31.6$ Hz, $J_{\text{CF}_x\text{C}^1} = 7.8$ Hz; $\delta_\beta = 114.5$ ppm, d-t, $J_{\text{C}_2\text{FX}} = 282.4$ Hz, $J_{\text{C}_2\text{FB}} = 23.2$ Hz.

Mass spectrum (electron impact, 70 eV, mass, fragment, >1%): 250, $M^+ (32\text{S})$, 4.6; 190, (M-COS)$^+$ (rearr.), 2.1; 184, (-COF$_2$)$^+$, 1.3; 172, (M-COSF)$^+$ (rearr.), 1.1; 170, (M-SO$_3$)$^+$, 21.7; 165, $\text{F}_3\text{SCSO}_2^+$, 2.4; 151, (M-SO$_3$-F)$^+$, 10.0; 145, $\text{C}_2\text{F}_3\text{S}_2^+$, 1.6; 144, CHFO$_3\text{S}_2^+$, 1.5; 143, CFO$_3\text{S}_2^+$, 1.4; 127,
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SF$_5^+$, FSCSO$_2^+$, 5.3; 113, C$_2$F$_3$S$^+$, 1.6; 105, OSF$_3^+$, 3.2; 94,
C$_2$F$_2$S$^+$, 1.5; 91, C$_2$FOS$^+$, 4.4; 89, SF$_3^+$, 100.0; 83, SO$_2$F$^+$
(rearr.), 2.5; 82, CF$_2$S$^+$, 1.6; 70. SF$_2^+$, 10.9; 69, CF$_3^+$,
4.7; 67, FOS$^+$, 6.6; 66, CF$_2$O$^+$, 1.9; 64, SO$_2$+, 8.5; 63, CF$_5^+$,
4.9; 62, C$_2$F$_2^+$, 1.0; 60, COS$^+$, 1.2; 51, SF$^+$, 3.4.

Infrared spectrum (neat, KBr, cm$^{-1}$): 1415 ,s; 1379, vs;
1282, m; 1235, s; 1225, s; 1207, s; 1192, s; 1066, s; 969,
m; 874, vs; 857, vs; 763, s; 671, m; 661, s; 624, m; 606, w;
572, w; 535, w; 493, w; 472, s; 439, w.

Anal.calcd for C$_2$F$_6$O$_3$S$_2$: C:9.60; F:45.6; S:25.63.
Found: C:9.23; F:46.1; S:25.39.

**Reaction of 23 with HF (1):**

A 30 ml Kel-F vessel was charged via vacuum transfer
with 0.04 g of gaseous HF (2mmol) and then with 0.34 g (1.4
mmol) of 23. After keeping the vessel at room temperature
for 16 h it was cooled to -78°C and the excess HF was pumped
away through a cold trap (-196°C). A total of 0.31 g of
product was left behind. The $^{19}$F n.m.r. spectrum of this
material (CDCl$_3$) showed two major products, whose resonances
were coincident with those of sultone 1 and the compound
F$_5$SCH$_2$SO$_2$F. Sultone 1 accounted for 80% of the total
intensity of the integrated spectrum and the latter compound
for 18%. No 23 was left behind. The infrared spectrum of the
neat sample was identical with the one of 1.
Reaction of 23 with HCl (42):

Into a 40 ml Carius tube containing 1.08 g of 23 (4.32 mmol) 19.8 mmol of HCl (Matheson) was condensed. The vessel was heated to 68°C (92 h) and chilled (-196°C). Excess HCl was removed by pumping off the volatile constituents at -78°C through a cold trap (-196°C). Finally, the reaction product was collected by vacuum transfer at room temperature with occasional warming with a heat gun. The crude product (0.93 g) contained two impurities, which were probably 1 and SF₅CH₂SO₂F. They could be removed by trap-to-trap distillation (-30, -75, -100, -196°C), leaving 0.78 g of 42 in the -30°C trap. Yield = 63.3%.

Infrared spectrum (neat, KBr, cm⁻¹): 2992 (m); 1477 (vw); 1430 (s); 1317 (m); 1275 (m); 1214 (vs); 1178 (w); 1097 (m); 1076 (s); 958 (w); 886 (w); 853 (s); 835 (vs); 814 (s); 786 (s); 770 (msh); 743 (s); 658 (s); 642 (w); 610 (w); 589 (s); 561 (vw); 549 (w); 527 (m); 516 (m); 492 (vw); 478 (vw); 452 (w); 438 (w).

¹⁹F n.m.r. (neat sample, CCl₃F ext), ABCDXY system: Φₐ = 141.7 ppm (quartet, int. = 1.0); Φₐ = 120.0 ppm (m, int. = 2.0); Φ = 67.0 ppm (m, int. = 1.2); Φₓ(ᵧ) = -74.9 ppm (d, int. = 1.0); Φᵧ(x) = -78.9 ppm (d-m, int. = 1.0). Jₓᵧ = 100.8 Hz.

¹H n.m.r. (300 MHz, CDCl₃, Si(CH₃)₄): δ = 7.09 ppm (m).

¹³C n.m.r. spectrum (neat sample, CDCl₃ external): ¹H
coupled, $\delta = 113.4$ ppm (t, $J_{C^2F} = 294$ Hz), $\delta = 108.8$ ppm (d-m, $J_{C^1H} = 157.5$ Hz); $^1H$ decoupled, $\delta \beta = 113.4$ (t), $\delta \alpha = 108.8$ ppm (m). In the $^1H$-coupled spectrum the lines of the triplet show further signs of coupling but were not resolved.

Mass spectrum (negative ion, mass, species, >1%): 269, (M-O)$^-$, 3.0; 179, (M+H-Cl-O-3F)$^-$, 1.6; 177, (M-HCl-O-3F)$^-$, 2.9; 167, (M-HCl-SO$_2$F)$^-$ (rearr.), 13.5; 166, (M+H-HCl-F-COF$_2$)$^-$, 3.9; 165, (M-HCl-F-COF$_2$)$^-$, 100.0; 161,(M-HCl-3F-20)$^-$, 1.2; 144, CHFO$_3$S$_2^-$, 2.4; 143, (M-ClSF$_4$)$^-$, CFO$_3$S$_2^-$, 6.1; 142, (M-HCl-SF$_4$)$^-$, 43.1; 127, SF$_5^-$, FSCSO$_2^-$, 17.0; 123, (M-HCl-5F-20)$^-$, 2.0; 120, SF$_4^-$, 2.1; 108, SF$_4^-$, 5.5; 105, C$_2$HOS$_2^-$, 3.7; 101, CSF$_3^-$, 3.2; 99, CHF$_2$OS$^-$, 11.3; 96, CHFS$_2^-$ 4.2; 94, C$_2$F$_2$S$^-$, 11.8; 89, SF$_3^-$, 11.4; 85, CF$_3$O$^-$ (rearr.), 5.1; 83, SO$_2$F$^-$ (rearr.),CHF$_2$S$^-$, 100.0; 79, C$_2$HF$_2$O$^-$, CFOS$^-$, 1.4; 78, C$_2$F$_2$O$^-$, 45.2; 76, C$_2$HFS$^-$, CSO$_2^-$, 12.6; 70, SF$_2^-$, 2.2.

Anal.calcd for C$_2$HClF$_6$O$_3$S$_2$: C, 8.38; H, 0.35; Cl,12.37; F, 39.8; S, 22.37. Found: C, 8.51; H, 0.40; Cl, 12.51; F, 39.7; S, 22.18.

39 by Reaction of 23 with ClF:

Acid-washed apparatus was used, due to the easy rearrangeability of the product.

An oven-dried 75 ml steel bomb was charged with 2.95 g of CCl$_3$F (dried over P$_4$O$_{10}$), 0.31 g 23 (1.24 mmol) and 0.17
g (3.1 mmol) ClF. After attaining room temperature slowly, the bomb was kept there for 2 days. Fractional condensation in an acid-washed glass apparatus yielded (-196, -40 °C) 0.23 g of product in the -40°C trap. The $^{19}$F n.m.r. spectrum showed still 23 to be present. Both cold trap fractions were transferred back into the bomb, another 0.19 g ClF (3.5 mmol) were added and the bomb kept at room temperature for 6 days. Fractional condensation afforded 0.19 g in the -50°C trap, and after repeating the procedure with the -196°C fraction another 0.06 g were collected. The product was free of 23 but contained some small impurities (71, 53, 48, -70 ppm in $^{19}$F n.m.r. spectrum). Yield 0.25 g (66.2%).

The infrared spectrum was run quickly, because rearrangement to the acyl fluoride took place very readily on the KBr plates, as could be judged from the slow increase of the CO band (1855 cm$^{-1}$) $^{19}$F n.m.r. spectrum (neat sample, CCl$_3$F ext.): $\phi_{SF_5} = 62.8$ ppm (m), int. = 5.00; $\phi_{CF_2}$ (AB system): $\phi_A = -77.2$ ppm, d-p, int. = 1.04, $J_{ASF_4} = 3.6$ Hz (average); $\phi_B = -79.0$ Hz, $\approx$ d-p-m, irregular spacing, int. = 1.09. $J_{AB} = 94.3$ Hz.

Infrared spectrum (neat sample, KBr, cm$^{-1}$): 1481, m; 1461, m, sh; 1437, s-vs; 1351, vw; 1276, m-s; 1234, s, 1202, vs; 1151, m-s; 1129, m-s; 1090, s-vs; 1046, w-m; 970, m-s; 934, s; 890, vs; 836, s-vs; 819, s; 782, m-s; 738, vw; 692, m; 666, m-s; 624, m; 609, m-s; 588, m; 572, m; 551, m; 535, m-s; 503, w; 483, w; 454, m; 434, w-m; 414, w-m.
Mass spectrum (c.i., mass, species, % < 1): 205, (M-Cl-SO₂)⁺, 1.9; 191, (M-Cl-CF₂-CO)⁺ (rearr.), 10.6; 163, C₂H₂F₃OS₂⁺, 1.6; 127, SF₅⁺, 2.2; 107, C₂FO₂S⁺, 1.8; 91, C₂FOS⁺, 5.9; 89, SF₃⁺, 100.0; 85, CF₃O⁺ (rearr.), 3.0; 83, SO₂F⁺ (rearr.), 1.8; 81, C₂F₃⁺ (rearr.), 2.4; 79, CFOS⁺, 1.6; 73, C₂H₂OS⁺, 2.6; 71, ?, 5.2; 70, SF₂⁺, 1.9; 69, CF₃⁺ (rearr.), 4.5; 67, , FSO⁺, 2.5; 63, CFS⁺, 1.9; 61, CHOS⁺, 1.9.

Anal. calcd for C₂ClF₇O₃S₂: C, 7.89; F, 43.7; S, 21.05; Cl, 11.64. Found: C, 7.94; F, 44.2; S, 21.36; Cl, 11.51.

Synthesis of F₅SCHClCF₂Cl:

F₅SCH=CF₂ (52, 8.68 g, 45.7 mmol) and chlorine was irradiated with stirring with a 250 watts GE sunlamp in a 300 ml Pyrex-quartz vessel. The chlorine was added in portions with the following irradiation times: 1. 1.79 g (25.2 mmol), 15 cm distance, 4.5 hours (the liquid was still yellow). 2. 1.65 g (22.8 mmol), 19 hours, 10 cm distance. The product contained by ¹⁹F n.m.r. spectroscopy 52 (25.2 %), but also SF₅Cl (11.9%), S₂F₁₀ (6.7 %), F₅SCHClCF₂Cl (47.4 %) and an unknown compound (probably F₅SCCl₂CF₂Cl, 8.9 %) and more Cl₂ was added: 1. 0.92 g, 16 hours of irradiation, 2. 0.33 g, 4 hours. Only a trace of 52 was left. After reaction, the volatile materials were transferred off at -78°C and collected in a -196°C trap (5 hours, 2.12 g); S₂F₁₀
and SF₅Cl, SiF₄ were identified by i.r. spectroscopy. The remaining product (11.00 g) was by ¹⁹F n.m.r. spectroscopy 83.3 % F₅SCHClCF₂Cl, 4.3 % S₂F₁₀ and 12.4 % of the alleged F₅SCCl₂CF₂Cl (φₐ ≈ 63.5 ppm, φₖ = 42.7 ppm). Only S₂F₁₀ could be separated by rectification, but not the two SF₅ compounds, and the mixture was thus used without further purification to prepare F₅SCCl=CF₂. An analytically pure sample was prepared by preparative scale gas chromatography.

¹⁹F n.m.r. spectrum (neat sample, external CCl₃F):
AB₄XY, φₐ = 71.2 ppm, 9 lines, area = 1.00; φₖ = 59.3 ppm, d-m, area = 3.79; φₓ = -55.5 ppm, asymmetric d-p-d, area = 1.16; φₜ = -62.7 ppm, asymmetric d-d-p, area = 1.05. Jₐₖ = 148.5 Hz; Jₓₚ = 176.2 Hz; Jₚₜ = 12.8 Hz; Jₚₖ = 14.1 Hz; Jₓₜ = 2.3 Hz; Jₓₚ = 10.5 Hz.

Anal. calcd for C₂HClF₇S: C, 9.20; H, 0.39; Cl, 27.17; F, 51.0; S, 12.28. Found: C, 9.32; H, 0.41; Cl, 27.22; F, 51.1; S, 12.14.

F₅SCCl=CF₂:
The 83 % pure F₅SCHClCF₂Cl (10.25 g) was added by vacuum condensation into a 300 ml Carius tube that contained 15.12 g of powdered KOH. The tube was brought to room temperature and shaken for 5 minutes (warming was observed). After keeping it at room temperature for 15 minutes, 6.95 g of product was transferred out. It was combined with the product of a similar reaction to give 10.34 g of crude
product. Distillation gave two fractions, room temperature-44°C (4.76 g, 94% FSCl=CF2, 6% S2F10 by 19F n.m.r. spectroscopy), and 44-50°C (5.20 g) which was pure olefin.

**Reaction of FSCl=CF2 and SO3:**

A mixture of 2.03 g of SO3 (25.4 mmol) and 3.48 g of FSCl=CF2 (14.8 mmol) was heated in an 80 ml Carius tube to 160°C for 96 hours. The product was condensed out and was found to consist (integrated 19F n.m.r. intensity) to ≈ 50% of the olefin and 50% of a material with a multiplet at ≈ 50 ppm. The material was heated for 4 more days to 170-180°C, and its composition was found almost unchanged. More SO3 (1.70 g, 21.3 mmol) was added and after heating for 20 hours at 165°C all olefin was gone. The product was distilled, and three fractions were obtained, room temperature - 48°C, 48-65°C and 65-68°C, all were fuming liquids. They had all a different composition, and were not analyzed further. None contained an SF5-compound, however (i.r., 19F n.m.r. spectroscopy), and a sultone was apparently not formed (no CF2 resonance ("AB"), although the crude product showed some bands in the 19F resonance range for CF2, which were singlets and thus atypical for an AB system. The major product had a 19F resonance at 48 ppm (singlet).
Anhydrous methanol (0.05 g, 1.516 mmol) was added under swirling dropwise to 0.32 g of 23 (1.28 mmol) in a 25 ml pear-shaped flask. The methanol was allowed to flow down the side walls. Heat was liberated. The flask was stoppered and kept at room temperature for 140 minutes. An i.r. spectrum of the heavy yellow oil showed the presence of an ester band (1765 cm⁻¹, medium-strength). Separation from the alleged \( \text{CH}_3\text{OSF}_4\text{CH(SO}_2\text{F})\text{COOCH}_3 \) was effected by repeated trap-to-trap distillation, first with a -45°C and a -196°C trap, and then, using the material from the -45°C trap, with a -45°C and a -17°C trap. The material in the -17°C trap was subjected again to the same procedure, and after this third process, 0.21 g of a white solid was collected in the -17°C trap. The infrared spectrum showed that there was only a very weak ester band. Yield = 58.2 %. M.p. = 20.5-22.0 °C (sealed tube).

\(^1\text{H} \text{ n.m.r. spectrum (CDCl}_3, \text{ Si(CH}_3)_4, 300 MHz):} \delta_1 = 4.11 \text{ ppm, s (signs of shoulders), area = 3.00 (CH}_3); \delta_2 = 6.37 \text{ ppm, 8 lines with signs of further splitting, area = 0.96 (CH-CF}_2). \)

\(^1\text{H} \text{ n.m.r. spectrum (neat sample, external CCl}_3\text{F):} \)

ABCDXY system \( \phi_A = 68.4 \text{ ppm, d-d-d-d, area = 1.00; } \phi_B = 73.6 \text{ ppm, d-d-d-d-d, area = 1.00; } \phi_C = 55.3 \text{ ppm, d-d-d, area = 0.93; } \phi_D = 88.2 \text{ ppm, d-d-d-d, area = 1.00; } \phi_X = -80.3 \text{ ppm, d-m, area = 1.02; } \phi_Y = -74.3 \text{ ppm, d, area = 0.97. } J_{AB} = J_{AX} \)
= 25.1 Hz; \( J_{AC} = J_{BC} = 137.2 \) Hz; \( J_{AD} = J_{BD} = 115.7 \) Hz; \( J_{CD} = 149.6 \) Hz; \( J_{AY} = J_{CX} = J_{CY} = J_{DX} = J_{DY} = J_{CH^2} \approx J_{YH^2} = 0.0 \) Hz; \( J_{AH^2} = 10.58 \) Hz; \( J_{BH^2} = 7.82 \pm 0.39 \) Hz; \( J_{DH} = 20.21 \) Hz; \( J_{XY} = 102.7 \) Hz; \( J_{BX} = 14.47 \pm 0.63 \); \( J_{XH} \approx 6.8 \) Hz.

**I.r. spectrum (neat sample on KBr, cm\(^{-1}\)):** 2997, m; 2865, vw; 1459, m; 1424, vs; 1319, s; 1281, s; 1216, vs; 1103, s; 1081, vs; 1020, vs; 975, m; 830, vvs; 744, vs; 671, m; 659, s; 614, m-w; 596, m-w; 583, m-w; 574, w, sh; 556, vw; 529, s; 518, m-w, sh; 492, vw; 453, m.

**Mass spectrum (e.i. mass, species, % > 1):** 251, (M-CH\(_3\)O\(^+\)), 3.6; 231, (M-CH\(_3\)O-HF\(^+\)), 2.8; 227, (M-2F-H-O\(^+\)), 1.2; 201, (M-SO\(_3\))\(^+\), 9.9; 197, (M-SO\(_2\)-F-2H\(^+\)), 2.1; 183, (M-SO\(_3\)-F\(^+\)), 13.0; 182, (M-SO\(_3\)-F-H\(^+\)), 1.5; 173, (M-4F-20-H\(^+\)), 3.1; 172, (M-4F-20-2H\(^+\)), (M-5F-CH\(_3\))\(^+\), 1.6; 171, (M-4F-20-3H\(^+\)), (M-5F-CH\(_3\)-H\(^+\)), 47.6; 168, (M-6F\(^+\)), (M-SO\(_3\)-CH\(_3\)-F\(^+\)), 4.4; 163, (M-SO\(_3\)-2F-H\(^+\)), 3.5; 149, (M-SO\(_3\)-2F-CH\(_3\))\(^+\), 9.4; 145, (M-3F-SO\(_3\))\(^+\), 3.3; 144, (M-3F-H-SO\(_3\))\(^+\), (M+H-SF\(_4\)OCH\(_3\))\(^+\), 4.3; 139, SF\(_4\)OCH\(_3\), 9.8; 133, C\(_2\)HF\(_4\)S\(^+\), 1.2; 127, (M+H-4F-SO\(_3\))\(^+\), 3.1; 124, CO\(_3\)S\(_2\)\(^+\), 15.7; 109, CH\(_2\)O$_2$S\(^+\), 1.1; 105, C\(_3\)H\(_2\)OFOS\(^+\), C\(_2\)HO$_3$S\(^+\), 23.6; 101, CH$_3$OF$_2$\(^+\), 12.4; 94, C\(_2\)SF\(_2\)\(^+\), 4.7; 91, C\(_2\)FOS\(^+\), 3.4; 89, SF\(_3\)\(^+\), 100.0; 85, C\(_3\)HOS\(^+\), 1.5; 83, CHSF\(_2\)\(^+\), 2.6; 82, CSF\(_2\)\(^+\), CH$_3$OSF\(^+\), 2.8; 81, CH$_2$OSF\(^+\), 2.6; 80, CHOSF\(^+\), SO$_3$\(^+\), 3.5; 79, COSF\(^+\), 5.1; 75, CH$_3$OSC\(^+\), 2.3; 70, SF\(_2\)\(^+\), 6.1; 69, CF\(_3\)\(^+\) (rearr.), 2.2; 67, FO$_3$\(^+\), 21.0; 66, CF\(_2\)O\(^+\), 1.6; 65, SFCH\(_2\)\(^+\), 9.2; 64, SO$_2$\(^+\), CH$_3$OSH\(^+\), SFCH\(^+\), 12.8 (two signals); 63, CFS\(^+\), CH$_3$OS\(^+\), 9.1 (two signals); 60, COS\(^+\), 2.0.
Anal. calcd for C$_3$H$_4$F$_6$O$_4$S$_2$+: C, 12.77; H, 1.43; F, 40.4; S, 22.72. Found: C, 12.87; H, 1.33; F, 40.0; S, 22.38.

23 and (CH$_3$)$_3$COH:

The reaction was run either in a solvent (CCl$_3$F) or with the pure reactants; in neither case an ABCD spectrum ($^{19}$F) could be detected. It was not possible to separate the constituents by fractional condensation, and the apparent instability of F$_5$SCH(SO$_2$F)COOC(CH$_3$)$_3$ excluded other methods. Inferences are thus made from $^{19}$F n.m.r. and i.r. data.

To 0.067 g of (CH$_3$)$_3$COH (0.91 mmol, Mallinckrodt, reagent grade) in a 50 ml pear-shaped flask with a Kontes Teflon valve connector, 0.22 g of 23 (0.88 mmol) was vacuum transferred. The bulb was allowed to attain room temperature slowly in a cold Dewar container (initially -196°C), and swirled when the constituents started to melt. After complete admixture, the vessel was kept at room temperature for one more hour. An i.r. spectrum of this crude material showed a strong ester band at 1760 cm$^{-1}$, and was otherwise also very similar to the i.r. spectrum of F$_5$SCH(SO$_2$F)COOC(CH$_3$)$_3$. The $^{19}$F n.m.r. spectrum (in CCl$_3$F) showed bands corresponding to F$_5$SCH(SO$_2$F)COOC(CH$_3$)$_3$ and probably 1 (F$_5$SCHCF$_2$OSO$_2$). When the solution was kept at room temperature for several days, the bands corresponding to 1 diminished, while bands corresponding to F$_5$SCH(SO$_2$F)COOC(CH$_3$)$_3$ (major product) and most likely
F₅SCH₂SO₂F) increased.

F₅SCH(SO₂F)COOC(CH₃)₃ (21d):

To 0.10 g of tertiary butanol (1.35 mmol, Mallinckrodt, reagent grade), in a 50 ml pear-shaped flask with a Kontes Teflon valve connector, 0.36 g (1.36 mmol) of 21 in 2.5 ml CCl₃F (Dupont) was vacuum transferred. The flask was swirled while it warmed up, and when all (CH₃)₃COH had dissolved, the mixture was cooled again by dipping the flask into liquid nitrogen, and was then allowed to warm up slowly in a cold Dewar container. The material that was obtained was used as such, as attempts to isolate it earlier by distillation resulted in the formation of F₅SCH₂SO₂F; trap-to-trap distillation was also not capable to furnish pure ester. In this case, an originally colorless product was collected that assumed a purple color within hours, and left finally a purple residue upon evaporation.

1H n.m.r. spectrum (400 MHz, CCl₃F, Si(CH₃)₄): δ = 1.58 ppm, s, int. = 9.94 (COOC(CH₃)₃); δ = 5.88 ppm, m, int. = 1.00 (SF₅CH); an impurity at δ = 1.29 and δ = 2.06 was identified as (CH₃)₃COH by adding a drop of an authentic sample to to original sample and which led to a large proportional increase in these two bands. (CH₃)₃COH accounted for 17.3 % of the t-butyl concentration in the crude ester.

19F n.m.r. spectrum (CCl₃F): ϕSF₅ = 71.7 ppm, narrow
multiplet, area = 5.00; \( \phi_{SO_2F} = 59.8 \) ppm, m. area = 0.93.

Infrared spectrum (neat sample, on KBr, cm\(^{-1}\)) of the crude product: 2990, m; 2954, w,sh; 1760, s-vs; 1481, w; 1442, s-vs; 1401, w; 1377, m-s; 1299, s; 1266, m-s; 1222, m-s; 1149, s-vs; 1040, vw; 965, w; 879, vs; 835, s; 801, m-s; 788, m-s; 773, m-s; 698, w-m,br; 664, w; 643, w,br; 612, m; 571, s-vs; 490, w-m; 458, w; 452, w-m; 446, w-m.

G.c. mass spectrum (e.i., 70 eV, mass, species, % > 1): 251, (m-(CH\(_3\))\(_3\)CO\(^+\)), 0.4; 231, (M-(CH\(_3\))\(_3\)CO-HF\(^+\)), 2.7; 127, SF\(_5^+\), 15.4; 105, C\(_2\)HO\(_3\)S\(^+\), 1.1; 97, CFO\(_2\)S\(^+\), C\(_6\)H\(_9\)O\(^+\), 1.0; 91, C\(_2\)FOS\(^+\), 1.1; 89, SF\(_3^+\), 17.3; 70, SF\(_2^+\), 2.0; 67, FSO\(^+\), 11.5; 64, SO\(_2^+\), 3.5; 60, COS\(^+\), 3.3; 59, C\(_3\)H\(_7\)O\(^+\), 100.0; 58, C\(_3\)H\(_6\)O\(^+\), 6.2; 57, C\(_3\)H\(_5\)O\(^+\), 68.6; C\(_3\)H\(_4\)O\(^+\), 21.0; 55, C\(_3\)H\(_3\)O\(^+\); 53, C\(_3\)HO\(^+\); 51, SF\(^+\), 1.9.

Reaction of 23 and CsF:

Finely ground cesium fluoride (2.80g, 18.4 mmol) was dried (3d, 115°C, high vacuum) in a 20 ml Pyrex glass reaction vessel, equipped with a Kontes Teflon valve. Then 0.19 g of 5 (0.76 mmol) were added by vacuum transfer (~196°C) and the vessel was maintained at 40°C (19h). The weight gain of the vessel was 0.16 g, after the volatile material had been removed by vacuum transfer. An infrared spectrum of the solid showed vibrational bands that were attributable to the complex Cs\(^+\)[F\(_5\)SC(SO\(_2\)F)COF\(^-\)]. A \( ^{19F} \) n.m.r. spectrum of the acetone extract, which was obtained
as described below, was also consistent with only the presence of the cesium compound. The reaction mixture was rinsed several times with acetone (4x3ml), the acetone extract was filtered, the filter residue was washed with several small portions of acetone, and the solution was brought to dryness, leaving 0.15g of the cesium salt. Yield = 49.1%.

\[ [\text{NH}_4]^+ [\text{F}_5\text{SC(SO}_2\text{F)COOCH(CH}_3\text{)}_2]^-(34) \]

Ammonia was bubbled into a solution of 0.55g of F\textsubscript{5}S\textsubscript{C}(SO\textsubscript{2}F)COOCH(CH\textsubscript{3})\textsubscript{2} (1.8 mmol) in 14 ml of anhydrous ether, cooled to -25°C, in a 25 ml round bottom flask, fitted with a Claisen-head, Drierite tube and a gas inlet tube. The temperature was allowed to rise continuously to +15°C, when a small amount of precipitation was observed. \textsuperscript{19}F n.m.r. spectroscopy showed the starting ester to be gone, but attempts to isolate the product failed. In the solid state the compound decomposed rather quickly upon reaching room-temperature, but was stable in solution. An i.r. spectrum was obtained by allowing the ether to evaporate on a KBr disk and immediately recording the spectrum.

\textsuperscript{19}F n.m.r. (in ether, CCl\textsubscript{3}F ext.): AB\textsubscript{4}X \( \phi = 96.1 \text{ ppm}, 9 \text{ lines}, \text{ int.}= 1.07, (F_A); \phi = 82.0 \text{ ppm}, \text{ d-m}, \text{ int.} = 4.00 (F_B); \phi = 65.8 \text{ ppm}, \text{ p}, \text{ int.} = 1.02 (F_X, \text{ SO}_2\text{F}); J_{AB} = 151.3 \text{ Hz}; J_{BX} = 10.9 \text{ Hz}.

I.r. (neat sample, KBr, cm\textsuperscript{-1}): 3648,m; 3585,m; 3400-
2900, s, br, peaks at 3037, 2995; 1743, w; 1636, m, sh; 1589, s; 1543, s, sh; 1493, m-s; 1453, s; 1436, s, sh; 1436, m-s, sh; 1399, s; 1366, vs; 1292, s-vs; 1191, s; 1181, s; 1145, m; 1100, s; 1047, m; 910, m-s; 868, s, sh; 850, vs; 810, vs; 760, vs; 710, m-s; 668, s; 620, s; 582, s; 573, m-s; 494, m; 486, m; 462, m; 424, m-s; 413, m-s.

$\text{F}_5\text{SCH(SO}_2\text{NH}_2\text{)COOCH(CH}_3\text{)_2} \quad (35)$:

Gaseous ammonia was introduced slowly into a stirred solution of $\text{F}_5\text{SCH(SO}_2\text{F)COOCH(CH}_3\text{)_2}$ (2.4 g, 7.7 mmol) in 5 ml dioxane in a 50 ml round-bottomed flask. Initially, a temperature-rise was observed, and the rate of NH$_3$-introduction was kept so low that the temperature stayed below 50°C (charring is possible). When the temperature of the mixture dropped, NH$_3$ addition was stopped, 2ml of water as added and most of the solvent was boiled off. The product contained still the starting ester and the treatment with NH$_3$ was repeated. The volume of the solution was further diminished and the product was obtained by twice recrystallizing it from dilute HCl ($H_2O: HCl = 5:1$). Yield: 1.2 g (dried over $P_2O_5$ in the desiccator), 50.5 %, cream-colored amorphous mass. M.p. 244-246°C, darkens; 254°C, solidifies, forms shiny flakes; 270°C, melts again, 280°C, darkens.

$^1H$ n.m.r. ($D_2O$, 300 MHz, Si(CH$_3$)$_4$ int.): $\delta_1 = 1.30$ ppm, d, int = 6.00 (CH(CH$_3$)$_2$), $\delta_2 = 5.13$ ppm, septet, int. =
0.97 \((\text{CH(\text{CH}_3)}_2); \delta_3 = 5.88 \text{ ppm, p, int.} = 0.77 \) (\text{F}_5\text{SCH}); \text{NH}_2
is not observed; \(J_{12} = 6.30 \text{ Hz; } J_{HB} = 5.85 \text{ Hz.} \)

\(^{19}\text{F} \text{ n.m.r. (H}_2\text{O, CCl}_3\text{F ext.)}: \text{AB}_4 \phi = 79.8 \text{ ppm, 9 lines, int.} = 1.0 \) (\(\text{F}_A\)); \(\phi = 69.2 \text{ ppm, d-m, int.} = 3.6 \) (\(\text{F}_B\)); \(J_{AB} = 151.7 \text{ Hz.} \)

\(\text{I.r. (KBr, neat, cm}^{-1}): 3184, \text{ m-s; } 3087, \text{ m-s; } 2986, \text{ m-s; } 2880, \text{ m; } 1733, \text{ s; } 1684, \text{ m,sh; } 1460, \text{ m-s; } 1440, \text{ m-s; } 1393, \text{ m; } 1380, \text{ m; } 1353, \text{ m; } 1310, \text{ m-s; } 1265, \text{ s; } 1211, \text{ s-vs; } 1181, \text{ s,sh; } 1149, \text{ m,sh; } 1104, \text{ m-s; } 1051, \text{ s; } 945, \text{ vw; } 904, \text{ s; } 871, \text{ s; } 859, \text{ vs; } 810, \text{ s; } 783, \text{ m-s; } 741, \text{ m-s; } 682, \text{ w; } 648, \text{ m-s; } 610, \text{ w; } 592, \text{ m-s; } 572, \text{ m; } 550, \text{ m; } 521, \text{ m; } 483, \text{ m-w; } 417, \text{ w.} \)

Mass spectrum (negative ionization, mass, species, \% for int. > 1%): 307, \(M^-\), 0.14; 249, \((\text{M-C}_3\text{H}_7\text{O})^-\), 2.3; 180, \((\text{M-SF}_5)^-\), 1.2; 139, \(\text{C}_5\text{HNO}_2\text{S}^-\), 2.1; 129, \(\text{C}_4\text{H}_3\text{SO}^-\), 4.0; 127, \(\text{SF}_5^-\), 59.7; 124, \(\text{C}_2\text{HFO}_3\text{S}^-\), \(\text{C}_2\text{HOS}_2^-\), \(\text{CH}_2\text{NO}_2\text{S}_2^-\), 7.1; 123, \(\text{C}_2\text{FO}_3\text{S}^-\), \(\text{C}_2\text{FOS}_2^-\), \(\text{CHNO}_2\text{S}_2^-\), 2.4; 122, \(\text{CNO}_2\text{S}_2^-\), 2.4; 118, \(\text{C}_2\text{NO}_3\text{S}^-\), 2.2; 113, \(\text{C}_2\text{F}_3\text{S}^-\), 2.2; 110, \(\text{C}_2\text{F}_2\text{OS}^-\), 4.5; 108, \(\text{SF}_4^-\), 100.0; 105, \((\text{CCHSO}_2\text{NH}_2)^-\), 1.6; 98, \((\text{CCOOC}_3\text{H}_6)^-\), 1.8; 96, \(\text{CHFS}_2^-\), 8.1; 95, \(\text{CFS}_2^-\), 2.5; 94, \(\text{CH}_2\text{SO}_2\text{NH}_2^-\), \(\text{CC(O)OC}_3\text{H}_2^-\) two fragments), 9.0, 6.9; 93, \(\text{CHSO}_2\text{NH}_2^-\), 23.6; 89, \(\text{SF}_3^-\), 15.4; 85, \(\text{C}_4\text{H}_5\text{O}_2^-\), 1.0; 83, \(\text{CHSF}_2^-\), 34.8; 82, \(\text{CSF}_2^-\), 7.4; 80, \(\text{SO}_2\text{NH}_2^-\), 2.9;

\text{Anal. calcd for } \text{C}_5\text{H}_{10}\text{F}_5\text{NO}_4\text{S}_2: C, 19.54; H, 3.28; F, 30.9; N, 4.56; S, 20.87. \text{Found: C, 19.41; H, 3.37; F, 30.2; N, 4.44; S, 20.62.}
\((\text{C}_2\text{H}_5)_3\text{NH})^+\left[\text{F}_5\text{SC(SO}_2\text{F)}\text{COOCH}_3\right]^- \quad (32b): \]

To 0.3385 g of \(\text{F}_5\text{SC(SO}_2\text{F)}\text{COOCH}_3\) (1.2 mmol) in 2.0 ml anhydrous ether in a 10 ml flask, fitted with a Claisen-head, Drierite tube and dropping funnel, 0.15 g \(\text{N(C}_2\text{H}_5)_3\) (1.49 mmol) in 2.0 ml anhydrous ether was added dropwise during 30 seconds with swirling at \(-45^\circ\text{C}\). An immediate separation of a lower pale yellow layer was observed. The cold-bath was removed and the mixture was allowed to attain room-temperature slowly with occasional swirling. After removal of the solvent in vacuo 0.4288 g (1.12 mmol) or 93 % of an amorphous, slightly yellow solid was obtained, which was analytically pure.

\(^1\text{H n.m.r. (CDCl}_3, \text{Si(CH}_3)_4 \text{ ext.}): \delta_1 = 1.79 \text{ ppm, t, int. } = 10.1 \ (\text{HN(CH}_2\text{CH}_3)_3^+)\; \delta_2 = 3.59 \text{ ppm, quart.}, \text{int. } = 6.3 \ (\text{HN(CH}_2\text{CH}_3)_3^+); \ \delta_3 = 4.04 \text{ ppm, s}; \ \text{int. } = 3.0 \ (\text{COOCH}_3); \ \delta_4 = 7.46 \text{ ppm, s, broad, int. } = 1.0 \ (\text{HN(C}_2\text{H}_5)_3^+); \ J_{\text{H-H}} = 7.40 \text{ Hz (ethyl)}.\]

\(^19\text{F n.m.r. (CDCl}_3, \text{CCl}_3\text{F ext.}): \text{AB}_4\ \phi = 95.3 \text{ ppm, 9 lines, int. } = 1.0 \ (\text{F}_\text{A}); \ \phi = 82.1 \text{ ppm, d-m, int. } = 4.1 \ (\text{F}_\text{B}); \ \phi = 67.0 \text{ ppm, p, int. = 1.0 (F}_X, \text{SO}_2\text{F); J}_{\text{AB}} = 154.2 \text{ Hz; J}_{\text{BX}} = 10.2 \text{ Hz.}\]

I.r. (neat sample, KBr, cm\(^{-1}\)): 3093, m-s; 2995, m; 2952, m; 2875, w-m; 2748, w; 2495, vw; 1640, s-vs; 1523, vw; 1474, m-s; 1462, m; 1442, s; 1423, m; 1394, m-s; 1367, s,sh; 1355, s; 1306, vs; 1289, vs; 1179, s; 1093, m; 1052, m; 1020, w-m; 939, w-m; 900, vw; 864, s; 839, vs; 805, vs; 757,
Mass spectrum (electron impact, 70 eV, mass, species, >2): 251, (M-N(C2H5)3-CH3O)+, 11.7; 231, (M-N(C2H5)3-HF-CH3O)+, 5.1; 168, (M-N(C2H5)3-SO2F-CH3O)+, 4.5; 155, (M-N(C2H5)3-SF5)+, 6.5; 149, (M-N(C2H5)3-SO2F-CH3O-F)+, 2.3; 143, (M-N(C2H5)3-4F-CH3O-20)+, 5.2; 139, (M-N(C2H5)3-SO2F-COOCH3-H)+, 3.6; 135, (M-N(C2H5)3-SF5-HF)+, 3.7; 129, (M-N(C2H5)2C2H4-SF5-COOCH3)+, 2.7; 127, SF5+, 52.1; 125, (M-N(C2H5)2C2H4-SF5-CH3O)+, 19.5; 124, (M-N(C2H5)3-SF5-CH3O)+, 3.9; 122, (M-N(C2H5)2C2H4-SO2F-F-COOCH3)+, 5.6; 105, (M-N(C2H5)3-SF5-CH3O-F)+, (C3H2FOS)+ (two fragments, 8.8, 10.3; 102, HN(C2H5)3+, 2.5; 101, N(C2H5)3+, 29.1; 100, N(C2H5)2C2H4+, 11.5; 91, C2FOS+, 2.4; 89, SF3+, 48.3; 87, HN(C2H5)2CH2+, 11.3; 86, N(C2H5)2CH2+, 100.0; 79, CFOS+, 4.0; 72, N(C2H5)2+, CHCOOCH3+, 7.5; 71, N(C2H5)C2H4+, CCOOCH3+, 2.8; 70, N(C2H5)C2H3+, CCOOCH2+, SF2+ (two fragments), 5.7, 3.7; 69, N(C2H5)C2H2+, CCOOCH+, 3.0; 67, FSO+, C4H5N+, 35.1; 63, CFS+, 3.3; 62, C4N+, 2.3.

Anal. calcd for C9H19F6N04S2: C, 28.19; H, 5.00; F, 29.7; N, 3.65; S, 16.73. Found: C, 28.40; H, 4.89; F, 30.1; N, 3.78; S, 16.43.

\[(\text{C}_2\text{H}_5)_3\text{NH}^+ [\text{F}_5\text{SC}(\text{SO}_2\text{F})\text{COOCH(CH}_3)\text{)_2}^- (32a)\]:

To 2.11 g of F5SCH(SO2F)COOCH(CH3)2 (6.8 mmol), dissolved in 3.0 ml of anhydrous ether, in a 25 ml round-
bottom flask, equipped with a magnetic stirring bar, Claisen-head, dropping funnel and Drierite tube, 0.80 g of triethyl amine, dissolved in 4 ml of anhydrous ether, was added dropwise with stirring at -60° to -50°C (acetone/dry ice). At -45°C bath-temperature two layers were observed in the reaction flask. Stirring was continued for two minutes, the cold-bath was removed and the solvent was pumped away through a cold trap (-196°C), leaving 2.80 g of an analytically pure amorphous white solid. Yield = 100%. M.p. 94-97°C. The solid could be handled in air without difficulty and formed large irregular crystals upon recrystallization from chloroform.

1H n.m.r. (CD3COCD3, Si(CH3)4 ext.): δ1 = 6.45 ppm, s, broad, int. = 0.93 (NH); δ2 = 4.85 ppm, sept., int. = 0.71 (CH(CH3)2); δ3 = 3.23 ppm, quart., int. = 6.00 (N(CH2CH3)3); δ4 = 1.24 ppm, t, int. ≈ 8.6 (overlap, N(CH2CH3)3); δ5 = 1.09, d, int. ≈ 6.6 (overlap, CH(CH3)2). J25 = 6.36 Hz; J34 = 7.14 Hz.

19F n.m.r. (CD3COCD3, CCl3F ext.): AB4X, φ = 97.0 ppm, 9 lines, int. = 1.00 (FA); φ = 82.1 ppm, d-m, int. = 4.25 (FB); φ = 66.3 ppm, p, int. = 1.00 (FX, SO2F); JAB = 154.0 Hz; JBX = 10.6 Hz.

Infrared spectrum (neat, KBr, cm⁻¹): 3060, vw; 2992, vw, 2941, vw; 2754, vw; 2677, vw; 2498, vw; 1615, s; 1467, m; 1401, m; 1380, s; 1291, s; 1194, s; 1179, s; 1144, m; 1111, m-s; 1093, s; 1041, s; 910, m; 846, vs; 836, vs, sh;
Mass spectrum (chemical ionization, mass, species, > 2 %): 311, (M-N(C₂H₅)₂C₂H₄)⁺, 4.3; 269, C₂H₃F₆O₄S₂⁺, 9.8; 251, (M-N(C₂H₅)₃-C₃H₇O)⁺, 3.3; 219, CF₃O₃S₂⁺ (rearr.), 3.6; 209, (M-N(C₂H₅)₃-2F-HF-C₃H₇)⁺, 3.1; 205, CH₂F₅O₂S₂⁺, 6.4; 143, (M-N(C₂H₅)₃- SO₂F-2HF-CO₂⁺); 127, SF₅⁺, 12.4; 125, C₂H₂FO₃S⁺, 8.9; 105, C₄H₆FS⁺ (rearr.), 3.1; 103, C₄H₄FS⁺ (rearr.), 3.3; 102, NH(C₂H₅)₃⁺, 59.5; 101, N(C₂H₅)₃⁺, 93.6; 100, (N(C₂H₅)₃-H)⁺, 75.4; 99, (N(C₂H₅)₃-2H)⁺, 11.5; 91, C₂FOS⁺, 3.4; 89, SF₃⁺, 76.0; 87, NH(C₂H₅)₂CH₂⁺, 11.4; 86, N(C₂H₅)₂CH₂⁺, 100.0; 79, CF₃S⁺, 4.3; 74, C₅N⁺, (NH₂(C₂H₅)₂)⁺, 4.6; 72, N(C₂H₅)₂⁺, 11.7; 71, (N(C₂H₅)₂-H)⁺; 2.2; 70, SF₂⁺, (N(C₂H₅)₂-2H)⁺ (two fragments), 4.2, 4.4; 67, FSO⁺, 11.8; 65, CH₂FS⁺, 21.4; 61, C₃H₇OH₂⁺, 3.5; 59, C₃H₇O⁺, 53.7; 58, C₃H₆O⁺, 31.6; 57, C₃H₅O⁺, 4.2; 56, C₃H₄O⁺, 7.4; 55, C₃H₃O⁺.

Anal. calcd for C₁₁H₂₃F₆N₀₄S₂: C, 32.11; H, 5.64; F, 27.7; N, 3.40; S, 15.59. Found: C, 32.04; H, 5.45; F, 27.5; N, 3.31; S, 15.80.
minutes, 0.50 g of this oil was pipetted off. Its i.r. spectrum coincided with the spectrum of $\text{F}_5\text{SCH(SO}_2\text{F)}\text{COOCH(CH}_3\text{)}_2$, except for a few very weak bands from 2500-2700 cm$^{-1}$ and at 1616 cm$^{-1}$. Yield: 94.7% 

$\text{Na}^+ [\text{F}_5\text{SC(SO}_2\text{F)}\text{COOCH(CH}_3\text{)}_2] (33)$, general procedure:

Freshly prepared isopropoxide was stirred in anhydrous ether in a 50 ml flask, equipped with a magnetic stirring bar and fitted with a Claisen head, Drierite tube and a rubber septum. The ester, dissolved in 1-2 ml of ether was added with a syringe through the septum. The isopropoxide dissolved in a vigorous reaction, causing the ether to boil. The rate of addition of the ester was reduced, when necessary, in order to moderate the reaction. When addition was complete, the yellowish-brown solution was stirred for another 30-60 minutes at room temperature. The amount of ether used was 5-10 ml for 4-7 mmol of isopropoxide; the ester was used in $\approx$0.7-0.9 fold quantity.

The salt could be prepared by the same method in isopropanol. When the solvent was pumped away a brownish glassy mass remained, which upon re-dissolution did not show the spectrum of the pure anion anymore ($^{19}\text{F n.m.r.}$), nor did it show only one CO band (i.r.). It was also not possible to purify it by successive dissolution (in ether) and precipitation by adding hexane to the solution. In this case a white amorphous mass was obtained, which showed two C=O
bands in the i.r. spectrum: 1738 (m, broad) and 1637 cm\(^{-1}\) (s) and was very moisture-sensitive.

\(^{19}\text{F} n.m.r.\) (in isopropanol, CCl\(_3\)F ext.): AB\(_4\)X \(\phi = 95.3\) ppm, 9 lines, int. = 1.0 (F\(_A\)); \(\phi = 82.8\) ppm, d-m, int. = 4.0 (F\(_B\)); \(\phi = 67.3\) ppm, p, int. = 1.0, (F\(_X\), SO\(_2\)F); \(J_{AB} = 150.4\) Hz; \(J_{BX} = 10.6\) Hz. In ether: \(\phi_A = 93.5\) ppm, \(\phi_B = 81.7\) ppm, \(\phi_X = 66.2\) ppm.

Na\(^+\)F\(_5\)SC(SO\(_2\)F)COOCH(CH\(_3\))\(_2\)\(^-\) (33), stability:

The sodium salt was prepared as described above from 0.33 g (4.0 mmol) of sodium isopropoxide and 1.21 g (3.9 mmol) of F\(_5\)SCH(SO\(_2\)F)COOCH(CH\(_3\))\(_2\) in 5.0 ml of ether and stirred at room temperature. \(^{19}\text{F} n.m.r.\) spectroscopy showed that the anion signal, which was initially the only observed resonance, disappeared slowly with the reversion to the signal attributable to the starting ester. After six days this process was complete, with the formation of a white precipitate. Distillation (room temperature - 60 °C) gave one uniform fraction (0.64 g, 52.3 %) which was identified by i.r. and \(^{19}\text{F} n.m.r.\) spectroscopy as the starting ester.

F\(_5\)SCCl(SO\(_2\)F)COOCH(CH\(_3\))\(_2\) (37):

Na\(^+\)F\(_5\)SC(SO\(_2\)F)COOCH(CH\(_3\))\(_2\)\(^-\) was prepared freshly from sodium isopropoxide (0.5745 g, 7.0 mmol) and F\(_5\)SCH(SO\(_2\)F)COOCH(CH\(_3\))\(_2\) (1.845 g, 6.0 mmol) in a 50 ml pear-shaped flask as described above. The septum on the Claisen-
head was replaced by a gas inlet tube that reached just above the liquid level and chlorine gas was introduced at a slow rate. A white precipitate formed instantly; chlorine was added slowly until a greenish-brown color persisted. A distillation unit was then attached to the flask in place of the Claisen head, the ether was removed at lowered temperature and frozen out in a \(-196^\circ\text{C}\) trap. Distillation at 2.5 torr afforded three fractions: room-temperature \(- 48.5 \, ^\circ\text{C}\) (0.360g), 48.5-49 °C (0.962 g and 48 \(- 28 \, ^\circ\text{C}\) (0.139g). The last two fractions were 97.7 and 97.0 % pure (gas chromatography, QF1, 1m, 130 °C), whereas fraction 1 was 82 % pure chlorinated ester. The impurity was not the starting ester. Yield \(\approx\) 54 %. An analytically pure sample was obtained by preparative gas chromatography. The solid residue in the flask (0.6323 g) was somewhat more than the expected NaCl (0.3482 g) and excess NaOCH(CH\(_3\))\(_2\) (0.0865 g).

\(\text{\begin{tabular}{l}
1H n.m.r. (CDCl}_3, \text{Si(CH}_3)_4 \text{ ext.}): \delta = 1.85\text{ppm (CH}_3, \\
\text{int. = 6.3)}; \delta = 5.80\text{ppm (CH, int. = 1.0)}, J_{H-H} = 6.47 \text{Hz.}
\end{tabular}}\)

\(\text{\begin{tabular}{l}
1^9\text{F n.m.r. (CDCl}_3, \text{CCl}_3\text{F, ext.}): AB}_4\text{X : } \phi_A = 68.8 \\
\text{ppm, 9 lines (F}_\text{ax, int. = 0.95; } \phi_B = 63.0 \text{ppm } \approx d-m \text{ (Feq,} \\
\text{int. = 4.25); } \phi_X = 49.8 \text{ ppm, p (SO}_2\text{F, int. = 1.0); } J_{AB} = \\
150.4 \text{Hz; } J_{AX} = 11.6 \text{Hz.}
\end{tabular}}\)

\(\text{\begin{tabular}{l}
I.r. (cm}^{-1}): 2993, w; 2945, vw; 1778, s;1766 ms; 1470, \\
w; 1444, s; 1393, w; 1391, w-m; 1347, vw; 1256, s; 1223, s; \\
1098, s; 1035, vw; 981, vw; 916, s; 887, vs; 877, vs; 840, \\
s; 807, s; 784, m; 737, m-w; 689, w; 673, vw; 659, w;611, m;
\end{tabular}}\)
Mass spectrum (e.i., 70 eV, mass, species, %) only the $^{35}\text{Cl}$-signals will be given: 343, (M-H $^{35}\text{Cl}$)$^+$, 1.0; 329, (M-H-CH$_3$)$^+$, 19.9; 285, (M-OCHCH$_3$)$_2$)$^+$, 26.5; 231, (M-Cl-OC$_3$H$_7$-F)$^+$, 2.1; 217, (M-Cl-4F-O)$^+$, 1.2; 209, (M-Cl-SO$_2$F-O-H)$^+$, 2.4; 203, (M-C$_3$H$_7$-F-Cl-CO$_2$)$^+$, 2.9; 202, (M-Cl-4F-O-CH$_3$)$^+$, 2.1; 177, (M-SF$_5$-HF-O-6H)$^+$, 9.5; 159, (M-SF$_5$-C$_3$H$_6$)$^+$, 3.8; 158, (M-SF$_5$-OC$_3$H$_7$)$^+$, 5.2; 129, (M-SF$_5$-HF-O-5H-C-Cl = C$_4$H$_9$O$_3$S$^+$, 1.3; 127, SF$_5^+$, 19.1; 96, C$_4$OS$^+$, 1.6; 94, C$_2$F$_2$S$^+$, 6.8; 91, C$_2$FOS$^+$; 89, SF$_3^+$, 100.0; 83, SO$_2$F$^+$, 2.7; 79, C$_5$H$_3$O$^+$, 1.8; 77, C$_5$HO$^+$, 1.8; 76, C$_5$O$^+$, 3.1; 75, C$_2$FS$^+$, 2.6; 70, SF$_2^+$, 8.2; 69, C$_4$H$_5$O$^+$, C$_3$H$_2$O$^+$, 2.4, 1.1 (two fragments); 68, C$_4$H$_4$O$^+$, C$_3$O$_2^+$, 3.8; 67, P$^+$O, 39.1; 66, C$_4$H$_2$O$^+$, 6.1; 65, C$_4$HO$^+$, 1.8; 64, SO$_2^+$, C$_4$O$^+$, 12.1; 63, CFS$^+$, 4.0; 61, CHOS$^+$, 8.1; 59, C$_3$H$_7$O$^+$, 13.3; 58, C$_3$H$_6$O$^+$, 2.4; 51, SF$^+$, 2.7.

_{Anal. calcd} for C$_5$H$_7$ClF$_6$O$_4$S$_2$: C, 17.42; H, 2.05; Cl, 10.28; F, 33.1; S, 18.60. Found: C, 17.56; H, 2.24; Cl, 10.50; F, 33.2; S, 18.55.

$^{5}$SCBr(SO$_2$F)COOCH(CH$_3$)$_2$ (38):

The sodium salt of the ester was prepared as described above from 0.517 g NaOCH(CH$_3$)$_2$ (6.3 mmol) in 10.0 ml ether and 1.37 g $^{5}$SC(SO$_2$F)COOCH(CH$_3$)$_2$ (4.4 mmol). Bromine was added dropwise with a syringe through the septum until a yellow color persisted. A white precipitate was immediately formed upon addition of Br$_2$. An n.m.r. spectrum ($^{19}$F) showed
that only one fluoro compound was present. The ether was pumped off from the reaction flask through a cold-trap and distillation at 3 mm Hg afforded one fraction, r.t. - 83°C (1.18 g), a slightly brownish heavy liquid. It was a mixture (19F n.m.r.) of probably F5SCH(SO2F)COOCH(CH3)2 (12 parts) and the alleged F5SCBr(SO2F)COOCH(CH3)2 (88 parts). No separation was achieved by repeated distillation.

19F n.m.r. spectrum: degenerate AB₄ spectrum φ = 64.9 ppm, m, int. = 5.18 (SF₅); φ = 49.0 ppm, p, int. = 1.00 (SO₂F); Jₓₓ = 12.3 Hz.

I.r. spectrum of the impure product (neat, KBr, cm⁻¹):
2995, m, 2945, w; 2889, vw; 1764, vs; 1470, m; 1441, vs; 1393, m; 1361, m; 1334, w; 1296, m; 1254, vs; 1222, vs; 1185, m; 1148, m; 1097, vs; 972, w; 889, vs; 860, vs; 838, s; 821, s-vs; 807, s-vs; 793, s-vs; sh; 785, s,sh; 742, w; 688, m; 648, m; 611, s; 574, vs; 497, w-m; 463, w-m; 415, w-m.

Further reactions of Na⁺[F₅SC(SO₂F)COOCH(CH₃)₂]⁻:

Reactions with other halides were tried by the same method as above. The reaction with acetyl chloride was performed with the white residue that is obtained from the salt solution after removal the solvent in vacuo, but in all cases only the starting ester could be recovered.

Na⁺ Anion⁻ + BrCN : 39.4% recovery

" + ClCOOCH₃: 49.9% "$
Reactions with CH2Br2 were also tested in an n.m.r. tube experiment, which indicated that only the original ester was formed. In all cases, however, a white precipitate was formed. With acetyl chloride small amounts of other products with fluorine n.m.r. signals in the SF region were also formed, but could not be identified.

F5SCH(SO2F)CONHCONH2 (31):

A solution of F5SCHCF2OSO2 (2.09g, 7.8 mmol) in 15 ml of dry ether was added dropwise to a suspension of urea (0.46g, 7.7 mmol) in 50 ml of anhydrous ether at 0°C. The solution turned quickly cloudy and a sticky precipitate formed. It was stirred at room-temperature for 2h, and then kept at room-temperature for 3 days. The ether was decanted and evaporated, leaving 1.78 g of a white amorphous mass with an acidic odor. It was dissolved in the minimum amount of hot ether and allowed to re-precipitate upon cooling. Yield = 1.54 g (60.3 %), m.p. = 168-172°C.

19F n.m.r. (in ether, CCl3F ext.): $\phi = 71.0$ ppm, m, int. = 5.0 (SF5); $\phi = 59.3$ ppm m, int. 1.1 (SO2F).

I.r.(neat sample, NaCl, cm$^{-1}$): 3400, 3341, 3254, peaks on broad feature, s, at $\approx 3300$; 2967, w; 1729, m-s; 1701, s; 1627, m; 1560, m; 1434, m-s; 1386, m; 1301, m; 1250, w; 1219, m-s; 1178, m; 1099, m-s; 1052, w; 976, m; 952, w; 891, s; 863, vs; 850, s; 822, m; 792, w-m; 783, m; 744, m; 736, m;
685, w; 645, w; 610, w; 600, w; 571, m.

Anal. calcd. for C\textsubscript{3}H\textsubscript{4}F\textsubscript{6}N\textsubscript{2}O\textsubscript{4}S\textsubscript{2}: C, 11.62; H, 1.30; F, 36.8; S, 20.7; N, 9.03. Found: C, 11.89; H, 1.30; F, 35.3; S, 18.81; N, 8.81.

F\textsubscript{5}SC\textsuperscript{−}(SO\textsubscript{2}F)CONHCONH\textsubscript{2}(C\textsubscript{2}H\textsubscript{5})\textsubscript{3}NH\textsuperscript{+} (36):

To 1.3085 g of F\textsubscript{5}SCH(SO\textsubscript{2}F)CONHCONH\textsubscript{2} (4.2 mmol) in 12 ml of anhydrous ether in a 50 ml pear-shaped flask, fitted with a Claisen-head, dropping funnel and a Drierite tube, 0.5187 g of N(C\textsubscript{2}H\textsubscript{5})\textsubscript{3} (5.1 mmol) in 10 ml of anhydrous ether was added dropwise at -70°C with stirring. At ≈ 0°C an oily layer was formed, while the ureide disappeared slowly. Removal of the solvent in vacuo left 1.714 g of an almost white solid (theor. 1.735 g), which contained a small impurity (\textsuperscript{19}F n.m.r.). Attempts to recrystallize the material resulted in decomposition.

\textsuperscript{19}F n.m.r. spectrum (aceton, C\textsubscript{Cl}\textsubscript{3}F ext.): AB\textsubscript{4}X \( \phi = 95.7 \) ppm, 9 lines with doublet splitting, \( J = 4.9 \) Hz, (F\textsubscript{A}), int. = 0.94; \( \phi = 82.0 \) ppm, d-m, int. = 4.00 (F\textsubscript{B}); \( \phi = 73.1 \) ppm, not resolved, int. = 0.99 (F\textsubscript{X}, SO\textsubscript{2}F).

I.r. of the solid obtained above (neat, KBr, cm\textsuperscript{-1}): 3367, s,br; 3220, m,br; 2980, s; 2938, m,sh; 2882, m; 2671, m, br; 2495, w, br; 2185, w-m (?); 1688, vs, 1652, vs; 1567, s; 1478, s; 1459, s, sh; 1370, vs; 1232, s-vs; 1167, vs; 1116, s; 1036, m; 945, w-m; 889, m; 854, vs; 834, vs; 754, s-vs; 734, s; 663, s; 626, m-s; 580, s,br; 480m-s.
\[
\text{F}_5\text{SCH}_2\text{CHClCH}_2\text{OC}(\text{O})\text{CH}_3 \quad (67): \\
\]
6.50 g of \(\text{CH}_2=\text{CHCH}_2\text{OC(O)CH}_3\) (Aldrich, 65 mmol), 11.69 g \(\text{SF}_5\text{Cl}\) (PCR, 72 mmol) and 25 ml \(\text{CCl}_3\text{F}\) (DuPont) were heated in a 75 ml steel bomb to 95°C for 15 days. After venting and bleeding 11.69 g off the gaseous products, the \(\text{CCl}_3\text{F}\) was distilled away at atmospheric pressure, and vacuum distillation (1-2 torr) resulted in four fractions: 1. 55°C (1.54 g); 2. 55-58°C (3.36 g); 3. 58-61°C (4.78 g); 4. 61-62°C (1.54 g). Fraction 1 was \(\approx 95\%\) pure adduct, the remainder being allyl acetate. Fractions 2, 3, and 4 had the same \(^1\text{H}\) n.m.r. spectra which showed only a trace of allyl acetate. The cold trap contained 1.96 g of liquid which was largely allyl acetate. Yield (2+3+4): 9.68 g (56.8 %).

\(^1\text{H}\) n.m.r. (neat sample, \(\text{Si(CH}_3\text{)}_4\) ext.): \(\delta = 2.50\) ppm, s, int. 3.0 (CH\(_3\)); \(\delta = 4-5.4\) ppm, m, int. 5.5.

\(^{19}\text{F}\) n.m.r. (neat sample, \(\text{CCl}_3\text{F}\) ext.): \(\text{AB}_4\) spectrum; \(\phi_A = 83.2\) ppm (9 lines), int. = 1.02; \(\phi_B = 67.7\) ppm (d-m), int. = 4.00. \(J_{AB} = 154.1\) Hz.

Infrared spectrum (neat sample, NaCl, cm\(^{-1}\)): 3029 (w); 2973 (w); 2855 (vw); 1752 (vs); 1460 (w-m); 1430 (m,sh); 1420 (m); 1385 (m-s); 1369 (m-s); 1318 (vw); 1302 (vw); 1232 (s-vs); 1111 (w); 1063 (m,sh); 1045 (m); 1026 (m); 945 (m); 873 (s); 846 (vs); 828 (vs); 728 (w); 715 (w-vw); 696 (vw); 636 (vw); 621 (w); 599 (m); 565 (w).

Mass spectrum (chemical ionization, mass, species, %): 263, (M\(^{35}\text{Cl}+\text{H})^+, 100; 181, (M\(^{35}\text{Cl})-\text{CH}_3\text{COOH-2H-F})^+, 11.5;
136, (M(35Cl)-SF5)+, 1.5; 135, (M(35Cl)-H-SF5)+, 29.9; 113, (M+H-Cl-3F-CH3COO)+, 1.9; 102, SF3CH+, 2.8; 101, SF3C+, 62.9; 89, SF3+, 1.6; 85, CH3COOH2C+, SCCCHO+, 6.4; 83, CH3COOC2+, SF2CH+, 4.4; 81, CHCOOC2+, 6.3; 79, C5H3O+, 3.3; 73, CH3COOCH2+, 9.5; 71, CH3COOC+, 10.8; 70, SF2+, C3H2S+, 4.4; 69, CHCOOC+, 12.0; 67, C4H3O+, 6.9.

Anal. calcd for C5H8ClF5O2S: C, 22.87; H, 3.07; Cl, 13.50; F, 36.2; S, 12.21. Found: C, 23.10; H, 3.19; Cl, 13.61; F, 36.5; S, 12.00.

F5SCH=CH2OH (68), by aqueous hydrolysis:

4.00 g of F5SCH2CHClCH2COOCH3 (5.2 mmol) were stirred at room temperature with 9.0 g of 23% KOH for 20 h. According to a 19F n.m.r. spectrum, all starting material was gone. The lower layer was pipetted off (2.59 g, 92.3%). There was apparently no significant amount of water, which had been suggested by the infrared spectrum, as drying over K2CO3 or CaO did not alter the infrared spectrum, which showed a broad feature at 1640 cm⁻¹. Vacuum distillation (2.5 mm Hg, 39-42°C) did also not lead to an altered infrared spectrum.

F5SCH=CHCH2OH (68), prepared in methanolic methoxide (anhydrous conditions):

To a solution of 67 (2.10 g, 8.0 mmol) in 2.0 g of anhydrous methanol in a 10 ml flask, equipped with magnetic
stirring bar, Claisen head, Drierite tube and dropping funnel, a solution of 0.23 g of sodium (0.01 mol) in 5.0 ml anhydrous methanol was added dropwise at -20°C (acetone and dry ice). There was no obvious reaction, so the cold-bath was removed and the reaction flask was allowed to attain room-temperature. When this temperature was nearly reached, a white precipitate formed quickly. Addition of more CH₃ONa solution caused more precipitate to form. After ≈ 0.8 times the total volume of methoxide solution had been added, stirring was stopped and the precipitate was allowed to settle. Addition of another drop of CH₃ONa to the clear colorless supernatent solution caused no more turbidness. After keeping the mixture for an additional 3 hours at room temperature, the methanol was removed by stirring the solution at ≈ 50 torr for 1 hour (the methanol was collected in a -196°C trap). Distillation at 6-6.5 torr afforded two fractions (54.5-57 °C, 0.54 g, 57-57.5 °C, 0.84 g) as colorless viscid liquids, which had identical infrared spectra. Yield = 1.38 g (93.8 %). A sample for ¹H n.m.r. spectroscopy was prepared by preparative gas chromatography (5% QF-1 on Chromosorb W, 1m. 120 °C).

¹H n.m.r. spectrum (CDCl₃, 400 MHz, Si(CH₃)₄): ABM₂X: δ₀A = 6.67 ppm (m), SF₅CH=; δ₀B = 6.55 ppm (m), SF₅CH=CH₂, area A+B = 2.00; δ₀M = 4.34 ppm (s), CH₂, area = 2.04; δ₀X = 1.75 ppm (s), OH, area = 1.13. J₀AB = 14.8 Hz.

¹⁹F n.m.r. spectrum (neat sample, external CCl₃F): AB₄
\( \varphi_A = 83.1 \) ppm, 9 lines with doublet splitting, area = 1.0; 
\( \varphi_B = 62.3 \) ppm, asymmetric doublet, area = 4.0; \( J_{AB} = 151.3 \) Hz; \( J_{AH} = 4.2 \) Hz.

Infrared spectrum (neat sample, film on KBr plates, cm\(^{-1}\)): 3664, w; 3353, m-s,br; 3103, w-m; 2923, w; 2879, w; 1644, w,br; 1449, w-m; 1434, w,sh; 1370, w; 1300, w; 1229, w-m; 1208, w,sh; 1097, m-s; 1030, m; 1009, w-m; 984, w-m; 940, s; 902, s-vs; 837, vs; 764, s; 726, m; 661, m; 648, m; 602, m; 571, m; 540, m; 441, w-m.

Mass spectrum (e.i., 70 eV, m/e, fragment, % > 2): 183, (M-H)\(^+\), 0.2 %; 135, (M-F-CH\(_2\)O)\(^+\), 48.7; 127, SF\(_5\)\(^+\), 16.1; 116, SF\(_3\)C\(_2\)H\(_3\)\(^+\), 2.4; 97, SF\(_2\)C\(_2\)H\(_3\)\(^+\), 5.7; 91, C\(_3\)H\(_4\)FS\(^+\), 3.7; 89, SF\(_3\)\(^+\); C\(_3\)H\(_5\)OS\(^+\), 100.0; 75, C\(_2\)FS\(^+\), 11.1; 73, (M+H-5F-OH)\(^+\), 2.1; 70, SF\(_2\)\(^+\), 16.5; C\(_2\)H\(_3\)S\(^+\) (M+H-5F-CH\(_2\)OH)\(^+\), 6.5; 58, C\(_2\)H\(_2\)S\(^+\), 4.9; 57, C\(_3\)H\(_5\)O\(^+\), 79.4; 55, C\(_3\)H\(_3\)O\(^+\), 22.8; 53, C\(_3\)HO\(^+\), 3.5; 51, SF\(^+\), 3.4.

\textbf{Anal.calcd} for C\(_3\)H\(_5\)F\(_5\)OS: C, 19.57; H, 2.74; F, 51.6; S, 17.41. Found: C, 19.69; H, 2.79; F, 51.5; S, 17.48.

\textbf{F\(_5\)SCF\(_2\)CF\(_2\)I (63):}

The preparation was similar to the method employed by Hutchinson, except that the reaction was run in a static system (bomb). Yields were comparable (\( \approx 35 \) %) to Hutchinson's (\( \approx 30 \) %). Thus, 13.06 g S\(_2\)F\(_{10}\) (51.0 mmol), 21.56 g of 82 % JCF\(_2\)CF\(_2\)J (rest was F\(_5\)SCF\(_2\)CF\(_2\)J, this mixture was recycled from earlier runs), (\( \approx 50 \) mmol) and 5.50 g of
terafluoroethylene (Peninsula Research Corporation, "PCR", 55 mmol), was heated at 120°C for 26 hours in a 75 ml stainless steel bomb behind an explosion shield. The product was collected by vacuum transfer (34.8 g after outgassing). Repetition in the same vessel with 16.14 g of ≈ 90 % C₂F₄J₂ (≈ 43 mmol), 10.99 g S₂F₁₀ (43.3 mmol), and 4.50 g of C₂F₄ at 131°C/16 hours yielded 31.30 g of crude product. Distillation of both products through a rectification column, 12.5x200mm, 0.05"x0.05"x0.1" steel helices, yielded disulfur decafluoride (room temperature-29°C, 16.66 g) and a purple liquid (80-96°C), which was (¹⁹F n.m.r.) 25% C₂F₄J₂ and 75% F₅SCF₂CF₂I, while the pot residue (28.97 g) was 12% F₅SCF₂CF₂J, 28% F₅S(CF₂CF₂)₂I and 60% C₂F₄J₂. The reaction was repeated with this pot residue, 14.03 g of S₂F₁₀ (54.9 mmol) and 8.78 g of C₂F₄ (heated overnight, 130°C). The product was vacuum transferred (40.0 g), combined with the above distillate, and rectified through the same column: S₂F₁₀, room temperature- 30°C, 8.70 g, F₅SCF₂CF₂⁻, 80-85°C (very slow distillation, circa 96 % pure). Yield (with respect to S₂F₁₀) ≈ 35 %. The bomb contained in the end ≈ 11 g of nonvolatile residue, which was rinsed out with acetone and filtered off: Waxy, off-white mass, shows CF and SF₅- bands in the infrared spectrum, darkens at circa 260°C under exhaustion of purple vapors. This material is probably F₅S(CF₂CF₂)ₙI.

¹³C n.m.r. spectrum of F₄B₄SCα₈₂Cβ₂Fβ₂J (100.6 MHz,
CDCl₃, Si(CH₃)₄: δₓ = 119.72 ppm (t-p); δᵧ = 91.30 ppm, t-t; Jₓₓᵧ = 25.9 Hz; Jₓᵧᵧ = 305.5 Hz; Jᵧᵧᵧ = 323.0 Hz; Jₓᵧᵧ = 41.0 Hz; Jₓᵧᵧ = 34.5 Hz.

**Sulfurtrioxodation of F₅SCF₂CF₂J (65):**

In an all-glass apparatus, consisting of a 25 ml pear-shaped flask with a magnetic stirring bar, a short reflux condenser, and a connector to a cold trap (-78°C, Drierite tube), 2.25 g of F₅SCF₂CF₂J (6.3 mmol) and 21.7 g of oleum (~ 30%) was heated to 130°C for 12 hours. The product (1.8 g) was condensed into a capillary tube (2 mm inner diameter) and cooled to -78°C for 1 hour. Two layers were formed, an upper (circa 1/3) and a lower layer. The tube was immersed in liquid nitrogen to somewhat less than the upper layer, and it could be removed by vacuum transfer. Repeating the -78°C bath procedure, some more of the upper layer material was obtained and it was removed in the same way. A total of 0.97 g of a colorless liquid was collected that did not separate anymore at -78°C. This material was not quite pure, but contained much less (by i.r.) of the impurity than samples subjected to fractional condensation. The product showed a strong CO absorption at 1890 cm⁻¹, and the impurity an absorption at 1360-1390 cm⁻¹. Yield ≈ 68%.

**F₅SCF=CF₂ (50):**

To 85 g cooled (ice-bath) ≈30% KOH (30 g 85% KOH
pellets + 60 ml H₂O) in a 250 ml round-bottomed flask, equipped with a reflux condenser and a magnetic stirring bar, 77.06 g of F₅SCHCF₂Br (267 mmol) were added. The top of the reflux condenser was connected with rubber tubes to two cold-traps (-78°C), the first of which was a 100 ml Schlenk flask, which was connected to the second trap. Cold tap water was passed through the condenser. With vigorous stirring, the reaction flask was heated very slowly to ≈40°C in 45 minutes. The reaction started already with the cold solution and the product (b.p. 19°C) escaped as a gas, which was chiefly trapped in the first -78°C cold-trap. After 1 hour, the bottom layer in the reaction flask had disappeared. The reaction flask was heated to 95°C for 3 more hours. Gas chromatography showed that the product (55.0 g) in the first cold-trap (the second one was almost empty) was ≈85% olefin and 15% F₅SCHCF₂Br. The base treatment was repeated, and in order to minimize loss by evaporation, the reaction vessel (no new base was added) was cooled by swirling in liquid nitrogen until its contents started to congeal. The product, which was kept in the -78°C bath, was poured in, and the procedure was repeated as above. Most of the product was collected in ≈30 minutes. The oil bath temperature was raised (140°C), and the cooling water was turned off. When the refluxing water had reached the top of the condenser the reaction was stopped. This allowed the collection of ≈1 more g of product. The product in both
cold-traps was combined and shaken with \( \text{P}_2\text{O}_5 \). Yield: 53.57 g or 96.5 % (g.c. pure).

The preparation of \( \text{F}_5\text{SCH=CF}_2 \) (52) is described at the beginning of the experimental part.

**Synthesis of \( \text{F}_5\text{SC(H,F)CF}_2\text{O} \) (53, 54):**

Methyl-tri-n-octylammonium chloride (1.50 g, 3.7 mmol, TCI) and 41.0 g of freshly prepared \( \text{NaOCl+NaOH} \) solution (calc. 11.8% in \( \text{NaOCl} \), 5% in \( \text{NaOH} \)) were cooled in a 100 ml round-bottomed flask with stirring in an ice-salt bath to -12°C. Then, 4.72 g of \( \text{F}_5\text{SCF=CF}_2 \) (22.7 mmol) in 20 ml of \( \text{CCI}_2\text{FCClF}_2 \), precooled to -10°C, were added in three portions, with vigorous stirring. After each addition, 2 minutes elapsed before the next addition. The temperature had risen to -6°C after the third addition and had dropped to -10°C three minutes later. \(^{19}\text{F n.m.r.} \) indicated that all olefin was gone. The mixture was poured into a separatory funnel and the lower layer was transferred and dried at -12°C (\( \text{Na}_2\text{SO}_4 \), 16 h). The liquid was vacuum-transferred and distilled with a spinning band column; the product was removed from the head of the column and condensed in a \(-78°C\) trap. Access of moisture to the cold-trap was prevented with a Drierite tube. After 5 h, 2.67 g (53%) of product was collected. This material was virtually pure and contained only a trace of Freon 113 (g.c.). The infrared spectrum was identical with the one of a g.c. pure sample of \( \text{F}_5\text{SCFCF}_2\text{O} \).
Infrared spectrum: (gas phase, 10 cm path length, KBr, 20 torr): (cm\(^{-1}\)); 1585 (m); 1530 (s); 1274 (vs); 1193 (s); 1133 (s); 1031 (m-s); 960 (w); 936 (m-s); 905 (vs); 863 (vs); 824 (m); 795 (w); 724 (m-w); 696 (m); 622 (m); 599 (m-s); 581 (m); 544 (w); 418 (w).

Mass spectrum (chemical ionization, mass, fragment, %): 209, (M+H-O)\(^+\), 0.9; 127, SF\(_5\)\(^+\), 9.1; 108, SF\(_4\)\(^+\), 6.5; 98, (M+H-SF\(_5\))\(^+\), 3.4; 97, (M-SF\(_5\))\(^+\), 72.2; 89, SF\(_3\)\(^+\), 43.8; 70, SF\(_2\)\(^+\), 5.6; 59, C\(_2\)F\(_7\)O\(^+\), 4.1; 51, SF\(^+\), 12.1; 50, CF\(_2\)\(^+\), 2.9.

**Anal.calcd** for C\(_2\)F\(_7\)OS: C, 10.72; F, 67.8; S, 14.31.

**Found**, C, 10.54; F, 68.2; S, 14.37.

F\(_5\)SCHC\(_2\)(\(_2\)O was obtained in a similar fashion with CCl\(_4\) at -10°C. The dried (Na\(_2\)SO\(_4\)) CCl\(_4\) layer was distilled twice through a 12 cm Vigreux column and gave a fraction with a boiling range 42.4-46.0 °C that consisted of SF\(_5\)CHC\(_2\)O (87% by g.c.) and CCl\(_4\) (13%). Further purification was achieved with preparative gas chromatography (30% SE-30, 3m, 95°C). Yield ≈ 30%.

Infrared spectrum (path length 10 cm, 10 torr, KBr, cm\(^{-1}\)): 3065(w); 1535(m); 1509(s-vs,sh); 1497(vs); 1297(s); 1250(vs); 1206(w); 1174(vw); 1128(s); 1049(s); 1003(w); 965(vs); 889(vs); 831(w-m); 767(w); 738(s); 677(m); 651(m); 613(s); 572(m); 534(w); 505(w); 435(w).

**Anal.calcd** for C\(_2\)HF\(_7\)OS: C, 11.66; H, 0.49; F, 64.5; S, 15.56. **Found**: C, 11.76; H, 0.56; F, 63.7; S, 15.65.

The samples were generally stored at -196°C, since for
F_5SCFCF_2O the observation was made that storage at room-temperature in glass vessels resulted in the formation of COF_2.

**Reaction of 54 with CsF in CH_3CN:**

Cesium fluoride (0.05 g, 3.3·10^{-4} mol) was dried overnight in a 40 ml Carius tube at 115 °C under high vacuum. Then 0.63 g of 54 (2.8 mmol) was added by vacuum transfer. The tube was heated for 45 minutes to 115 °C, and although the CsF had darkened somewhat and it seemed to stick together, the i.r. spectrum of the contents of the vessel showed only 54. To this, 0.34 g of dry CH_3CN was condensed and the tube was kept at 65 °C for 30 minutes (i.r., still some 54, but also a strong band at 1898 cm^{-1}). Heating was continued for 1 hour, when no more 54 was detected in the i.r. spectrum. Fractional condensation (- (35-16) °C, -(78-66) °C, -(100-95) °C, -196 °C) yielded some separation. The -35 °C trap was empty, the -78 °C trap contained 0.16 g of CH_3CN, the -100 °C (0.14 g) trap contained CH_3CN in mixture with other (unidentified) material. The 19F n.m.r. spectrum showed several very weak bands, but no SF_5 or CFO group was detectable. The -196 °C trap contained a mixture of SF_4 and CF_3CFO (0.48 g, 76 %). This latter mixture was stored in a 50 ml Pyrex vessel with rubber septum at room temperature for several weeks, and then subjected to preparative gas chromatography.
The effluent fractions were sucked into an evacuated gas infrared cell. The major band (≈ 77 %, Rf = 10.4 minutes) showed several irregularities and the i.r. samples were taken at various places of the band. Although these samples showed a continuous change, it was clear from comparisons that initially SiF₄, then SiF₄ + CF₃CFO and finally SiF₄ + CF₃CFO + SOF₂ were eluted. The minor band (23 %, Rf = 12.9 minutes) showed the following bands (cm⁻¹): 1212, s; 1179, vs; 1118, vs; 1047, vs; 980, vw, broad; 909, vs; 890, m,sh; 816, vs; 655, vw; 533, vw.

Reaction of 54 and H₂O:

To 1.71 g of distilled H₂O (65 mmol) in a 50 ml Pyrex-glass reaction vessel that contained a magnetic stirring bar and that could be closed with a Kontes Teflon valve, 1.07 g of 54 (4.8 mmol) was added by vacuum transfer. After stirring at room temperature overnight (18 h), the bottom layer was gone. A gas infrared spectrum showed a pressure drop of 20 torr in a total volume of 530 ml, as compared to the calculated value for 1.07 g of epoxide. The infrared spectrum (10 cm, 10 torr, KBr) showed only the epoxide and a weak band at 1374 cm⁻¹. The volatile material was re-condensed into the reaction vessel, and stirring was continued (room temperature, 24 h; 50 °C, 18 h). The pressure had dropped by another 10 torr after this time, and
only the epoxide was detected in the i.r. spectrum. After stirring another 18 h at 60 °C, a white gel-like mass was observed and three more days of stirring at 60°C, then keeping it at room temperature (24 h) did not show any change in the i.r. spectrum, neither was the pressure altered upon expansion of the material in the vessel in the same volume (530 ml). The volatile material was removed (vacuum transfer, 0.81 g of 54, 76 % recovery). The vessel was severely etched as seen after removing the water and the white inorganic mass that remained.

Reaction of 54 and BF₃:

To 0.30 g of 54 (1.3 mmol) in a 50 ml round bottomed flask, equipped with a Kontes Teflon valve, 6.7·10⁻⁴ mol of BF₃ (Matheson) was condensed. No reaction was observed upon warming to room temperature, neither showed the i.r. spectrum any other products than BF₃ and 54 after keeping the sample at room temperature for 18 hours.

F₅SCH=CHOCH₃ from F₅SCH=CHBr:

In a 250 ml round bottom flask, equipped with a magnetic stirring bar, 11.59 g of sodium ( 504 mmol) were dissolved in 120 ml of anhydrous methanol. A Claisen head, dropping funnel and a Dryerite tube were attached, the solution was cooled in an ice-bath and 80.45 g (345.3 mmol) of F₅SCH=CHBr were added dropwise under stirring to the
sodium methoxide solution. Sodium bromide started soon to precipitate and the solution had a yellow tinge. Addition was finished after 30 min and the ice-bath was removed. The solution became warm slowly and was cooled with a water bath (room temperature). After 1 h the Claisen head was replaced by a Drierite tube. Stirring was continued for 96 h at room temperature, when $^{19}$F n.m.r. indicated that all the bromide had gone. The NaBr was filtered off (25.3 g, 71%) and the reaction mixture was poured into 300 ml of cold water, the lower layer was separated and saturated NaCl solution (200 ml) was added to the aqueous layer. After standing overnight a little bit more product had precipitated. It was washed with a small amount of saturated NaCl solution and combined with the original product; after drying (MgSO$_4$), 54.7 g (86%) of crude product was obtained. Distillation through a Vigreux column at atmospheric pressure yielded 17.7 g of pure F$_5$SCH=CHOCH$_3$ (b.p. 114°C) and 36.2 g of a mixture the vinyl ether and the acetal F$_5$SCH$_2$CH(OCH$_3$)$_2$ in a ratio of about 9:1 (by $^{19}$F n.m.r.), boiling range 114-122°C.

**Acid hydrolysis of** F$_5$SCH=CHOCH$_3$

In a 100 ml round bottom flask, fitted with a 30 cm reflux condenser, 53.7 g of a mixture of F$_5$SCH=CHOCH$_3$ and F$_5$SCH$_2$CH(OCH$_3$)$_2$ (ca 94:6, by $^{19}$F n.m.r.) were heated to 80 - 90°C with 27 ml of conc. HCl and vigorous stirring. After 3 h all starting material appeared to have been
consumed ($^{19}$F n.m.r.), but heating was continued for another hour. Yield 51.6 g, mixture of two compounds in the ratio of 34 (F$_5$SCH$_2$CHO) : 66 (F$_5$SCH$_2$CH(OH)$_2$ ($^{19}$F n.m.r.). Yield 51.6 g. This material was directly used for the preparation of the aldehyde by dehydration with P$_4$O$_{10}$ (next paragraph). In this fashion 22.1 g P$_{50_{10}}$ were used to dehydrate 39.6 g of the above mixture, leaving a product (29.55 g) that contained still the dimethyl acetal and some unreacted vinyl ether. The mixture was thus distilled through a Vigreux column to obtain 20.98 g of a mixture of the vinyl ether (20%) and the aldehyde, boiling range 55-96 °C. This fraction was heated with an excess of concentrated HCl for 15 hours in a 50 ml round bottom flask with a 30 cm reflux condenser to 80-89 °C. A 25 : 75 mixture of the aldehyde and the aldehyde hydrate was obtained (19 g).

(F$_5$SCH$_2$CHO)$_3$:

To 18.99 g of a mixture of F$_5$SCH$_2$CHO and F$_5$SCH$_2$CH(OH)$_2$ (25 : 75 % by $^{19}$F n.m.r.) 16 g (56.3 mmol) of powdered P$_{40_{10}}$ were added under cooling in a 50 ml round bottomed flask. The mixture was swirled occasionally and after ca. 30 min the aldehyde was removed by vacuum transfer (15.64 g).

The residual phosphorous pentoxide was dissolved in 50 ml of distilled water and extracted with chloroform (3x20 ml); the combined extracts were dried (Na$_2$SO$_4$) and brought to dryness, leaving an off-white solid, 0.87 g after repeated
recrystallization from hexane, m.p. 110.5-111.5 °C. An analytically pure sample was obtained by sublimation (0.001 torr, 50 °C).

$^1$H n.m.r. spectrum (CDCl$_3$, external Si(CH$_3$)$_4$): $\delta_1 = 3.81$ ppm, p - d, area = 2.0 (CH$_2$); $\delta_2 = 5.58$ ppm, t, area = 1.0 (CHO); $J_{1FB} = 7.9$ Hz, $J_{12} = 4.8$ Hz.

$^{19}$F n.m.r. spectrum (CDCl$_3$, external CFCl$_3$): AB$_4$ $\phi_A = 82.0$ ppm, area = 1.0; $\phi_B = 69.5$ ppm, area = 4.0, $J_{AB} = 152$ Hz.

I.r. spectrum (sample squeezed on KBr plates): 3044 (w); 2988 (vw); 2932 (vw); 1450 (vw); 1423 (m); 1405 (m); 1368 (ms); 1301 (vw); 1227 (m); 1142 (vs); 1068 (s); 1049 (ms); 901 (vs); 867 (s); 9838 (vs); 830 (vsh); 715 (ms); 651 (s); 607 (ms); 581 (m); 556 (s); 516 (m).

Mass spectrum (e.i., m/e, % > 1, species): 341, 1.8, (F$_5$SCH$_2$CHO)$_2$H$^+$; 321, 1.1, ((F$_5$SCH$_2$CHO)$_2$H-HF)$^+$; 171, 14.0, F$_5$SCH$_2$CHOH$^+$; 170, 1.2, F$_5$SCH$_2$CHO$^+$; 169, 26.8, F$_5$SCH$_2$CO$^+$; 153, 5.7, F$_5$SCH$_2$C$^+$; 152, 3.2, F$_5$SC$_2$H$^+$; 151, 100.0, F$_5$SC$_2$; 127, 2.9, SF$_5$$^+$; 122, 8.6, C$_3$H$_3$FO$_2$S$^+$; 105, 1.4, C$_3$H$_2$FOS$^+$; 103, 2.2, C$_3$FOS$^+$; 91, 9.6, FSC$_2$O$^+$; 89, 40.3, SF$_3$$^+$; 77, 1.3, FSCH$_2$C$^+$; 70, 1.3, SF$_2$$^+$; 65, 2.6, CH$_2$FS$^+$; 63, 11.5, F$^+$; 62, 13, CH$_2$OS$^+$; 61, C$_3$H$_3$FO$_2$S$^{2+}$.

Anal. calcd for C$_6$H$_9$F$_{15}$O$_3$S$_3$: C, 14.12; H, 1.78; F, 55.9; S, 18.85. Found: C, 14.28; H, 1.78; F, 55.9; S, 19.04.
$\text{F}_{5}\text{SCH}_2\text{CHO} + \text{H}_2\text{SO}_4$:

Into an n.m.r. tube, 0.5180 g (3.05 mmol) of $\text{F}_{5}\text{SCH}_2\text{CHO}$ (purified by preparative gas chromatography) were added. Four drops of 96% $\text{H}_2\text{SO}_4$ were added at room temperature, and an immediate formation of a white compound was observed. The tube was capped and gently shaken, whereby it warmed somewhat, and within a short time there was only a white solid left in the tube. The typical odor of the aldehyde was gone after 20 min. After keeping the sample at room temperature overnight the product was dissolved in acetone ($\approx 2\text{ml}$). The volume of the solution was reduced in a stream of air to $\approx 0.5\text{ ml}$, $\text{CH}_2\text{Cl}_2$ (3ml) was added, then saturated $\text{NaHCO}_3$ solution with swirling until $\text{CO}_2$ formation ceased. Removing the lower layer (pipet), extracting the remaining aqueous layer with methylene chloride (1 ml), combining the two solutions and drying ($\text{Na}_2\text{SO}_4$) followed. The dried solution was filtered through a filter pipet, the $\text{Na}_2\text{SO}_4$ was rinsed with $\text{CH}_2\text{Cl}_2$ (1 ml), the rinse solution was combined with the filtrate and after evaporation of the solvent in a stream of air and drying in the oven (120°C, 15 min) an off-white product (solidified melt) was obtained. The infrared spectrum of this material was identical to the spectra of the products obtained either by the extraction of the $\text{P}_4\text{O}_{10}$ residue from the dehydration procedure or from the precipitate collected from the aldehyde after some time of storage in the cold. Yield: 0.4460 g (0.87 mmol), 86%.
\( \text{F}_5\text{SCH}_2\text{CHBrOAc} \):

Into a 300 ml dry glass-vessel containing a stirring bar and equipped with a Kontes-Teflon valve, 100 ml of CCl\(_3\)F, previously dried over \( \text{P}_4\text{O}_{10} \), were vacuum-transferred. Then 12.67 g of SF\(_5\)Br (51.5 mmol) were condensed to the solvent. The CCl\(_3\)F was allowed to melt and dissolve the SF\(_5\)Br. Freshly distilled vinyl acetate (Aldrich) (10.0 g, 116.3 mmol) was added by vacuum-transfer, and the reaction vessel was allowed to warm slowly in a cold-bath until the solution started to melt; it was then swirled so as to achieve complete mixing. Another 7.8 g (37.7 mmol) of SF\(_5\)Br was added by the same method. The reaction mixture was now allowed to attain room-temperature slowly in the air with stirring. Only slight bubbling was observed. The solvent was then removed (rotary evaporator), leaving 19.8 g of a lightly yellowish liquid. Distillation through a 12 cm Vigreux column at gave 8.5 g of a main fraction, b.p. 45-48°C/6 mm Hg. The foreshot and the fraction 48-49°C (2.0 g together) had the same composition (\(^1\text{H n.m.r.}\)) as the main fraction. Yield 10.5 g, 36.2 %. The 48-49°C fraction was used for analysis.

\(^1\text{H n.m.r.} \text{(CDCl}_3, \text{Si(CH}_3)_4 \text{ ext.}): \delta_1 = 2.10 \text{ ppm, s, int.} = 3.17 (\text{CH}_3); \delta_2 = 4.74 \text{ ppm, m, int.} = 1.92 (\text{CH}_2); \delta_3 = 7.15 \text{ ppm, d-d, int.} = 1.00 (\text{CH}). J_{23}(\text{cis}) = 9.6 \text{ Hz, } J_{23}(\text{trans}) = 3.0 \text{ Hz.}

\(^19\text{F n.m.r.} \text{(neat sample, CCl}_3\text{F ext.): (AB}_4 \phi = 81.4\)
ppm, 9 lines, int. = 0.96 (F\textsubscript{A}); \phi = 65.9 ppm, d-m, int. = 4.0 (F\textsubscript{B}); J\textsubscript{AB} = 146.4 Hz.

I.r. spectrum (NaCl, neat, cm\textsuperscript{-1}): 3042, w; 3023, w; 2988, w-vw; 1778, vs; 1419, m; 1377, m-s; 1358, w-m; 1312, w; 1203, vs; 1193, vs, sh; 1103, s-vs; 1040, vs; 998, m; 941, m-s; 879, vs; 844, vs; 817, s-vs; 735, w; 675, w; 645, vw; 614, w; 602, m; 577, w; 564, vw.

Anal. calc'd for C\textsubscript{4}H\textsubscript{5}BrF\textsubscript{5}O\textsubscript{2}S: C, 16.39; H, 2.06; Br, 27.27; F, 32.42; S, 10.94. Found: C, 17.19; H, 2.18; Br, 28.86; F, 30.4; S, 9.90.

\textit{F\textsubscript{5}SCH\textsubscript{2}CH(OC\textsubscript{2}H\textsubscript{5})\textsubscript{2}i}

To 22.86 g of F\textsubscript{5}SCH\textsubscript{2}CHClOAc (92.0 mmol) in a 100 ml flask 32.46 g of absolute ethanol (705.6 mmol, 2.56-fold excess) were added at room-temperature. The lightly stoppered flask was set aside. After two days there was still some F\textsubscript{5}SCH\textsubscript{2}CHClOAc (i.r.), but after 48 h all of the compound was consumed. The smell of ethyl acetate was noticeable. The mixture was poured into 200 ml of water, washed again with water (2 x 30 ml), dried (Na\textsubscript{2}SO\textsubscript{4}) and distilled through a 12 cm Vigreux column. 16.47 g (73.4 %) of product boiling from 66-72\degree C / 19-20.5 torr was collected.

\textsuperscript{1}H n.m.r.(CDCl\textsubscript{3}, Si(CH\textsubscript{3})\textsubscript{4}): 1.27 ppm, t, (6H, CH\textsubscript{3}), J\textsubscript{H-H} = 7.1 Hz; 3.67 ppm, m, (6H, H\textsubscript{3}CH\textsubscript{2} + F\textsubscript{5}SCH\textsubscript{2} ); 4.92 ppm, t, (1H, CH), J\textsubscript{H-H} = 5.2 Hz).
$^{19}$F n.m.r. (neat sample, CCl$_3$F ext.): (AB$_4$), $\phi = 83.7$ ppm ($F_A$), 9 lines, (1F); $\phi = 66.8$ ppm ($F_B$), m, (4F); $J_{AB} = 143.8$ Hz.

I.r. (cm$^{-1}$): 3035, vw; 2985, s; 2936, m; 2904, m; 2893, m; 2805, vw; 1484, w; 1458, w, sh; 1448, w-m; 1417, m; 1378, m-s; 1361, m; 1349, m; 1300, w; 1229, w; 1159, m, sh; 1125, s-vs; 1063, s-vs; 1023, s, sh; 1000, w-m, sh; 935, w-m, 906, s; 865, s-vs; 840, vs; 817, vs; 737, w-m; 710, s; 667, w; 643, s; 630, m; 620, m; 600, s; 566, m; 539, w.

Mass spectrum (c.i., 70 eV, mass, fragment, %> 2):

243, (M-H)$^+$, 0.15; 201, (M+H-C$_2$H$_5$-CH$_3$)$^+$, 5.0; 200, (M-C$_2$H$_5$-CH$_3$)$^+$, 4.9; 199, (M-C$_2$H$_5$O)$^+$, 100; 179, C$_4$H$_7$F$_4$OS$^+$, 14.2; 178, C$_4$H$_6$F$_4$OS$^+$, 2.5; 177, C$_4$H$_5$F$_4$OS$^+$, 40.2; 171, (M-C$_2$H$_5$-C$_2$H$_4$)$^+$, 26.5; 153, (M-C$_2$H$_5$O-C$_2$H$_5$OH)$^+$, 3.0; 152, (M-2C$_2$H$_5$OH)$^+$, 2.0; 151, C$_2$SF$_5^+$, 54.3; 127, SF$_5^+$, 1.2; 107, C$_6$H$_3$O$_2^+$, 2.5; 104, C$_6$O$_2^+$, CH$_3$SF$_3^+$, 3.5; 103, CH$_2$SF$_3^+$, 59.7; 92, C$_6$H$_4$O$^+$, C$_2$HFOS$^+$, 2.7; 91, C$_2$FOS$^+$, 57.4; 89, SF$_3^+$, 19.5; 87, C$_2$H$_2$O$_2$C$_2$H$_5^+$, 2.4; 79, FSC$_2$H$_4^+$, 2.9; 77, FSC$_2$H$_2^+$, 11.8; 75, FSC$_2^+$, C$_2$H$_3$OS$^+$, 8.0; 73, C$_2$HOS$^+$, C$_3$H$_5$O$_2^+$, 7.9; 72, C$_2$OS$^+$, C$_3$H$_4$O$_2^+$, 2.9; 70, two fragments, SF$^+$, C$_3$H$_2$O$_2^+$, 2.2, 1.9; 69, C$_3$H$_2$O$_2^+$, C$_2$OC$_2$H$_5^+$, 3.2; 65, C$_4$HO$^+$, 1.5; 64, C$_4$O$^+$, 2.0; 63, CFS$^+$, 71.3; 61, (SC$_2$H$_3$+2H)$^+$, 11.4; 59, SC$_2$H$_3^+$, CH$_2$OC$_2$H$_5^+$, 2.3; 57, SC$_2$H$^+$, COC$_2$H$_5^+$, 4.7; 56, SC$_2^+$, COC$_2$H$_4^+$, 3.6; 55, COC$_2$H$_3^+$, 6.8; 54, COC$_2$H$_2^+$ 1.1.

Anal.calcd for C$_6$H$_{13}$F$_5$O$_2$S: C, 29.51; H, 5.37; F, 38.9; S, 13.13. Found: C, 29.50; H, 5.23; F, 39.0; S, 13.03.
The compound was obtained by a procedure similar to the one for $F_5SCH_2CH(OCH_3)_2$ with methanol. Yield: 81.2%. B.p. 83-84°C/ 96 mm Hg.

$^{19}$F n.m.r. (neat sample, CCl$_3$F ext.): AB$_4$ $\phi = 83.1$ ppm, 9 lines, int. = 1.0 (F$_A$); $\phi = 66.5$ ppm, d-m, int. = 4.0 (F$_B$); $\text{J}_{AB} = 144.7$ Hz.

I.r. (neat sample, NaCl, cm$^{-1}$): 2960, vw; 2945, w; 2847, vw; 1465, w; 1451, w; 1418, w; 1387, w; 1387, w; 1187, w-m; 1126, m; 1078, m; 1060, w-m, sh; 1023, w-m; 976, w-m; 879, m-s; 840, vs; 817, vs; 725, w-m; 712, w-m; 641, w.

To 2.96 g of $F_5SCH_2CH(OCH_3)_2$ (13.7 mmol) in a 50 ml round-bottomed flask with a stirring bar 2.96 g of m-chloroperbenzoic acid (≈ 85%, ≈ 14 mmol, Aldrich) were added and a reflux condenser was attached. The mixture was heated slowly with stirring to 100°C and a homogeneous melt was obtained. After ≈ 2.5 h the melt was solidified, but after heating overnight (16h) it was liquid again. $^{19}$F n.m.r. indicated the reaction to be incomplete. Another 1.5 g of peracid was added and heating was continued at 96°C for 20h. From the mixture 1.82 g of crude product was collected by vacuum-condensation. This product was 99% pure by g.c.. Yield 66.4 %. An analytically pure sample was obtained by preparative gas chromatography on a SE-30 column at 100°C.
The ester is a clear, colorless pleasant-smelling liquid.

$^1$H n.m.r. (CDCl$_3$, Si(CH$_3$)$_4$ ext.): $\delta = 4.58$ ppm, s, int. = 3.0 (CH$_3$); $\delta = 5.08$ ppm, p, int. = 2.0 (CH$_2$). $J_{HF} = 7.7$ Hz.

$^{19}$F n.m.r. (neat sample, CCl$_3$F ext.): AB$_4$ $\phi = 78.2$ ppm, 9 lines, int. = 1.02 (FA); $\phi = 69.0$ ppm, d-m, int. = 4.00 (FB). $J_{AB} = 145.7$ Hz.

I.r. (neat, NaCl, cm$^{-1}$): 3058, w; 3002, w; 2966, w; 2854, vw; 1757, vs, (C=O); 1441, s; 1321, s; 1272, m; 1166, s; 1009, m; 948, m; 836, vs; 787, m; 709, m; 660, m; 611, w; 569, vw.

Mass spectrum (electron impact, 70 eV, mass, species, %>1): 181, (M-F)$^+$, 1.5; 180, (M-HF)$^+$, 1.2; 171, C$_2$H$_4$F$_5$OS$^+$, 4.0; 170, (M+H-CH$_3$O)$^+$, 2.9; 169, (M-CH$_3$O)$^+$, 1.7; 168, (M-CH$_3$OH)$^+$, 2.1; 149, (M-CH$_3$O-HF)$^+$, 2.1; 131, (M-CH$_3$O-2F)$^+$, 2.3; 127, SF$_5^+$, 20.8; 122, FSCCOOCH$_3^+$, 3.1; 119, C$_3$FO$_2$S$^+$, 2.2; 91, C$_2$FOS$^+$, 5.1; 89, SF$_3^+$, C$_2$HO$_2$S$^+$ (two fragments), 100, 30.2; 74, CH$_2$COOCH$_3^+$ (13C), 1.7; 73, CH$_2$COOCH$_3^+$ (12C), 46.3; 72, CH$_2$COOCH$_2^+$, 11.0; 70, SF$_2^+$, 6.8; 69, C$_3$HO$_2^+$, 8.6; 65, CH$_2$FS$^+$, 3.1; 62, CH$_2$OS$^+$ (rearr.), 2.7; 61, CHOS$^+$ (rearr.), 64.6; 60, COS$^+$, C$_2$H$_4$O$_2^+$, 4.3; 59, C$_2$H$_3$O$_2^+$, 36.9; 51, SF$^+$, H$_2$SO$^+$ (two fragments), 2.2, 2.1.

Anal.calcd. for C$_3$H$_5$F$_5$O$_2$S: C, 18.00; H, 2.52; F, 47.5; S, 16.02. Found: C, 18.17; H, 2.58; F, 48.2; S, 15.96.
Into a thoroughly dry 100 ml Pyrex-glass Carius tube that contained a magnetic stirring bar, 10.01 g of AcO\text{CH=CHCOOC}_{2}\text{H}_{5} (63.4 mmol) and 45 ml of CCl\text{3F} (Dupont, dried over P\text{4}O\text{10}) was added. The solution was degassed three times by freeze-pump-thaw-freeze cycles, and then 18.15 g (87.7 mmol) of bromine-free SF\text{5Br} was added via vacuum transfer. The lower part of the tube (circa 3 cm) was immersed in an ice bath, and the solution was stirred and irradiated (GE-250 W sunlamp, 80 cm distance) for 14 hours. When the volatile materials were condensed into a very dry cold trap, it was found that the residue contained still some olefin, and the volatile materials and an additional 4.36 g of SF\text{5Br} (21.1 mmol) were recondensed into the Carius tube. Irradiation at ice-temperature was continued for 7.5 hours (50 cm distance). A check of the i.r. spectrum showed still some olefin to be present, and the volatile materials were recondensed and irradiation was continued for another 10 hours (30 cm distance) at 0°C. The solution was by now distinctly yellow and a precipitate had appeared; no more C=C bond was detected in the infrared spectrum (the Carius tube was stored between the operations in an ice bath). After vacuum transfer of the volatile materials, 20.5 g of a lightly brown oil remained behind (yield (crude) = 88.7%). The $^{19}$F n.m.r. spectrum showed that there was very little of the impurity at -120-130 ppm; when the reaction is not
carried out a low temperature there is much more of this impurity. Part of this crude product was distilled (5.01 g, 0.017 torr), and a fraction, boiling range 55-70°C (1.81 g, first fraction, oil bath temperature 100°C), contained some of the C-F impurity (φ -120-130 ppm) and showed some enrichment of the φ ≈ 52 ppm impurity; the second fraction, b.p. 58-61°C, (1.20 g, oil bath temperature = 90°C, very slow distillation), a colorless oil, was free of the C-F impurity. There is another contaminant, an SF₅ compound φₐ ≈ 52 ppm (2% integrated intensity), and possibly another contaminant buried under the B₄-resonance of the main product (shoulder). The samples were stored in the refrigerator, as they turned yellowish at room temperature within hours. Yield (extrapolated) ≈ 56 %.

The purity of the second fraction is estimated from ¹⁹F n.m.r. measurement to 98 %.

¹H n.m.r. spectrum (neat sample, external Si(CH₃)₄):
δ₁ = 1.68 ppm, t, area = 3.1 (CH₃CH₂); δ₂ = 2.55 ppm, s, area = 3.0 (CH₃C(O)O); δ₃ = 4.74 ppm, quartet, area = 2.0 (CH₃CH₂); δ₄ = 5.53 ppm, d-p, area = 1.0 CHCOOC₂H₅; δ₅ = 7.63 ppm, d, area = 0.9, (CH₃C(O)OCH). J₁₃ = 7.22 Hz; J₄₅ = 10.8 Hz; J₄B = 5.7 ± 0.2 Hz (coupling to SF₅, AB₄). All lines are broadened.

¹⁹F n.m.r. spectrum (neat sample, external CCl₃F): φₐ = 79.0 ppm, 9 lines, area = 1.0; φₐ = 67.3 ppm, d-m, area 4.0; JₐB = 149.7 Hz.

Infrared spectrum (neat sample on KBr, cm⁻¹): 3025,
vw, sh; 2992, w; 2945, vw; 1784, s; 1755, s; 1470, w; 1449, w; 1440, w; 1373, m; 1354, w; 1308, m; 1262, m; 1195, s; 1164, m, sh; 1116, m, s; 1104, s; 1054, m, sh; 969, vw; 940, vw; 890, s; 854, vs; 795, m; 735, vw; 697, w, br; 671, w; 665, w; 621, w; 599, m; 570, m; 525, w.

Mass spectrum (e.i., 70 eV, mass, species, % > 1%):
285, (M-Br)\(^+\), 1.7; 197, (M-Br-COOC\(_2\)H\(_5\)-CH\(_3\))\(^+\), 2.2; 194, (M-Br-COOC\(_2\)H\(_5\)-O-2H)\(^+\), 1.1; 168, F\(_5\)SCHCO\(^+\), 1.3; 166, C\(_3\)H\(_3\)F\(_5\)S\(^+\), 2.2; 149, C\(_2\)HF\(_4\)OS\(^+\), 7.1; 146, C\(_5\)H\(_3\)FO\(_2\)S\(^+\), 1.9; 127, SF\(_5\)\(^+\), 1.4; 89, SF\(_3\)\(^+\), 6.7; 88, C\(_2\)O\(_2\)S\(^+\), C\(_3\)HFS\(^+\), 1.0; 87, C\(_3\)FS\(^+\), 3.1; 73, C\(_3\)H\(_5\)O\(_2\)\(^+\), 1.1; 71, C\(_3\)H\(_3\)O\(_2\)\(^+\), 5.1; 70, SF\(_2\)\(^+\), C\(_3\)H\(_2\)O\(_2\)\(^+\), 1.9; 69, C\(_3\)HO\(_2\)\(^+\), 3.2; 45, C\(_2\)H\(_5\)O\(^+\), 3.8; 44, C\(_2\)H\(_4\)O\(^+\), 3.9; 43, CH\(_3\)CO\(^+\), C\(_2\)H\(_3\)O\(^+\), 100.0.

Anal. calcd for C\(_7\)H\(_{10}\)BrF\(_5\)O\(_4\)S: C, 23.03; H, 2.76; Br, 21.88; F, 26.0. Found: C, 22.77; H, 2.67; Br, 22.21; F, 25.4.

**Acidic Hydrolysis of F\(_5\)SCH\(_2\)CHClOAc:**

The adduct F\(_5\)SCH\(_2\)CHClOAc (3.02 g, 12.2 mmol) was stirred vigorously with 5 ml of concentrated HCl in a 50 ml round bottomed flask with a short reflux condenser at 70-73°C for 6 hours. The product (0.96 g, 46.5 %) contained only a small amount of acetic acid and had only a weak OH band, but was otherwise identical with F\(_5\)SCH\(_2\)CHO (i.r. spectrum). The product was not further purified.
To 7.92 g of \( \text{F}_5\text{SCH}_2\text{CHClO(C(O)CH}_3 \) (31.9 mmol) in 10 ml of anhydrous \( \text{CH}_3\text{OH} \), kept in a 100 ml round bottomed flask, equipped with magnetic stirring bar, Claisen head, Drierite tube and dropping funnel, was added 2.15 g of 85% KOH, dissolved in 10 ml anhydrous methanol (32.6 mmol). A slight excess is necessary because the starting material and the product cannot be separated by distillation when the reaction is not complete. The base solution was added within 20 minutes at \(-78^\circ\text{C} \), and the solution was then brought slowly to room temperature. The methanol was distilled off at atmospheric pressure. A small amount of water was then added and the precipitate was removed (5.50 g). No starting material was present. Distillation (3-4 torr, 30-37°C) gave 1.58 g of \( \text{F}_5\text{SCH}_2\text{CH(OH)OCH}_3\text{OC(O)CH}_3 \). The contents of the cold trap (\(-196^\circ\text{C} \)) were distilled to give 2.85 g of a mixture \( \text{F}_5\text{SCH}_2\text{CH(OH)}_2 \) + \( \text{F}_5\text{SCH}_2\text{CHO} \). Yield \( \approx 45 \% \), and some \( \text{F}_5\text{SCH}_2\text{CH(OH)OCH}_3\text{OC(O)CH}_3 \) (0.36 g). Yield \( = (1.58 + 0.36)g = 1.94 g \) (24.8 %). An analytically pure sample was obtained by preparative g.c. on a Carbowax (20 %) column, 3 ft, 100 °C, 1 ml He/sec.

\( ^1\text{H n.m.r. spectrum (CDCl}_3, \text{Si(CH}_3)_4 \text{ ext.}): \delta_1 = 2.65 \text{ ppm, s, 1.00 H (OCH}_3); \delta_2 = 4.03 \text{ ppm, s, 1.00 H, (C(O)CH}_3); \delta_3 = 4.37 \text{ ppm, d-p, 2.07 H, (SF}_5\text{CH}_2); \phi_4 = 6.70 \text{ ppm, t, 0.98 H, (CH)}. \)

\( J_{\text{CH}_2-B} = 8.23 \text{ Hz}; J_{\text{H-H}} = 5.13 \text{ Hz}. \)
\(^{19}\text{F}\) n.m.r. spectrum (CDCl\(_3\), external CCl\(_3\)F): AB\(_4\) system, \(\phi_A = 83.67\) ppm, 9 lines, 1.07 F; \(\phi_B = 68.79\) ppm, d-m, 4.00 F. \(J_{AB} = 147.4\) Hz.

Infrared spectrum (neat sample, KBr, cm\(^{-1}\)): 3041, vw; 3013, vw; 2973, w; 2948, w; 2858, w; 1755, s; 1453, w-m; 1420, w-m; 1377, m; 1360, w,sh; 1300, vw; 1255, m, sh; 1230, s; 1190, m-s; 1142, m; 1128, m; 1083, m,sh; 1078, m; 1050, m; 1014, s; 992, m; 941, m,broad; 879, s; 842, vs; 824, s-vs; 714, w-m; 655, w, sh; 641, m; 604, m; 581, w; 564, w-m; 558, w-m; 549, w,sh; 581, vw; 566, w; 557, w; 550, vw.

Mass spectrum (e.i., mass, species, % > 1): 185, (M-CH\(_3\)COO\(^+\)), 90.7 (+ peaks at 187 (3.8) and 186 (4.9), S + C isotope peaks); 184, (M-CH\(_3\)COOH\(^+\)), 19.7; 169, (M-CH\(_3\)COOH-CH\(_3\))\(^+\), 1.6; 165, (M-CH\(_3\)COO-HF\(^+\)), 12.3; 163, (M-CH\(_3\)0-CH\(_3\)-O-F\(^+\)), 3.0; 153, (M-CH\(_3\)COOH-CH\(_3\)0\(^+\)), 5.3; 149, (M-CH\(_3\)COOH-CH\(_3\)-HF\(^+\)), 2.3; 127, SF\(_5\)\(^+\), 12.0; 122, C\(_3\)F\(_2\)OS\(^+\), 3.4; 103, C\(_4\)H\(_7\)O\(_3\)\(^+\), 1.4; 101, C\(_4\)H\(_8\)O\(_3\)\(^+\), 2.5; 96, F\(_2\)SC\(_2\)H\(_2\)\(^+\), 1.1; 91, FSCCO\(^+\), 1.9; 89, SF\(_3\)\(^+\), 43.6; 87, C\(_3\)H\(_3\)O\(_3\)\(^+\), 2.3; 83, C\(_4\)H\(_3\)O\(_2\)\(^+\), F\(_2\)SCH\(^+\), 1.3; 81, C\(_4\)HO\(_2\)\(^+\), 1.9; 78, C\(_2\)H\(_2\)FS\(^+\), 3.7; 77, C\(_2\)H\(_2\)FS\(^+\), 74.5; 76, C\(_2\)HFS\(^+\), 11.9; 75, C\(_2\)H\(_3\)O\(_3\)\(^+\), C\(_2\)FS\(^+\), 8.7; 74, C\(_2\)H\(_2\)OS\(^+\), 7.6; 73, C\(_2\)HO\(_2\)\(^+\), 3.3; 70, C\(_3\)H\(_2\)O\(_2\)\(^+\), SF\(_2\)\(^+\), 6.2; 69, C\(_3\)HO\(_2\)\(^+\), 1.1; 65, CH\(_2\)FS\(^+\), 3.9; 64, CHFS\(^+\), 1.0; 63, CFS\(^+\), 100.0; 61, (CH\(_3\)OCHO·H\(^+\)), 13.2; 60, CH\(_3\)COOH\(^+\), 18.5; 59, CH\(_3\)COO\(^+\), 1.2; 58, CH\(_2\)COO\(^+\), 17.0; 57, CHCOO\(^+\), 17.2.

F$_5$SCHBrCH$_2$OCH$_3$i

Sodium (0.67 g, 29.1 mmol) was dissolved in 11.5 ml of anhydrous methanol in a 50 ml pear-shaped flask. To this solution 5.86 g of SF$_5$CBr=CH$_2$ (25.2 ml) were added dropwise (1 minute, dropping funnel) with (magnetic) stirring. The dropping funnel was replaced by a Drierite tube. After ≈ 10 minutes, the mixture turned yellow and became warm. The $^{19}$F n.m.r. spectrum showed that the reaction was only ≈ 95% complete after 24 h at room temperature. After 20 more hours of stirring at room temperature, 10 ml of 4 m HCl was added dropwise. A thick precipitate appeared that changed quickly to an oil. The oily lower layer was removed (4.69 g), and the remaining phase was extracted (3 x 1 ml CH$_2$Cl$_2$), the extracts were combined with the first obtained oil, and the CH$_2$Cl$_2$ was then distilled off at atmospheric pressure through a 12 cm Vigreux column. Distillation of the residue at 39-42°C (9 torr) yielded 3.98 g (59.7 %) of product. A sample for analysis was obtained in at 41-42°C. The pot residue (0.79g) had the same i.r. spectrum as the main product.

$^1$H n.m.r. (CDCl$_3$, Si(CH$_3$)$_4$, FF$_4$SCH$_2$BrCH$_2$OCH$_3$': c = cis, t = trans): $\delta_1$ = 4.00 ppm, s, 3.10 H (CH$_3$); $\delta_2$ = 4.55 ppm (center), m, 2.07 H, (CH$_2$); $\delta_3$ = 6.10 ppm, septet, 1.00 H, (CHBr). $J_{CH_3-B} = 5.75$ Hz; $J_{Ha-Hb}$ = 11.50 Hz; $J_{Ha-Hbc} = 0$ Hz; $J_{HbC-Hbt} = -$.

$^{19}$F n.m.r. (neat sample, CCl$_3$F external): AB$_4$-system;
$\phi_A = 78.67$ ppm, 9 lines, 1.0 F; $\phi_B = 57.22$ ppm, d-m, 3.8 F; $J_{AB} = 146.6$ Hz.

Infrared spectrum (neat sample, KBr, cm$^{-1}$): 3004, w; 2942, w; 2901, w, sh; 2840, w; 1471, w; 1458, w; 1389, w; 1298. vw, br; 1263, vw; 1210, w, sh; 1196, w; 1127, m; 1110, w-m, sh; 1075, w; 1054, w; 965, w; 936, vw; 923, vw; 842, vs; 766, w, br; 713, w-m; 663, sh, w; 654, w-m; 597, m; 587, w; 571, w; 551, w; 528, vw; 511, vw.

Mass spectrum (c.i., mass, species, % > 1): (for Br-fragments only listing of $^{79}$Br) 233, M-CH$_3$O$^+$, 1.7; 185, (M-Br)$^+$, 3.6; 165, (M-Br-HF)$^+$, 2.2; 139, (M-HBr-CH$_3$OCH$_2$)$^+$, 95.7; 138, C$_2$H$_3$$^{79}$Br$^+$, 4.4; 135, (M-Br-CH$_3$O-F)$^+$, 1.8; 127, SF$_5^+$, 6.5; 125, (M-Br-3HF)$^+$, 5.5; 123, (M-Br-3HF-2H)$^+$, 1.2; 109, (M+H-Br-3HF-CH$_3$)$^+$, 1.7; 108, SF$_4^+$, 2.4; 107, FSC$_3$H$_4$O$^+$, 3.5; 106, FSC$_3$H$_3$O$^+$, 2.2; 105, FSC$_3$H$_2$O$^+$, 2.8; 101, CSF$_3^+$, 4.0; 95, C$_2$H$_4$FOS$^+$, 3.5; 94, C$_2$H$_3$FOS$^+$, 1.3; 93, C$_2$H$_2$FOS$^+$, 3.3; 91, C$_2$FOS$^+$, 1.1; 89, SF$_3^+$, 21.9; 87, SC$_2$OCH$_3^+$, 1.7; 79, FSC$_2$H$_4^+$, 1.5; 77, FSC$_2$H$_2^+$, 7.9; 76, FSC$_2$H$^+$, 1.3; 75, FSC$_2^+$, 3.9; 70, SF$_2^+$, 2.8; 69, CF$_3^+$, 1.5; 67, FOS$^+$ (rearr.), 1.4; 63, FSC$^+$, 6.1; 61, CHOS$^+$, (rearr.), 1.7; 59, C$_2$H$_3$S$^+$, 4.2; 58, C$_2$H$_2$S$^+$, 11.9; 57, C$_2$HS$^+$, 2.4; 56, C$_2$S$^+$, 1.9; 55, C$_3$H$_3$O$^+$, C$_3$F$^+$, 3.1 and 1.9.

Anal. calcd for C$_3$H$_6$BrF$_5$O$_3$: C, 13.59; H, 2.28; Br, 30.15; F, 35.8; S, 12.10. Found: C, 13.73; H, 2.33; Br, 30.04; F, 35.5; S, 12.05.
A mixture of 3.34 g silver acetate (19.8 mmol), 10 ml of glacial acetic acid and 4 g of \( \text{F}_5\text{SCH}_2\text{CH}_2\text{Br} \) (17.0 mmol) was refluxed in a 25 ml round bottomed flask, while being magnetically stirred, for 36h. A Drierite tube was fitted to the top of the reflux condenser. The insoluble salt was filtered off and rinsed with 10 ml of diethyl ether. Water was added to the filtrate, and the bottom layer was separated and neutralized with \( \text{NaHCO}_3 \). Extraction of the aqueous phase with ether (3 x 10 ml), combining the extracts with the original ether layer, adding 5 ml of water and small amounts of solid \( \text{NaHCO}_3 \) until bubbling ceased completely, removing the top layer (separatory funnel), drying (\( \text{MgSO}_4 \)) and distilling the product at atmospheric pressure resulted 4 cuts, of which the first two (-48°C) were ether (g.c.), and the cuts from 48-98°C, and from 98-112°C, as well as the pot residue showed two bands in g.c. (\( R_f = 13.3 \) min = \( \text{F}_5\text{SCH}_2\text{CH}_2\text{OC}(\text{O})\text{CH}_3 \), 61%; \( R_f = 5.4 \) min = \( \text{F}_5\text{SCH}_2\text{CH}_2\text{Br} \) (39%), Carbowax, 20%, 1m, 90°C). A total of 1.58 g of product was obtained. A sample for analysis was obtained with preparative gas chromatography (20% Carbowax, 1m, 90°C). The compound is a colorless liquid with a pleasant odor.

\(^1\text{H} \) n.m.r. spectrum (CDCl\(_3\), external \( \text{Si(CH}_3)_4 \)):

- \( \delta_1 = 2.56 \) ppm, s, 3.00 H (CH\(_3\));
- \( \delta_2 = 4.40 \) ppm, m (≈ sextet), 1.96 H (SF\(_5\)CH\(_2\));
- \( \delta_3 = 4.98 \) ppm, 1.92 H (SF\(_5\)CH\(_2\)CH\(_2\));
- \( J_{23} = \)
5.7 Hz; $J_{2B} \approx 7.3$ Hz (B refers to the AB₄ system of SF₅).

$^{19}$F n.m.r. spectrum (CDCl₃, CCl₃F external reference): AB₄ system, $\phi_A = 83.57$ ppm, 9 lines, 1.00 F; $\phi_B = 66.57$ ppm, d-m, 3.94 F; $J_{AB} = 144.2$ Hz.

I.r. spectrum (neat sample, KBr, cm⁻¹): 3039, vw; 2966, vw, br; 2930, w; 2858, w; 1750, s; 1465, w; 1449, w; 1421, w; 1393, w; 1370, w-m; 1334, vw; 1283, w, sh; 1240, m-s; 1134, w, sh; 1082, w-m; 1052, m-s; 979, w; 947, vw; 835, vs; 816, s; 751, vw, br; 711, vw, br; 672, vw; 639, w; 605, w; 587, w; 566, w; 522, w; 534, vw; 520, vw; 493, w.

Mass spectrum (c.i., mass, fragment, % > 1): 215, (M+H)+, 2.2; 195, (M+H-HF)+, 4.6; 155, F₅SCH₂CH₂⁺, 1.1; 153, F₅SC₂H₂⁺, 1.4; 135, C₂H₃F₄S⁺, 23.1; 133, C₂HF₄S⁺, F₃SCH₂CH₂O⁺, 100.0; 127, SF₅⁺, 8.7; 122, F₄SCH⁺, 2.4; 109, HSF₄⁺, 1.4; 97, C₂H₃F₂S⁺, 1.6; 91, C₂FOS⁺, 2.4; 89, SF₃⁺, 51.9; 87, C₂H₄OC(O)CH₃⁺, 100; 86, C₂H₃OC(O)CH₃⁺; 4.7; 75, C₂H₃OS⁺, 15.6; 73, C₂HOS⁺, 1.2; 70, SF₂⁺, 5.0; 67, C₄H₅O⁺, 1.7; 65, C₄HO⁺, 1.4; 63, CFS⁺, 1.9; 61, C₂H₅S⁺, 24.0; 59, CH₃COO⁺, 1.5; 57, CHCOO⁺, 1.4; 55, C₃H₃O⁺, 1.5.

Anal. calcd for C₄H₇F₅O₂S: C, 22.43; H, 3.29; F, 44.4; S, 14.97. Found: C, 22.72; H, 3.32; F, 44.0; S, 14.91.

Reaction of F₅SCHBrCH₂Br with AgCH₃COO in CH₃COOH:

A mixture of 4.81 g of F₅SCHBrCH₂Br (15.3 mmol), 5.30 g of AgCH₃COO (28.5 mmol) and 15 ml of glacial acetic acid in a 50 ml round bottomed flask, fitted with a short reflux
column and a Drierite tube, was refluxed for 4 days at an oil bath temperature of 120-130°C. A $^{19}$F n.m.r. spectrum taken after 3 days of refluxing showed the presence of three SF$_5$ compounds, one of them being the starting material. The volatile materials were vacuum transferred out, 10 ml of water was added, and the precipitated liquid (2.79 g) was separated. Gas chromatography showed that only little CH$_3$COOH was present. The product had a faint ester-like odor, but a sample of the main product, as collected by preparative g.c. (30% SE-30, 140°C, 3m), was identical with the compound SF$_5$CBr=CH$_2$, as shown by infrared spectroscopy. Yield = 72%.

F$_5$SCH$_2$CHBrCOOC$_2$H$_5$:

Crude SF$_5$Br (41.42 g, 200.1 mmoles) was transferred to a 75 ml steel vessel held at -196°C, then 19.78 g (197.8 mmol) of ethyl acrylate, dried over MgSO$_4$, were condensed in at the same temperature and the vessel was allowed to attain slowly room temperature. When the vessel had reached (10-15)°C, it was noticed that the temperature climbed rapidly to about 50 °C, whence it dropped slowly to room temperature. The vessel was kept at room temperature for 2h, then warmed to 55 °C for 19h. The contents of the vessel were then condensed out (58.93g). From the vessel 4.72 g of a viscous liquid could also be poured out; the reaction had been run in the same vessel before, without noticing the
non-volatile oil, total mass balance = 104%. The combined products from the bomb were distilled with a 12 cm Vigreux column (3 torr), yielding one fraction, 21.55 g, at (66.5-67.8)°C. Upon raising the temperature the residue turned yellow; the pressure was lowered to 0.005-0.020 torr (dynamic vacuum) and 3.07 g of a fraction boiling > 110 °C (out of thermometer range) was collected. The cold trap contained 1.08 g of material, which was mainly ethyl acrylate (1H n.m.r.). Yield = 35.5% of theory.

1H n.m.r. spectrum (neat sample, external Si(CH3)4): δ = 1.33 ppm, t, (CH3), relative area = 3.00 H; δ = 4.30 ppm, (CH3CH2), J = 7.2 Hz; δ = 3.6 - 5.0 ppm, m (SF5CH2CH + CH3CH2), total area (quartet + multiplet) = 5.04 (theoretically 5.00).

19F n.m.r. spectrum (CHCl3, external CFCl3): AB4 spectrum, δA = 81.49 ppm, δB = 65.83 ppm, d-m; JAB = 147.4 Hz. Neat sample: δA = 81.00 ppm, δB = 65.17 ppm.

I.r. spectrum (neat sample on KBr plates, cm⁻¹): 3042, w; 2988, w-m; 2946, w-vw; 2912, vw; 2881, vw; 1750, vs; 1477, w,sh; 1469, w-m; 1456, w-m; 1417, m; 1398, m; 1378 m-s; 1356, m; 1311, m-s; 1262, m-s; 1241, s; 1206, w,sh; 1187, m-s; 1158, m-s; 1116, w,sh; 1097, w; 1089, w,sh; 1034, m; 1018, m; 988, m; 965, w-vw; 926, w; 877, s-vs; 851, vs; 832, vs; 802, m-s; 740, w,br; 698, vw; 673, vw; 648, vw; 623, vw; 602, m; 577, s; 564, m; 481, w.

Mass spectrum (e.i., 70 eV, mass, fragment, % > 1):
179, C₅H₇BrO₂⁺, 6.3; 155, SF₅C₂H₄⁺, 18.6; 153, SF₅C₂H₂⁺, 28.1; 151, S₅C₂⁺, 2.1; 135, C₂⁷⁹BrO₂⁺, 3.2; 135, SF₄C₂H₃⁺, 2.9; 127, SF₅⁺, 13.5; 126, SF₃C₃H⁺, 1.5; 125, SF₃C₃⁺, 15.0; 109, C₃H₃F₂S⁺, 9.1; 109, SF₄⁺, 66.2; 107, C₃H₄FOS⁺, 12.5; 106, C₃H₃FOS⁺, 65; 105, C₃H₂FOS⁺, 10.2; 99, C₅H₇O₂⁺, 9.2; 89, SF₃⁺, 32.7; 85, C₄H₅O₂⁺, 2.1; 73, C₃H₅O₂⁺, 10.6; 71, C₃H₃O₂⁺, 2.5; 70, SF₂⁺, 5.3; 67, C₄H₃O⁺, 3.6; 56, C₃H₃O⁺, C₂S⁺ (two fragments), 4.9, 3.7; 55, (?), 30.9; 55, C₃H₃O⁺, 100.0; 54, C₃H₂O⁺, 5.2; 53, C₃HO⁺, 6.3; 51, SF₂⁺, 2.1.

**Anal. calcd** for C₄H₈BrF₅O₂S: C, 19.56; H, 2.63; Br, 26.02; F, 30.9; S, 10.45. Found: C, 20.10; H, 2.58; Br, 28.68; F, 29.4; S, 9.86.

**Attempted dehydrohalogenation of F₅SCH₂CHBrCOOC₂H₅:**

No tractable products were obtained from the reaction with aqueous KOH. Both Br⁻ and F⁻ could be detected in the solution after a while. It was thus tried to carry out the reaction with organic nitrogen bases:

1. With N,N-dimethyl aniline:

To 5.00 g of F₅SCH₂CHBrCOOC₂H₅ (16.3 mmoles) in a 50 ml round bottomed flask, equipped with a magnetic stirring bar, 2.99 g of N,N-dimethylaniline (24.7 mmoles) were added at room temperature. A 24 cm reflux condenser was then fitted on the flask and the stirred mixture was slowly heated in an oil bath. When the oil bath temperature had reached 80-90 °C a reaction occurred suddenly and much
material was blown out through the condenser. The dark blue residue was kept at that temperature for another 15 minutes. After the mixture had cooled down to room temperature, it was treated repeatedly with CHCl₃ until the rinses were only slightly blue. There was a large amount of almost black tarry residue. The chloroform solution was repeatedly washed with 5% HCl until the acid layer was almost colorless. The extract was dried (MgSO₄), concentrated (rotary evaporator) and finally transferred on a vacuum line, yielding a small amount of an almost colorless oily product. ¹H n.m.r. spectrum (CDCl₃): d-t, centered at δ = 1.77 ppm, CH₃, intensity ≈ 1:1; two doublets, δ = 6.66 ppm, δ = 7.27 ppm and a multiplet, δ = 4-5 ppm. ¹⁹F n.m.r. (CDCl₃): showed only starting material to be present. The product that was formed was apparently a mixture of starting material and a compound that does not contain an SF₅-group. The reaction was repeated by heating slowly a similar base - starting material mixture (C₆H₅N(CH₃)₂); the result was the same.

2. Reaction with diazabicyclononane:

Reaction of diazabicyclononane (neat) with the ester resulted in charring and destruction of all starting material. The ¹⁹F n.m.r. spectrum of this mixture showed a singlet at 73.6 ppm, which is probably SF₄. (δ = 72.7 ppm for SF₄ in CDCl₃). Employing ethereal solutions of both the ester and diazabicyclononane at -10 °C resulted in formation of a brown solution. The reaction was thus finally run at -78 °C:
In a 50 ml round bottomed flask equipped with magnetic stirring bar, 1.0 g of SF₅CH₂CHBrCOOC₂H₅ (3.26 mmoles) was dissolved in 15 ml anhydrous ether. A dropping funnel containing 0.40 g diazabicyclononane (Aldrich) (3.26 mmoles) in 2ml of ether (anhydrous) with a Drierite tube was fitted on the flask and it was cooled in an acetone-dry ice bath to -78 °C. The base solution was added dropwise during 10 minutes. A white precipitate formed immediately upon addition of the base. When addition was completed the mixture was allowed to attain slowly room temperature (≈ 45 min). The white precipitate had dissolved during the warming up process, leaving an almost colorless clear solution. It was kept in the refrigerator overnight, when a light brown oil had separated the next morning (i.r.spectrum: no SF₅, 0.22g). The remaining solution was freed of solvent by distillation, leaving 0.82 g of a brownish oil. It showed only a weak SF₅ band in the i.r.spectrum and traces of starting material in the fluorine n.m.r.spectrum. No other fluorine compound was found. The ¹H n.m.r.spectrum shows as the only product the product which was obtained earlier in the attempted dehydrohalogenation with N,N-dimethylaniline and some diethyl ether. This material was identified as 2-bromoacrylic acid by comparison with an authentic sample. The product polymerized rather quickly to a rubbery translucent mass.
Crystal Data and Methods used in X-ray analysis:

\([\text{HN(C}_2\text{H}_5)_3]^+\text{[F}_5\text{SC(SO}_2\text{F)COOCH}_3]^-\):

Monoclinic, \(P2_1/n\), \(a = 8.758(2)\,\text{Å}, b = 9.645(2)\,\text{Å}, c = 19.167(4)\,\text{Å}\), \(\beta = 97.92(3)^\circ\); \(V = 1603.6(6)\,\text{Å}^3\); \(Z = 4\); density = 1.588 g/cm\(^3\); crystal size = 0.5x0.4x0.4 mm; reflexions collected: 2998; independent reflexions: 1991 (\(R_{\text{int}} = 1.81\%\)); observed reflexions: 1683 (\(F > 3.0\sigma(F)\)); \(R = 4.62 \%\) (obs. data), \(R = 5.59 \%\) (all data); hydrogen atoms: riding model, fixed isotropic \(U\).

Diffractometer: Siemens-Stoe AED2 (MoK\(\alpha\), graphite monochromator); program used: SHELXTL PLUS (PC Version).

\([\text{F}_5\text{SC(SO}_2\text{F)CH}_2]_2\text{O}\):

Tetragonal, \(P4\), \(a = 11.069(2)\,\text{Å}, c = 6.0780(10)\,\text{Å}\); \(V = 744.7(2)\,\text{Å}^3\); \(Z = 2\); density = 2.187 g/cm\(^3\); crystal size = 0.2x0.2x0.5 mm; reflexions collected: 2164; independent reflexions: 970 (\(R_{\text{int}} = 1.56 \%\)); observed reflexions: 898 (\(F > 3.0\sigma(F)\)); \(R = 3.82 \%\) (obs. data), \(R = 4.22 \%\) (all data); hydrogen atoms: riding model, fixed isotropic \(U\). Apparatus and methods as above.

\(\text{F}_5\text{SC(SO}_2\text{F)CON(C}_2\text{H}_5)_2\):

Monoclinic, \(P21/a\) (#14), \(a = 10.945(2)\,\text{Å}, b = 8.932(2), c = 12.564(1)\,\text{Å}\), \(Z = 4\); density 2.19 g/cm\(^3\); crystal size: 0.3x0.25x0.25 mm; total reflexions: 796. \(R_w = 0.079\).

Diffraktometer: Rigaku AFC6 (MoK\(\alpha\) graphite monochromator).
[F₅SCH₂CHO]₃:

Rhombohedral, R₃m, a = 10.266(2) Å, c = 26.336(8) Å; V = 2404 Å³, Z = 18; density = 2.13 g/cm³; crystal size: 0.5 mm maximal dimension; total reflexions: 916; Rₓ(F₀) = 0.0763, Diffractometer: upgraded Syntex P₂₁ (MoKα graphite monochromator); program used: Nicolet SHELXTL (version 5.1, 1985).
REFERENCES

PART I


24. References 18 and 23.


39. These values are: \( F_5S-Cl \approx 2.0 \text{ eV}, F_5S-F \approx 3.2-3.4 \text{ eV}; \) Gmelins Handbuch der anorganischen Chemie, Bd. 2 (1978); Springer-Verlag.


53. Described in the second half of this manuscript.


55. F$_5$SCF$_2$SO$_2$F has been made recently by the direct fluorination (with F$_2$) of F$_5$SCHF$_2$SO$_2$F: G. L. Gard, A. Waterfeld, R. Mews, J. Mohtasham, R. Winter, to be published.

56. Reference 27.


71. It will be noted that electron density as an intensive quantity cannot be redistributed. This terminology, although common, is clearly incorrect.


78. The iterative program LAOCN5 (quantum chemical exchange service, Bloomington, Indiana) was used to calculate the spectral parameters; the spectrum was then generated with the program PCPMT (Serena Software, Bloomington, Indiana).


83. In CH₂=SF₄ this band is found at 1167 cm⁻¹, S. Sunder, H.H. Eysel, G. Kleemann, K. Seppelt, Inorg. Chem., 18, 3208 (1979). In CH₃CH=SF₄ the band is found at 1200 cm⁻¹, B. Pötter, K. Seppelt, Inorg. Chem., 21, 3147 (1982).

84. The crystal structures of (F₅SCH(SO₂F)CH₂)₂O [29] and F₅SC(SO₂F)COOCH₃⁻HN(C₂H₅)₃⁺ [32b] were kindly determined by Dr. Mathias Noltemeyer, Inst. f. anorganische Chemie, Universität Göttingen, and of F₅SCH(SO₂F)CON(C₂H₅)₂ [27] by Dr. D.A. Keszler, Oregon State University, Corvallis, Oregon.


PART II


3. Examples are given in ref. 5.


6. (F₃C)₂PF₂ was first described by W. Mahler, Inorg. Chem., 2, 230 (1963), and an example and further references are given in ref. [5b].


14. These examples are taken from ref 7, pages 93 and 94.
16. Ref. 2, ref 165.
20. Ref. 17 b.) p. 182.


40. The program PCPMR (Serena Software, Bloomington, Indiana) was used. Approximate values for J₆F², JₓF¹ and φₓ⁻φ₆² were extracted from the spectrum and successively changed.


PART III


45. S. Akiyoshi, K. Okuno, J. Am. Chem. Soc. 74, 5759 (1952) (JCH2CH(OR)2); P. Bedoukian, J. Am. Chem. Soc. 66, 651 (1944) (BrCH2CH(OR)2).


49. AcOCH=CHCOOC2H5 was first obtained by H.v. Pechmann, Chem. Ber. 25, 1040 (1892). It was made according to P. Freeman et al., J. Org. Chem. 39, 546 (1974), but without cooling. Both Freeman et al. and M. Perezamador, M. Salmon and F. Walls, Chem. Abst. 67, 43386 (1967) describe β-acetyl ethylacrylate as yellow liquids. It was found that slow distillation resulted in a completely colorless oily product.


APPENDIX

COMPOUND REFERENCE TABLE

1. \( \text{F}_5\text{S} \text{C} = \text{CF}_2 \text{SO}_2 \)
   
2. \( \text{F}_3\text{C} \text{C} = \text{C} = \text{OSO}_2 \)

3. \( \text{F}_3\text{C} \text{C} = \text{C} = \text{O} \)
   
4. \( \text{F}_3\text{C} \text{C} = \text{H} \)

5. \( \text{F}_3\text{C} \text{C} = \text{C} = \text{O} \)
   
5a. \( \text{F}_5\text{S} \text{C} = \text{C} = \text{O} \)

6. \( \text{F}_3\text{C} \text{O} \text{C} = \text{O} \text{SO}_2 \)

7. \( \text{F}_3\text{C} \text{C} = \text{C} = \text{F} \)

8. \( \text{F}_4\text{S} = \text{C} = \text{COF} \)
   
8a. \( \text{F}_4\text{S} = \text{C} = \text{COF} \)

9. \( \text{(CF}_3\text{)}_2\text{C} = \text{C} - \text{C} - \text{C} - \text{C} - \text{O} - \text{C} = \text{O} \)

10. \( \text{F}_3\text{C} \text{C} = \text{C} = \text{COOR} \)

11a. \( \text{F}_3\text{C} \text{COOR} \)

b. \( \text{Br} \)

11b. \( \text{F}_3\text{C} \text{COOR} \)

b. \( \text{Cl} \)

12. \( \text{F}_3\text{C} \text{C} = \text{C} = \text{Cl} \)

13. \( \text{F}_3\text{C} \text{H} \text{COOR} \)

14. \( \text{(CF}_3\text{)}_2\text{C} = \text{S} = \text{O}_2\text{F}^{\ -} \)

15. \( \text{(CF}_3\text{)}_2\text{CHO}_2\text{F}^{\ -} \)

16. \( \text{CF}_2 = \text{C} = \text{SO}_2\text{F}^{\ -} \)

17. \( \text{Cs}^+ \text{(CF}_3\text{)}_2\text{C} = \text{S} = \text{O}_2\text{F}^{\ -} \)

18. \( \text{F}_3\text{B} \cdot \text{N} (\text{C}_2\text{H}_5)_3 \)
47 $\text{F}_3\text{CCFCF}_2$

$\text{O}$

48 $(\text{F}_3\text{CCFCF}_2)_n$

49 $\text{O}$

50 $\text{CF}_2[\text{CFOCF}_2]_{n-1}\text{CF}_2\text{SO}_2\text{F}$

51 $\text{F}_5\text{SOCF}_2\text{CFO}$

52 $\text{F}_5\text{S}$

53 $\text{C}^\circ\text{C}^\circ\text{F}$

54 $\text{C}^\circ\text{C}^\circ\text{F}$

55 $\text{F}_5\text{SCF}_2\text{C}=\text{CF}_2$

$x = \text{H, halogen}$

56 $\text{F}_5\text{SCF}_2\text{CX}=\text{CF}_2$

56a $\text{F}_5\text{S(CF}_2)_n\text{CX}=\text{CF}_2$

57 $\text{FO}_2\text{SCF}_2\text{CF}=\text{CF}_2$

58 $\text{F}_5\text{SCH}_2\text{CH}_2\text{Br}$

59 $\text{F}_5\text{SCH}_2\text{CH}_2\text{OC(O)}\text{CH}_3$

60 $\text{F}_5\text{SCHBrCH}_2\text{Br}$

61 $\text{F}_5\text{SCH}_2\text{CH}_2\text{OC(O)}\text{CH}_3$

62 $\text{F}_5\text{SCHBrCH}_2\text{OC(O)}\text{CH}_3$

64 $\text{F}_5\text{SCF}_2\text{C}=\text{CF}_2$

65 $\text{Cl}$

66 $\text{F}_5\text{SCF}_2\text{CFCF}_2$

67 $\text{F}_5\text{SCF}_2\text{CF}_2\text{J}$

68 $\text{F}_5\text{SCF}_2\text{CF}_2\text{O}$

69 $\text{F}_5\text{SCF}_2\text{CF}_2\text{O}$

69 $\text{F}_5\text{SCF}_2\text{CF}_2\text{O}$