Trifluoromethyl sulphur pentafluoride, SF$_5$CF$_3$ : atmospheric chemistry and its environmental importance via the greenhouse effect

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Trifluoromethyl sulphur pentafluoride, SF₅CF₃: atmospheric chemistry and its environmental importance via the greenhouse effect

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Abstract  One molecule of SF₅CF₃, an adduct of the SF₅ and CF₃ free radicals, causes more global warming than one molecule of any other greenhouse gas yet detected in the atmosphere, i.e. it has the highest per molecule radiative forcing of any greenhouse pollutant, and the value of its global warming potential is only exceeded by that of SF₆. Using tunable vacuum-UV radiation from a synchrotron and coincidence spectroscopy, the strength of the central S−C bond in SF₅CF₃ is determined to be 3.86 ± 0.45 eV or 372 ± 43 kJ mol⁻¹, and this molecule is very unlikely to be removed from the earth’s atmosphere by UV photolysis in the stratosphere. Complementary laboratory-based experiments have shown that the main sink route of this greenhouse gas is low-energy electron attachment in the mesosphere, with Lyman-α photodissociation at 121.6 nm being only a minor channel. By comparison with data for SF₆, the lifetime of SF₅CF₃ in the earth’s atmosphere is estimated to be ca. 1000 years. The principal reason for the current low level of concern about the impact of SF₅CF₃ on our environment is that the concentration levels are still very low, at the sub parts per trillillion level. The high growth rate of ca. 6% per annum, however, should cause concern for policymakers.

Keywords: SF₅CF₃, greenhouse gas, atmospheric lifetime, synchrotron radiation, photoionisation, TPEPICO, mesospheric reactions, ion-molecule, electron attachment, Lyman-α radiation
1. Introduction

Trifluoromethyl sulphur pentafluoride, SF$_5$CF$_3$, has recently been termed the super greenhouse gas. This claim to fame arises because, per molecule, it makes more contribution to global warming via the greenhouse effect than any other molecule yet detected in the atmosphere. Within this headline phrase, however, caution must be exercised because the amounts of SF$_5$CF$_3$ in the earth’s atmosphere are still very low at the sub parts per trillion level by volume, ca. 0.15 pptv; 1 pptv is equivalent to a number density of $2.46 \times 10^7$ molecules cm$^{-3}$ for a pressure of 1 bar and a temperature of 298 K. By contrast, carbon dioxide has a current concentration in the earth’s atmosphere of ca. 370 parts per million by volume, over nine orders of magnitude greater than that of SF$_5$CF$_3$. Despite a much lower environmental impact per molecule, CO$_2$ makes a much greater impact to global warming, simply because its concentration is much higher. Nevertheless, the Kyoto protocol of 1997, now ratified as an International Treaty in 2005, to limit emissions of greenhouse gases has meant that scientists and industry are now legally bound to determine the most environmentally-benign and cost-effective materials in large-scale industrial applications. It is therefore essential to understand the reactive and photochemical properties of molecules used in industry that are emitted into the atmosphere, even gases that at first sight might seem benign and harmless. It is within this near-worldwide political consensus for control of global warming that one can understand the huge upsurge in interest in SF$_5$CF$_3$ since its potential for global warming was first raised by Sturges et al. [1] five years ago.

SF$_5$CF$_3$ is a gas at room temperature, with a boiling point of 253 K and an enthalpy of vapourisation of 20.2 kJ mol$^{-1}$ over the temperature range 223–253 K [2], and such thermochemical data has been available for some time. The paper by Sturges et al. [1], however, was the first to describe detection of SF$_5$CF$_3$ in the earth’s atmosphere. Its source was believed to be anthropogenic in nature, and most likely a breakdown product of SF$_6$ in high-voltage equipment. Since the trends in concentration levels of SF$_6$ and SF$_5$CF$_3$ have tracked each other very closely over the last 30-40 years, Sturges et al. suggested that SF$_5$CF$_3$ has mainly been produced in the electronics industry via the recombination of SF$_5$ and CF$_3$ free radicals. This claim has since been disputed [3]. Infrared (IR) absorption measurements showed that SF$_5$CF$_3$ has the highest radiative forcing per molecule of any gas found in the atmosphere to date (0.57 W m$^{-2}$ ppb$^{-1}$, a figure since updated to 0.60 ± 0.03 W m$^{-2}$ ppb$^{-1}$ [4,5]). Antarctic firn measurements suggested that it has grown from a concentration of near zero in the late 1960s to ca. 0.12 pptv in 1999 with a current growth rate of ca. 6% per annum, and stratospheric profiles suggested that the lifetime of this species in the atmosphere is between several hundred and a few thousand years. It was estimated that the global warming potential (GWP) of SF$_5$CF$_3$ is 18,000 relative to CO$_2$, with only SF$_6$ having a higher value. Sturges et al. concluded that, whilst still a relatively minor problem, nevertheless it was important to control the source(s) of SF$_5$CF$_3$ into the atmosphere in order to guard against an undesirable accumulation of this potent greenhouse gas. These comments remain as true now as they were then.
Furthermore, in historical terms, the story of the chlorofluorocarbons, and their ‘conversion’ from industrially-produced benign molecules to serious ozone-depleting molecules in the stratosphere over a period of less than 20 years, is still fresh in the memory of many atmospheric scientists. Small problems have a tendency to become big problems in atmospheric science. Thus, although the best estimate two years ago was that SF₅CF₃ only contributes 0.003 % to the total greenhouse effect [6], it is not surprising that there has been huge interest in the reactive and photochemical properties of SF₅CF₃.

The purpose of this chapter is to review the microscopic properties of SF₅CF₃, most of them measurements and calculations made since Sturges et al. published their paper. Emphasis will be on those measurements that pertain to the atmospheric properties of SF₅CF₃ and the processes that remove it from the earth’s atmosphere. Only limited descriptions of the macroscopic properties of SF₅CF₃ will be given. Since the readership of this chapter will be wide, the terms microscopic and macroscopic may mean different things to different readers. As a chemical physicist, I define a microscopic property to be that of one molecule of the system under study (e.g. a spectroscopic, kinetic, photochemical or thermodynamic property). A macroscopic property relates to the system as a whole (e.g. industrial production sources of SF₅CF₃, the geographical properties that determine the transport of SF₅CF₃ through the atmosphere via convection and diffusion). The chapter will conclude with a review of the properties of SF₅CF₃ that determine its lifetime in the earth’s atmosphere. The lifetime of a pollutant is another term that can mean different things to different scientists, and I will bring together all the many interpretations of this term in the context of the physical, chemical and geographical properties of SF₅CF₃ in our atmosphere.

2. Structure, spectroscopy and thermochemistry of SF₅CF₃

It is of some surprise to note that SF₅CF₃ is not a new molecule, and its microwave spectrum was recorded by Kisliuk and Silvey [7] as long ago as 1952. The S–C bond distance was determined to be 0.186 nm, and in 1985 all the geometrical parameters for SF₅CF₃ were reported by Marsden et al. [8] from an electron diffraction study. Figure 1 shows an unpublished structure of the highest occupied molecular orbital (HOMO) of SF₅CF₃ by Knowles [9] using the MOLPRO ab initio suite of programmes. The particular points to note are the substantial dipole moment of the molecule, 0.95 Debye, the significant σ-bonding electron density along the S–C bond, the S-C bond length of 0.187 nm in almost exact agreement with the microwave study, and the FSF and FCF bond angles of approximately 90° or 109.3°, respectively. SF₅CF₃ can therefore either be regarded as a perturbed SF₆ molecule in which one fluorine atom has been replaced by a CF₃ group of tetrahedral symmetry, or as a perturbed CF₄ molecule in which one fluorine has been replaced by an SF₅ group with octahedral symmetry. This will form a recurring theme of this chapter. The only major difference from SF₆ and CF₄ is that the HOMO of these two
molecules is a F 2p\pi non-bonding in character, whereas the HOMO of SF₅CF₃ is S–C σ-bonding in character. The gas-phase IR and liquid-phase Raman spectra over the extended range of 100-4000 cm⁻¹ were reported at low resolution over 40 years ago [10,11]. These measurements have been made at much higher resolution in the last three years, and all the important IR-active vibrations in the atmospheric window region of ca. 700-1500 cm⁻¹ have been determined [12-14]. Of particular note is the determination by two groups of the integrated absorption coefficient over this range as a function of temperature between 200 and 300 K [13,14], both studies demonstrating that it showed little variation with temperature. There were no other spectroscopic details (e.g. UV/visible electronic, vacuum-UV photoionisation) on this molecule before the Sturges et al. [1] paper was published. Since 2000, there have been a large number of new studies, some of direct importance to the atmospheric properties of SF₅CF₃, some of more general and pure scientific interest. They include VUV and XUV absorption [4,15-19], valence photoelectron [4,15,20-22], electron energy loss spectroscopy [4,23], electron scattering [24-26], protonation [27] and photofragmentation studies [20-22,28].

As with the spectroscopic situation, there was scant information on the thermochemistry of SF₅CF₃ prior to 2000. The JANAF tables of 1998 [29] quote an enthalpy of formation of −1700 ± 63 kJ mol⁻¹ at 0 K, −1717 ± 63 kJ mol⁻¹ at 298 K, although these data are indirect and it is not clear how these numbers are derived. The strength of the SF₅–CF₃ σ-bond will also be determined by the enthalpies of formation of the SF₅ and CF₃ radicals, where improved values since the tables were published are now available. Since the strength of this bond is related to possible photochemistry of SF₅CF₃ that may occur in the stratosphere, a direct determination of ΔfH° (SF₅CF₃) is therefore essential to explain any possible atmospheric reactions. Such an experiment is described in Section 5.3. As with experimental spectroscopic studies, there have been many theoretical studies on SF₅CF₃ since 2000 [30-34]. Their main aim has been to determine properties that relate to the atmospheric chemistry of SF₅CF₃, such as integrated infrared intensities, vibrational frequencies, and thermochemical and structural parameters.

3. The greenhouse effect and SF₅CF₃: how serious is the problem?

The greenhouse effect is usually associated with small polyatomic molecules such as CO₂, H₂O, CH₄, N₂O and O₃. The ‘natural’ greenhouse gases, notably CO₂ and H₂O, have been responsible for hundreds of years for maintaining the temperature of the earth at ca. 290 K, suitable for habitation. The ‘enhancing’ greenhouse gases, mainly CH₄, N₂O and O₃, have concentrations in the atmosphere which have increased in the last 50-100 years, have IR absorptions in the atmospheric window region where CO₂ and H₂O do not absorb, and are believed to be the main culprits for global warming. It is now clear, however, that there are larger polyatomic gases of low concentrations in the atmosphere which can contribute significantly to global warming because they possess exceptionally strong IR absorption bands.
in regions where other greenhouse gases do not absorb. Two notable examples are SF₆, which has a GWP of 22,200 relative to CO₂ over a time horizon of 100 years [35] and SF₅CF₃, the subject of this review. In qualitative terms, the physical and chemical properties that are necessary for a molecule to have a large GWP are: (a) it must absorb IR region in the black-body range of the earth’s emission, \( ca. 5–25 \ \mu\text{m} (2000–400 \ \text{cm}^{-1}) \) with a peak at 15 \( \mu\text{m} (667 \ \text{cm}^{-1}) \), in regions where CO₂ and H₂O do not absorb; in practice this means many C–F and C–Cl stretching vibrations around 900-1100 cm\(^{-1}\) contribute strongly, (b) the integrated absorption intensities of the IR-active vibrations must be strong due to large dipole moment derivatives, (c) the molecule must have a long lifetime (defined in Section 7) in the earth’s atmosphere; it must not be photodissociated by solar UV radiation in either the troposphere (\( \lambda > 290 \ \text{nm} \)) or the stratosphere (\( 200 < \lambda < 290 \ \text{nm} \)), and it must not react with either the OH or O*(1D) free radicals. Furthermore, a greenhouse gas whose concentration is increasing rapidly due to man’s activity, whilst not affecting directly an integrated absorption coefficient or the GWP value, is a special cause for concern for policymakers in Government.

SF₅CF₃ appears to obey all these criteria, and Table 1 shows data for five greenhouse gases which demonstrate how serious or otherwise the presence of SF₅CF₃ in the earth’s atmosphere as a pollutant should be regarded. The five compounds are CO₂ and CH₄ which contribute \( ca. 70\% \) to the overall greenhouse effect, N₂O, a chlorofluorocarbon, and SF₅CF₃. Most of the data is taken from the Intergovernmental Panel of Climate Change report of 2001 [35], an associated paper whose data fed into the IPCC2001 report [36], and one of many websites from an enviromental pressure group highlighting statistics on global warming [37]. Whilst some of the data may now be slightly out of date, the longterm trends should be unchanged. The term that is used to characterise the IR absorption properties of a greenhouse gas is the \textit{radiative forcing}, whilst the overall effect of this pollutant on the earth’s climate is described by the \textit{global warming potential}. The former measures the strength of the absorption bands over the infrared black-body region of the earth, \( ca. 400-2000 \ \text{cm}^{-1} \). Essentially, it is a (per molecule) microscopic property, is given the symbol \( a_o \), and is usually expressed in units of W m\(^{-2}\) ppbv\(^{-1}\). If this value is multiplied by the change in concentration of pollutant over a defined time window, usually the 250 years from before the Industrial Revolution to the current day, the macroscopic radiative forcing in units of W m\(^{-2}\) is obtained. One may then compare the radiative forcing of different pollutant molecules over this time window.

The GWP measures the radiative forcing, \( A_x \), of a pulse emission of a greenhouse gas, \( x \), over a defined time period, \( t \), usually 100 years, relative to the time-integrated radiative forcing of a pulse emission of an equal mass of CO₂ [35]:
\[ GWP_x(t) = \frac{\int_0^t A_x(t) \, dt}{\int_0^t A_{CO2}(t) \, dt} \quad (I) \]

The GWP is therefore a dimensionless number that informs how important one molecule of pollutant \( x \) is to the greenhouse effect via global warming compared to one molecule of CO\(_2\). The GWP of CO\(_2\) is defined to be unity. For most greenhouse gases, the radiative forcing following an emission at \( t = 0 \) takes a simple exponential form:

\[ A_x(t) = A_{o,x} \exp(-t / \tau_x) \quad (II) \]

where \( \tau_x \) is the lifetime for removal of species \( x \) from the atmosphere. For CO\(_2\), the lifetime ranges from 50 to 200 years \([37,38]\), a single exponential decay is not appropriate, and we can write:

\[ A_{CO2}(t) = A_{o,CO2} \left[ b_o + \sum_i b_i \exp(-t / \tau_i) \right] \quad (III) \]

where the response function, the bracket in the right-hand side of equ. (III), is derived from more complete carbon cycles. Values for \( b_i \) (\( i = 0-4 \)) and \( \tau_i \) (\( i = 1-4 \)) have been given by Shine et al. \([38]\). It is important to note that the radiative forcing, \( A_o \), in equs. (I)-(III) has units of W m\(^{-2}\) kg\(^{-1}\). For this reason, it is given a different symbol to the microscopic radiative forcing, \( a_o \), which has units of W m\(^{-2}\) ppbv\(^{-1}\).

Conversion between the two units is simple \([38]\). The time integral of the large bracket on the right-hand side of equ. (III), defined \( K_{CO2} \), has units of time, and takes values of 13.44 and 45.73 years for a time period of 20 and 100 years, respectively, the values of \( t \) for which GWP values are most often quoted. Within the approximation that the greenhouse gas \( x \) follows a single-exponential time decay in the atmosphere, it is then possible to parameterise equ. (I) to give an exact analytical expression for the GWP of \( x \) over a time period \( t \):

\[ \frac{GWP_x(t)}{GWP_{CO2}(t)} = \frac{MW_{CO2}}{MW_x} \cdot \frac{a_{o,x}}{a_{o,CO2}} \cdot \frac{\tau_x}{K_{CO2}} \left[ 1 - \exp\left(\frac{-t}{\tau_x}\right) \right] \quad (IV) \]

In this simple form, the GWP only incorporates values for the microscopic radiative forcing of greenhouse gases \( x \) and CO\(_2\), \( a_{o,x} \) and \( a_{o,CO2} \), in units of W m\(^{-2}\) ppbv\(^{-1}\); the molecular weights of \( x \) and CO\(_2\); the lifetime of \( x \) in the atmosphere, \( \tau_x \); the time period over which the effect of the pollutant is determined; and the constant \( K_{CO2} \) which can easily be determined for any value of \( t \). This equation is not given in this specific form in \([38]\), but is easy to derive from equations given therein. Note that a
similar equation given in Section 6 of reference [5] by Mason et al. has numerous typographical errors, and should be disregarded. The recent increase in concentration per unit time interval of a pollutant (e.g. the rise in concentration per annum over the last 5-10 years), one of the factors of most concern to policymakers, does not contribute directly to the GWP value. This and other factors [38] have caused some criticism of the use of GWPs in policy formulation. That said, the important point to note from Table 1 is that CO2 and CH4 contribute most to the greenhouse effect simply due to their high atmospheric concentration; the radiative forcing per molecule and GWP of both gases are relatively low. By contrast, SF\textsubscript{6}CF\textsubscript{3} has the highest radiative forcing per molecule of any greenhouse gas and its GWP value is therefore very high, but its contribution to the overall greenhouse effect is relatively small, because the concentration of SF\textsubscript{6}CF\textsubscript{3} in the atmosphere is still very low.

4. Kinetics and removal processes of SF\textsubscript{6}CF\textsubscript{3} from the earth’s atmosphere

Even if the presence of SF\textsubscript{6}CF\textsubscript{3} in our atmosphere contributes only a very small amount to global warming, nevertheless it is important to understand better the physical and chemical properties of this new greenhouse gas. It is especially important to determine what are the chemical and photolytic reactions that remove it from the atmosphere, since they will contribute to its lifetime and GWP value. The total rate of removal of SF\textsubscript{6}CF\textsubscript{3} per unit volume per unit time is given by

\[
\text{Rate} = \left[ \text{SF}_6\text{CF}_3 \right] \left( k_{1}[OH] + k_{2}[O^* (1D)] + \sum_{\text{ions}} k_{\text{ion}}[\text{ion}] + k_{e}[e^-] + \sum_{\lambda} \sigma_{\lambda} J_{\lambda} \Phi_{\lambda} \right)
\]  

(V)

where each of the five terms in the large bracket of equ. (V) is a pseudo-first-order rate constant. The first four terms represent reactions of SF\textsubscript{6}CF\textsubscript{3} with OH, O*(1D), cations and electrons, respectively ; \(k_1\), \(k_2\), \(k_{\text{ion}}\) and \(k_e\) are the corresponding second-order bimolecular rate coefficients. The first term will dominate in the troposphere (0 < altitude (h) < 10 km), the second term in the stratosphere (10 < h < 50 km), and the third and fourth terms in the mesosphere (h > 50 km). In the fifth term of the bracket, \(\sigma_{\lambda}\) and \(J_{\lambda}\) are the absorption cross section for SF\textsubscript{6}CF\textsubscript{3} and the solar flux at wavelength \(\lambda\), respectively, and \(\Phi_{\lambda}\) is the quantum yield for dissociation following photoabsorption at wavelength \(\lambda\). In the troposphere, the summation for \(\lambda\) is over the range ca. 290-700 nm, in the stratosphere ca. 200-290 nm, and in the mesosphere the solar flux at the Lyman-\(\alpha\) wavelength of 121.6 nm dominates all other vacuum-UV wavelengths. We note that equation (V) assumes that the ion-molecule and electron attachment reactions lead to the removal of every SF\textsubscript{6}CF\textsubscript{3} molecule by formation of dissociation products. Furthermore, secondary reactions of such products must not recycle SF\textsubscript{6}CF\textsubscript{3}. This assumption is true for ion-molecule reactions, but is not necessarily so for electron attachment (Section 6.2).
Having no hydrogen atoms, SF₅CF₃ is very unlikely to be removed by reaction with OH in the troposphere, and the rate coefficient for this reaction is almost certainly too small to measure by conventional laboratory techniques (e.g. discharge flow or pulsed photolysis with laser-induced fluorescence detection of OH). Likewise, the reaction of O*(1D) with SF₅CF₃ has not been measured. There are therefore no chemical processes that remove SF₅CF₃ from the troposphere and stratosphere. Furthermore, the electron energy loss spectrum of SF₅CF₃, in effect a pseudo-absorption spectrum, showed that no bound excited electronic states exist for hv < ca. 8 eV (i.e. λ > 150 nm) [23]. Therefore, σₜ is effectively zero in both the relevant wavelength ranges of the troposphere and stratosphere. These laboratory measurements confirm field measurements reported by Sturges et al. [1] from stratospheric profiling that the lifetime of SF₅CF₃ in the earth’s atmosphere is very long. In Section 5.2, an experiment is described to determine the strength of the SF₅–CF₃ bond, leading to an improved value for the enthalpy of formation of SF₅CF₃ [20,21]. This experiment was performed in the middle of 2000, the authors having obtained an early preprint of the Sturges paper. The principal concern of meterologists then was to determine this bond strength. The result, that the S–C bond has strong σ-character with a dissociation energy of ca. 4 eV or 400 kJ mol⁻¹, confirmed that UV photolysis in the stratosphere was very unlikely to contribute to the rate of removal of SF₅CF₃ from the atmosphere. In retrospect, and given experiments that have been performed since, this is an obvious result. Attention in recent years has therefore turned to laboratory-based measurements relevant to the mesosphere, where ionic processes involving cations, anions and electrons dominate, and where the only significant photolytic wavelength is 121.6 nm. These experiments are described in Section 6. In Section 7, I draw together all these data, and describe how the lifetime of SF₅CF₃, ca. 1000 years, is defined and determined.

5. The VUV photoionisation and photofragmentation spectroscopy of SF₅CF₃

5.1 Definition of the first dissociative ionisation energy: application to CF₄ and SF₆

Photodissociation generally occurs through excitation of a molecule to a repulsive electronic state which lies above a dissociation threshold. Close to threshold, the cross section for photodissociation can be negligibly small, and this makes the experimental determination of a dissociation energy notoriously difficult. Thus, CF₄ has a dissociation energy (to CF₃ + F) of 5.61 eV, but VUV photons with energies in excess of 12 eV are required to photodissociate CF₄. Likewise, the bond dissociation energy of SF₆ (to SF₅ + F) is 3.82 eV, but photodissociation is not observed until the photon energy exceeds ca. 10 eV. SF₅CF₃ seems to show similar properties. The electron energy loss spectrum [23] suggests that there are no excited electronic states of SF₅CF₃ lying less than ca. 8 eV above the ground state. In the stratosphere, the highest-energy solar photons have an energy of ca. 6 e (λ = ca. 200 nm). It is therefore very unlikely that SF₅CF₃ will be photodissociated in the stratosphere, irrespective of the strength of the S–C bond,
simply because dissociative electronic states at energies accessible with stratospheric UV photons are not present. The use of neutral excited states of SF₅CF₃ to determine its dissociation energy to SF₅ + CF₃ is therefore not viable.

CF₄, SF₆ and SF₅CF₃, however, also share the common property that the parent cation is not observed in a conventional 70 eV electron-impact mass spectrum [39]. Thus the ground electronic state of these cations is repulsive in the Franck-Condon region, dissociating on a time scale much faster than the transit time of an ion through a magnetic or quadrupole mass spectrometer. With CF₄ and SF₆, CF₃–F and SF₅–F bond cleavage to form CF₃⁺ or SF₅⁺ + F + e⁻ must occur. With SF₅CF₃, we assume that cleavage of the central S–C bond occurs, forming either SF₅⁺ + CF₃ + e⁻ or CF₃⁺ + SF₅ + e⁻. Since the threshold energy of the latter dissociation channel lies ca. 0.8 eV lower than that of the former, we assume dissociation within the mass spectrometer to CF₃⁺ + SF₅ + e⁻ dominates. (We note that dissociation to SF₄⁺ + CF₄ + e⁻ requires even less energy [20,22], driven by the large and negative enthalpy of formation of CF₄, but the significant re-arrangement of chemical bonds needed in the transition state makes this reaction kinetically unfavourable.) Taking SF₅CF₃ as an example, we can define the first dissociative ionisation energy (DIE) of these molecules to be the 0 K energy of CF₃⁺ + SF₅ + e⁻ relative to the ground vibronic level of the neutral molecule (Figure 2). This energy has alternatively been called the adiabatic ionisation energy of the neutral molecule, but it is this author’s belief that this latter terminology is inappropriate in cases such as CF₄, SF₆ and SF₅CF₃ where the Franck-Condon overlap at threshold is zero. The determination of the DIE of a molecule whose ground state of the parent ion is totally repulsive in the Franck-Condon region is a difficult problem, because its value is likely to be significantly less than the energy corresponding to the onset of ionisation of the neutral precursor; in the opinion of this author, onset of ionisation is a more appropriate phrase to use than adiabatic ionisation energy for such molecules, since the latter term has ambiguous meanings. Thus the photoelectron spectrum of the parent molecule can only give an upper bound to its first DIE.

This problem is well known for CF₄ and SF₆, and for many years the DIE of these molecules has been the subject of controversy. However, Figure 2 can also apply to dissociation of a polyatomic molecule to two molecular free radicals. From the figure, it is clear that:

\[
\text{DIE (ABCD)} = D_o(AB\text{–CD}) + \text{AIE (AB)} \quad (\text{VI})
\]

where ABCD refers to a general polyatomic molecule which dissociates along the central B–C bond, \(D_o(AB\text{–CD})\) is the dissociation energy of the AB–CD bond, and AIE (AB) is the adiabatic ionisation energy of the AB free radical; in this situation, the use of adiabatic ionisation energy is appropriate so long as the ground state of AB⁺ is bound in the Franck-Condon region. For SF₅CF₃, AB will clearly be
the CF₃, and CD the SF₅ radical. For SF₅CF₃, the estimation of its first DIE needs a knowledge of both the 
SF₅–CF₃ bond energy and the AIE of the CF₃ radical. The latter value is now well determined, 9.04 ± 0.04 eV [40], after many years of uncertainty, but when Sturges et al. [1] published their paper the former value was only known within a broad range of ca. 2-4 eV.

One method to determine the DIE of such molecules directly is to turn the problem around, and use the 
fact that, in the Franck-Condon region, the ground state of the parent cation lies above the DIE as a 
benefit rather than a hindrance. The fragments will be formed with translational kinetic energy, and we 
can perform a photoelectron – photoion coincidence (PEPICO) experiment to measure the mean 
translational kinetic energy, <KE>ₜ, released into the AB⁺ + CD fragments; CD is atomic F for 
dissociation of CF₄ or SF₆. Using tunable vacuum-UV radiation, it is most convenient to use threshold 
photoelectron detection, and the acronym for this mode of spectroscopy then becomes TPEPICO. From 
an analysis of the width and shape of the fragment ion (AB⁺) time-of-flight distribution in the TPEPICO 
spectrum measured at a photon energy h ν, it is possible to determine <KE>ₜ. This KE release will 
correspond to some fraction of the available energy, where

\[ E_{\text{avail}} = h \nu + (\text{thermal energy of ABCD}) - \text{DIE(ABCD)} \]  

(VII)

The thermal energy of the parent neutral molecule comprises contributions from rotational energy (3kB𝑇/2 per molecule) and vibrational energy \( (hν_i/\exp(hν_i/k_B𝑇)-1, \text{ summed up for all vibrational modes } ν_i) \). The size of this fraction, <KE>ₜ / Eavail, is governed by the dynamics of the decay mechanism [41]. The 
mechanism cannot unambiguously be determined from a measurement at one single photon energy. By 
measuring <KE>ₜ continuously as a function of (h ν + thermal energy of ABCD), however, and assuming 
that the fractional KE release is independent of energy, an extrapolation to a KE release of zero gives an 
intercept on the x-axis corresponding to the DIE of ABCD (Figure 3). As the AB⁺–CD bond breaks, if 
dissociation involves an impulsive release of energy so great that it results in intramolecular collisions 
between the recoiling fragments (the ‘pure’ impulsive model), then <KE>ₜ and Eavail are related only by 
the kinematics of the dissociation [42]:

\[ <\text{KE}>_\text{t} = \frac{\mu_{B,C}}{\mu_{AB,CD}} E_{\text{avail}} \]  

(VIII)

where \( \mu_{B,C} \) is the reduced mass of the two atoms in the dissociating bond, and \( \mu_{AB,CD} \) is the reduced mass of the two products of the dissociation. Thus the slope of the graph in Figure 3 is given simply by the 
ratio of two reduced masses. As a test of the experiment, we used this method to determine the first DIE
for CF$_4$ and SF$_6$, where there is now some degree of certainty in the correct values. From the former experiment, we determined the first DIE of CF$_4$ to be 14.45 ± 0.20 eV. Hence we were able to determine the 0 K enthalpy of formation of CF$_3^+$ to be 390 ± 20 kJ mol$^{-1}$ and, via $\Delta h^0 (\text{CF}_3)$ [40], the adiabatic ionisation energy of the CF$_3$ free radical. Our value, 8.84 ± 0.20 eV, was in good agreement with the now-accepted best experimental determination of 9.04 ± 0.04 eV [40]. The SF$_6$ experiment determined its first DIE to be 13.6 ± 0.1 eV, leading to a value for $\Delta h^0 (\text{SF}_5^+)$ of 29 ± 10 kJ mol$^{-1}$. Using the recommended value for $\Delta h^0 (\text{SF}_5)$ from the ion beam study of Fisher et al. [43], we obtained a value for the adiabatic ionisation energy of the SF$_5$ free radical of 9.8 ± 0.2 eV. Within error limits, this value is also in good agreement with the guided ion beam result of 9.60 ± 0.05 eV [43]. Following these test experiments, full details of which are given elsewhere [20,21], we measured the first DIE of SF$_5$CF$_3$ (Section 5.3). Using the now well-established ionisation energy of CF$_3$, we were able to determine, admittedly in an indirect manner, the dissociation energy of the SF$_5$–CF$_3$ bond.

5.2 Experimental details

The apparatus used for the TPEPICO study [44] is shown in Figure 4, the source of radiation being beamline 3.1 (1 m Seya-Namioka monochromator) of the UK Daresbury Synchrotron Radiation Source operating at a resolution of 0.3 nm. The monochromatised radiation is coupled into the interaction region via a capillary, and its flux is monitored from the fluorescence of a sodium salicylate coated window. Threshold photoelectrons and fragment cations from the interaction region are extracted in opposite directions by an electric field of 20 V cm$^{-1}$. The threshold electron analyser consists of a cylindrical electrostatic lens followed by a 127° post analyser, which rejects energetic electrons. The lens has a shallow depth of field and poor chromatic aberrations, so that only electrons with low initial energies produced in the centre of the interaction region focus efficiently at the entrance of the post analyser. Simulations suggest a high degree of space focusing, so a finite interaction volume is relatively unimportant. The resolution of the electron analyser, ca. 10 meV, is superior to that of the monochromator in these experiments, therefore the overall resolution of the experiment is limited by that of the photon source. Ions pass through a two-stage acceleration region followed by a linear time-of-flight (TOF) drift tube. This arrangement satisfies the space focusing condition, yielding sufficient TOF resolution that kinetic energy releases from dissociative ionisation events can be measured. Signals from the channeltron and microchannel plate detectors are discriminated and conveyed to a time-to-digital converter (TDC) card via pulse-shaping electronics. The electron signal provides the start pulse, the ion signal the stop pulse, and delayed coincidences can be recorded. Concurrently, the total ion yield and threshold photoelectron spectrum (TPES) can also be measured.
With this apparatus, three different spectra can be recorded. First, the TPES spectrum is obtained by recording the threshold electron signal as a function photon energy. Second, a TPEPICO spectrum is obtained by recording the coincidence spectrum continuously as a function of photon energy. The data record as a 3D map of coincidence counts vs. ion time of flight vs. photon energy. Sections from this map can yield either the time-of-flight mass spectrum at a defined photon energy or the yield of a particular ion. Third, with a fixed photon energy, high resolution TOF spectra can be recorded and values for $<\text{KE}>_T$ obtained \cite{45,46}. Briefly, for each TPEPICO-TOF spectrum a small basis set of peaks, each with a discrete energy release $\varepsilon_t$ is computed, and assigned a probability. The discrete energies are given by $\varepsilon_t(n) = (2n-1)^2 \Delta E$, where $n = 1, 2, 3, 4 \ldots \ 
$. $\Delta E$ depends on the statistical quality of the data; the higher the signal-to-noise ratio, the lower $\Delta E$ and the higher $n$ can be set to obtain the best fit. Each computed peak in the kinetic energy release distribution spans the range $4(n-1)^2 \Delta E$ to $4n^2 \Delta E$, centred at $\varepsilon_t(n) + \Delta E$. The reduced probability of each discrete energy, $P(\varepsilon_t)$, is varied by linear regression to minimise the least-squared errors between the simulated and experimental TOF peak. From the basis set of $\varepsilon_t$ and $P(\varepsilon_t)$, $<\text{KE}>_T$ is easily determined.

5.3 Determination of the first dissociative ionisation energy of SF$_5$CF$_3$ and the S–C bond strength

The three modes of experiment described above were performed for SF$_5$CF$_3$ \cite{20}. In addition, to determine the first DIE, important aspects of the second and third mode were combined. Thus the TOF spectrum for CF$_3^+$ was recorded over a narrow time window with a resolution of 16 ns, close to the optimum of the TDC card of 8 ns, and 64 wavelength channels were scanned over the energy range 12.9-15.7 eV, encompassing the Franck-Condon zone of the ground state of SF$_5$CF$_3^+$. The results are shown in Figure 5. Only those datapoints that define the Franck-Condon region, ca. 13.2-14.8 eV, were used to determine the slope and intercept of the graph. The onset of ionisation for SF$_5$CF$_3$, 12.9 $\pm$ 0.2 eV, lies significantly lower in energy than that in either CF$_4$ or SF$_6$, reflecting the different character of the HOMO. The mean KE releases over these channels range from 0.05 eV to 0.4 eV. As one example, Figure 6 shows the TPEPICO-TOF spectrum of CF$_3^+$/SF$_5$CF$_3$ recorded at 14.09 eV and the best simulated fit, from which a mean KE release of 0.24 $\pm$ 0.05 eV was determined. Within experimental error, the 35 lowest-energy data points which encompass the Franck-Condon region of the ground state of SF$_5$CF$_3^+$ fit to a straight line with a slope of 0.19, in excellent agreement with the prediction of the pure-impulsive model of 0.20. Extrapolation to a KE release of zero on the $y$-axis yields the first DIE of SF$_5$CF$_3$ to CF$_3^+$+$\text{SF}_5$+$\text{e}^-$ to be 12.9 $\pm$ 0.4 eV. The relatively large error in the DIE reflects the small slope of the $<\text{KE}>_T$ vs. photon energy graph, and hence the shallow nature of the extrapolation. We should note that the DIE, unlike that of CF$_4$ and SF$_6$, is coincidentally isoenergetic with the ionisation onset of the first photoelectron band of SF$_5$CF$_3$; for CF$_4$ and SF$_6$, in both cases the first DIE lies over 1 eV lower than the onset of ionisation, leading to a longer extrapolation to zero $<\text{KE}>_T$. 

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Two important thermochemical data can now be determined. First, using values for the 0 K enthalpies of formation of \( \text{CF}_3^+ \) \((409 \pm 3 \text{ kJ mol}^{-1})\) [40] and \( \text{SF}_5 \) \((-915 \pm 18 \text{ kJ mol}^{-1})\) [43], we determine \( \Delta H^0_0 \) (SF\(_5\)CF\(_3\)) to be \(-1750 \pm 47 \text{ kJ mol}^{-1}\), significantly lower than that quoted in the 1998 JANAF tables [29]. Second, using the now-accepted value for AIE (CF\(_3\)) of \(9.04 \pm 0.04 \text{ eV}\) [40], we determine the dissociation energy of the SF\(_5\)–CF\(_3\) bond at 0 K to be \(3.86 \pm 0.45 \text{ eV}\) or \(372 \pm 43 \text{ kJ mol}^{-1}\). (We note that these values are updates of those published in reference [20], following improvements in the published thermochemical values for CF\(_3\) and CF\(_3^+\).) Using the value for AIE (SF\(_5\)) with the highest claim for accuracy from Fisher et al. [43], 9.60 \(\pm\) 0.05 eV, the second DIE of SF\(_5\)CF\(_3\) (defined here to be SF\(_5^+\) + CF\(_3\) + e\(^-\)) is calculated to be \(13.46 \pm 0.45 \text{ eV}\). This energy is ca. 0.6 eV higher than the first DIE to CF\(_3^+\) + SF\(_5\) + e\(^-\), and presumably explains why only CF\(_3^+\) is observed for dissociation of the low-energy regions of the ground-state potential of SF\(_5\)CF\(_3\).

At this stage it is only proper to highlight the assumptions and limitations of this extrapolation technique. The errors quoted for the first DIE of CF\(_4\), SF\(_6\) and SF\(_5\)CF\(_3\) arise from random statistical errors in the data for \(<\text{KE}>_T\). Two factors have been ignored which might produce systematic errors. First, the theory assumes dissociation is 100% impulsive with no statistical component. The simple relation between \(<\text{KE}>_T\) and \(E_{\text{avail}}\) (equ. (VIII)) arises from classical mechanics, with linearity in the extrapolation graph (Figure 3) applying even for very low values of \(<\text{KE}>_T\). This is most likely to be true when the onset of ionisation to ABCD\(^+\) is significantly greater than the DIE of ABCD, i.e. the situation for CF\(_4\) and SF\(_6\), but not perhaps for SF\(_5\)CF\(_3\). Second, anisotropic effects are ignored in the analysis, despite the vacuum-UV radiation source being close to fully plane-polarised perpendicular to the time-of-flight axis. More details of these potential problems are given elsewhere [20].

Theoretical values for \(\Delta H^0_0\) (SF\(_5\)CF\(_3\)) are significantly less negative than the experimental value, yielding weaker SF\(_5\)–CF\(_3\) bond strengths. Ball [30] and Miller et al. [32], both using Gaussian-2 and Gaussian-3 Moller-Plesset perturbation methods, obtain \(\Delta H^0_0\) to be \(-1615\) and \(-1623 \text{ kJ mol}^{-1}\), respectively, the former being the average of G2 and G3 calculations. Using experimental values for the 0 K enthalpy of formation of CF\(_3\) and SF\(_5\) [40,43], they yield S–C bond strengths of 237 and 245 kJ mol\(^{-1}\), respectively. It is worth noting that, whilst there has been convergence in recent years between experiment and theory yielding an accepted value for \(\Delta H^0_0\) (CF\(_3\)), this has not been the case for the SF\(_5\) radical where theory [47] and experiment [43] differ by over 70 kJ mol\(^{-1}\). Using Irikura’s value for \(\Delta H^0_0\) (SF\(_5\)) [47], for example, Ball [30] predicts a S–C bond strength of 311 kJ mol\(^{-1}\), much closer to our experimental value of \(372 \pm 43 \text{ kJ mol}^{-1}\). Xu et al. [33], using varying density functional methods but without zero point vibrational energy corrections, obtain dissociation energies for SF\(_5\)CF\(_3\) \(\rightarrow\) SF\(_5\) + CF\(_3\) ranging between 1.9 and 3.1 eV.
(ca. 180-300 kJ mol\(^{-1}\)). Whilst there are no atmospheric implications for SF\(_5\)CF\(_3\) from these differing values for its enthalpy of formation, it is surprising and slightly concerning that experiment and theory differ by as much as 100 kJ mol\(^{-1}\), ca. 1 eV, for a microscopic property as fundamental as the 0 K enthalpy of formation. State-of-the-art \textit{ab initio} calculations on the SF\(_5\)–CF\(_3\) bond strength are currently being performed with full configuration interaction and a much larger basis set than has been used to date \cite{48}. The calculations are not yet complete, but early indications are that they predict an S–C bond strength much closer to the experimental value \cite{20} than the earlier calculations.

### 5.4 Threshold photoelectron spectrum of SF\(_5\)CF\(_3\)

In the final two parts of section 5, I describe additional and important information on an isolated SF\(_5\)CF\(_3\) molecule that resulted from this TPEPICO study. The valence threshold photoelectron spectrum (TPES) of SF\(_5\)CF\(_3\) was measured from 12.7 to 26.4 eV with a resolution of 0.3 nm (Figure 7). No vibrational structure was observed. The onset of ionisation, defined as the energy at which signal is first observed above the background noise, is 12.92 ± 0.18 eV. The vertical ionisation energy of this first band occurs at 14.13 eV. The low value of this vertical IE, ca. 2 eV lower than that in both CF\(_4\) and SF\(_6\) where the HOMO has F 2p\(\pi\) non-bonding character, has already been noted. The large difference between the onset of ionisation and the vertical IE suggests a significant change in geometry between neutral and cation, probably in the S–C bond length, compatible with a repulsive ground state of the parent cation along this coordinate. \textit{Ab initio} calculations on the structure of SF\(_5\)CF\(_3\) by Knowles \cite{9} at the Hartree-Fock level were described briefly in Section 2. The predicted structure, detailed in the caption to Figure 1, is in good agreement with that from gas-phase electron diffraction \cite{8}. No other structures of molecules with stoichiometry C\(_1\)S\(_1\)F\(_8\) are stable. The HOMO of SF\(_5\)CF\(_3\) has a large S–C \(\sigma\)-bonding character, whereas the next three orbitals lie ca. 2.7 eV lower in energy and are F 2p\(\pi\) non-bonding in character. No minimum-energy geometry of the ground state of SF\(_5\)CF\(_3^+\) can be obtained at either the Hartree-Fock or the MP2(full)/6-31g(d) level, giving further evidence that this state is indeed unbound. Results from these simple calculations have recently been confirmed using the GAMESS-UK suite of programs with a variety of basis sets \cite{15}. Higher-energy peaks in the TPES are observed at 15.68, 16.94, 17.86, 19.44, 21.34, 22.01 and 24.67 eV. Attempts have now been made to assign these peaks in the TPES and how they correlate with peaks in the (T)PES of both CF\(_4\) and SF\(_6\) \cite{15}. The evidence from the vacuum-UV absorption and valence photoelectron spectrum of SF\(_5\)CF\(_3\) is that this molecule behaves more like a perturbed SF\(_6\) than a perturbed CF\(_4\) molecule \cite{15}.

### 5.5 Ion yields and fixed-energy TPEPICO spectra of SF\(_5\)CF\(_3\)

The time-of-flight spectrum of the fragment ions from SF\(_5\)CF\(_3\), integrated over the whole range of VUV photoexcitation energies 12.7–26.4 eV, is shown in Figure 8. The parent ion is not observed. CF\(_3^+\) and SF\(_3^+\) are the dominant ions, then SF\(_5^+\), with CF\(_2^+\) and SF\(_4^+\) being very weak. Table 2 shows threshold
energies for their production. We consider first the major ions, defined as those formed by a single bond cleavage; reactions in which there is no barrier in the exit channel and hence, in the conventional sense, no transition state. The ion yield of CF$_3^+$ follows that of the TPES of SF$_5$CF$_3$ from the onset of ionisation to ca. 20 eV, and states of the parent ion with vertical energies below 20 eV dissociate predominantly to this ion. By contrast, the SF$_5^+$ signal is so weak that it is not possible to say whether its ion yield shows any correlation with electronic states of SF$_5$CF$_3^-$. Even well above the SF$_5^+$ threshold of 13.9 eV, its ion yield relative to that of CF$_3^+$ remains very weak. These thresholds and relative intensities are in good agreement with a tunable electron energy beam study over the same range of energies [22]. This behaviour is somewhat surprising for a large polyatomic cation with ten atoms and a correspondingly large density of vibronic states. These dissociation properties of SF$_5$CF$_3^+$ suggest that it behaves more like a pseudo-diatomic molecule with a much lower density of states, dissociating only on one repulsive potential energy surface with no curve crossings. By contrast, the minor ions are defined as those which can only form following intramolecular rearrangement within a transition state of the unimolecular reaction following photoexcitation. The threshold energy for the strongest minor ion, SF$_3^+$, is 14.94 ± 0.13 eV. Energetically, it can only form at this energy if the accompanying neutral fragments are CF$_4$ + F. Likewise, the two very weak minor ions, SF$_4^+$ and CF$_2^+$, can only form at their thresholds (Table 2) if migration of F$^-$ across the S–C bond occurs in the transition state to form CF$_4$ and SF$_6$, respectively. Such migrations in unimolecular reactions are usually only associated with H(D) atoms, but F-atom migration across a C–X bond has been observed in fragmentation of other long-lived greenhouse gases, e.g., the perfluorocarbon cations, C$_x$F$_y^+$ [49].

Table 3 shows values for the mean translational KE release into CF$_3^+$ + SF$_5$ when SF$_5$CF$_3$ is excited into the ground and first four excited states of the parent ion. The procedure is described in Section 5.2. Columns 5-7 show the experimental fractional release into translational energy ($f_{\text{expt}} = <\text{KE}>_T/E_{\text{avail}}$) and the predictions of statistical [50] and pure-impulsive [42] models. It was noted in Section 5.3 that for dissociation of the ground state of SF$_5$CF$_3^+$, the value for $f_{\text{expt}}$ at 14.25 eV, 0.21, is very close to that predicted by the pure-impulsive model. This is as expected, because the Franck-Condon maximum of the ground state of SF$_5$CF$_3^+$ lies ca. 1.3 eV higher than the dissociation threshold to CF$_3^+$ + SF$_5$ + e$^-$. Dissociation from this repulsive potential energy surface is therefore expected to occur rapidly on a sub-picosecond timescale, with a relatively large amount of the available energy being released into translation of the two fragments. As the photon energy increases to 19 eV, however, $f_{\text{expt}}$ decreases by a factor of ca. 3-4. It appears that as higher-energy electronic states of SF$_5$CF$_3^+$ are populated, there is a reduced coupling of the initially-excited vibrational modes into the reaction coordinate, behaviour that is normally associated with a multi-atom polyatomic species. This phenomenon, that the value of $<\text{KE}>_T$ does not increase as rapidly with photon energy as a pure-impulsive model would predict, has also been observed in dissociation reactions in other polyatomic molecules, CF$_3^+$ from CF$_4$ and SF$_5^+$ from SF$_6$ [51].
6. Removal processes of SF₅CF₃ from the earth’s atmosphere in the mesosphere

The larger-than-expected value of the S–C bond strength [20] and the absence of excited electronic states of SF₅CF₃ lying below ca. 8 eV [23] led scientists to study processes in the mesosphere that could remove this greenhouse gas from the atmosphere; reactions with cations, electrons and vacuum-UV photodissociation (the last three terms in the bracket of equ. (V)).

6.1 Reactions of SF₅CF₃ with cations and anions

Two papers have described the reactions of cations with SF₅CF₃ [52,53], one with anions [54]. All these studies have been made in a selected ion flow tube (SIFT), and a schematic of this apparatus is shown in Figure 9. Such an experiment can determine the rate coefficient and product ions and their branching ratios for the reactions of small anions or cations with neutral molecules, generically A± + BC → D± + EF where the ions can be atomic or molecular. Only fast reactions with rate coefficient greater than ca. 10⁻¹³ cm³ molecule⁻¹ s⁻¹ can be studied. In brief, a reagent ion of interest is produced in a high pressure electron impact ion source containing an appropriate source gas. The cation or anion is injected through a quadrupole mass filter into a flow tube holding ca. 0.5 Torr of high purity helium as a buffer gas. The neutral reactant of choice is then admitted through an inlet at one of various points down the flow tube. The resultant ionic products are detected using a quadrupole mass spectrometer. The loss of reagent ion signal, alongside the increase in the various product ion signal(s), is recorded as a function of neutral reactant concentration. The amount of neutral is altered between zero and a concentration that depletes the reactant ion signal by ca. 90 %. Since the experiment operates under pseudo-first-order conditions with [A±] « [BC], and with knowledge of the reaction length and ion flow velocity, a plot of the logarithm of the reagent ion signal vs. neutral molecule concentration allows the rate coefficient to be determined. Percentage branching ratios for each product ion are derived from graphs of the relative product ion counts vs. neutral molecule concentration, with extrapolation to zero neutral gas flow to allow for the effect of any secondary reactions. Most of the Birmingham, UK studies are made at 298 K, whereas the Air Force Research Laboratories, US apparatus can operate over the temperature range ca. 250-500 K.

Unrelated to the atmospheric importance of such reactions for SF₅CF₃, an additional motivation for such studies is to understand the importance of long-range charge transfer in cation-molecule reactions. Charge transfer can occur either at long range or at short range. In the long-range mechanism, assuming BC has a permanent dipole moment, A⁺ and BC approach under the influence of their charge-dipole interaction, until at some critical distance (Rc) the A⁺-BC and A-BC⁺ potential energy curves cross. At this point an electron jump can take place. We have shown [55] that Rc depends on the difference in energy between the recombination energy (RE) of A⁺ and the ionisation energy (IE) of BC; the smaller
this difference, the larger $R_c$. (The RE is defined as the energy that is released when a cation forms its neutral compound.) Furthermore, two important factors for a rapid electron transfer and an efficient long-range charge transfer process are a non-zero energy resonance connecting BC to an electronic state of BC$^+$ at the RE of A$^+$, and the transferring electron comes from a molecular orbital of BC that is not shielded from the approaching cation. So long as there is some overlap of vibrational wavefunction between BC and BC$^+$ at the RE of A$^+$, the magnitude of the photoionisation Franck-Condon factor for BC is not as important as originally thought in determining the efficiency of such a reaction. We note that if this long-range charge transfer mechanism operates, then the branching ratios for fragmentation of (BC$^+$)$^\ast$, where $\ast$ donates the possibility of BC$^+$ being in an excited electronic state, are expected to be independent of how this state is produced. Hence, we would expect similar product branching ratios from the ion-molecule study and from the TPEPICO photoionisation study, assuming the photon energy in the latter experiment matches the RE of A$^+$ in the former.

When long-range charge transfer is unfavourable, A$^+$ and BC move closer together. As their separation decreases, distortion of the potential energy surface of interaction occurs. Eventually, a curve crossing can occur through which efficient charge transfer takes place. This is called short-range charge transfer. As an intermediate complex has formed, a chemical reaction, defined as the breaking and making of new bonds, may, in addition, compete with short-range charge transfer. This means that it is unlikely that the product branching ratios from the ion-molecule and from the TPEPICO experiments will mimic each other. Thus, a comparison of the fragmentation patterns from the SIFT and TPEPICO experiments, together with an analysis of the TPES of BC at the energy of the RE of A$^+$, may indicate which mechanism, be it long-range or short-range, is dominant for the reaction of each cation. Agreement within ca. $\pm 15\%$ is taken as evidence for possible long-range charge transfer [56]. We have made this comparison for the reactions of SF$_5$CF$_3$ with those cations whose concentrations are the highest in the mesosphere, and might therefore be expected to contribute to the removal of SF$_5$CF$_3$ from the earth’s atmosphere.

The results of the SIFT study for reactions of SF$_5$CF$_3$ with the important atmospheric cations N$_2^+$, N$^+$, CO$^+$, CO$_2^+$, O$^+$, N$_2$O$^+$, H$_2$O$^+$ and O$_2^+$ are shown in Table 4. The RE values of these cations range from 15.58 (for N$_2^+$) to 12.07 eV (for O$_2^+$). The first five ions have RE values greater than the onset of ionisation of SF$_5$CF$_3$, so in principle long-range charge transfer may occur. The last three cations have RE values below the onset of ionisation, so can only react via a short-range process and the formation of a reaction intermediate. Seven of the eight ions react with a rate coefficient close to or equal to the Langevin capture value [57]. O$_2^+$ reacts significantly slower. The reactions of the six cations with RE in the range 12.89 to 15.58 eV (i.e. those equal to or greater than the onset of ionisation of SF$_5$CF$_3$) appear to occur by dissociative charge transfer, with CF$_3^+$ and SF$_3^+$ being the dominant product ions. It seems
likely that CF$_3^+$ forms by direct breaking of the S–C bond, whereas SF$_3^+$ may form from dissociation of (SF$_4^+$)$^*$ following intramolecular rearrangement within a reaction intermediate. The evidence for formation of such an intermediate lies in the SF$_3^+$ branching ratios. Significant values, ca. 10-20 %, are observed for reactions of N$^+$, CO$^+$, CO$_2^+$, O$^+$ and N$_2$O$^+$ despite the fact that these ions have RE values less than 14.94 eV, the appearance energy at 298 K ($AE_{298}$) of this cation in the photon-induced study (Section 5.5) [20]. Only the yield of SF$_3^+$ from the N$_2^+$ reaction, 28 %, gets close to approaching that from the photon-induced reaction at h$\nu$ = 15.58 eV, 25 %. Note also that the SF$_3^+$ ion is dominant for the H$_2$O$^+$ reaction, 92 %, although this ion has an RE value 0.3 eV below the onset of ionisation of SF$_5$CF$_3$. Thus the evidence is that, with the possible exception of N$_2^+$, all reactions yielding SF$_3^+$ proceed via the formation of an intermediate at short range, followed by intramolecular re-arrangement and migration of F$^-$ across the S–C bond to form CF$_4$ as an accompanying, thermochemically stable neutral partner. Table 4 also highlights the difficulties in using the experimental values of Fisher et al. [43] for the enthalpies of formation of SF$_x^+$, in that some ‘fast’ reactions are calculated to be significantly endothermic. Thus, column 5 of Table 4 also shows values for $\Delta_rH^o_0$ calculated using the theoretical values of Irikura [47] for $\Delta_fH^o_0 (SF_x^+)$, where his more negative values make such reactions exothermic.

In theory, the rate coefficients and products of such reactions should be used to model the fate of SF$_5$CF$_3$ in the mesosphere and hence its atmospheric lifetime. However, despite the fact that these reactions are fast, ca. 10$^{-9}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, they are still over an order of magnitude slower than low-energy electron attachment (Section 6.2). Furthermore, the concentrations of such positive ions in the mesosphere are also at least an order of magnitude lower than that of free electrons [58]. Therefore, the reactions of both cations and anions with SF$_5$CF$_3$, whilst yielding a huge amount of information at the microscopic level, makes very little contribution to the atmospheric chemistry of this molecule, because the reactions are too slow and the ion concentrations too low; the third term in the bracket in the right-hand side of equ. (V) is negligibly small. Although this section has concentrated on the reactions of cations with SF$_5$CF$_3$, mainly because of the comparison with the TPEPICO data and hence an understanding of reaction mechanisms, the arguments in this paragraph also hold true for the reactions of anions with SF$_5$CF$_3$.

### 6.2 Reactions of SF$_5$CF$_3$ with low-energy electrons

Since the mesosphere is electrically neutral, the total concentration of positive ions must be balanced by that of anions and free electrons, and Adams and Smith [58] estimate a mean electron number density of 10$^5$ cm$^3$ for altitudes in the range 100 < $h$ < 200 km, falling rapidly to 10$^3$ cm$^3$ for $h$ ≈ 80 km. The mesosphere is thermalised with a temperature in the range 170 to 250 K, the lowest temperature occurring at the mesopause, $h$ ≈ 85 km. The average thermal energy of the electrons, 3$k_B T/2$, is therefore low, ca.
0.02–0.03 eV, and laboratory studies of low-energy electron attachment to SF$_5$CF$_3$ can give an important
guide to reactions that may occur in the mesosphere.

Several groups have made such measurements. The group of Mayhew in Birmingham, UK [59] uses a
Swarm apparatus to determine absolute electron attachment cross-sections, rate coefficients and the
products of the low-energy attachment. The Air Force Research Laboratory group of Viggiano [32] use a
flowing afterglow Langmuir probe (FALP) apparatus whose temperature can be controlled between ca.
300-550 K. They can also measure the rate coefficient for low-energy electron attachment and the
products of such reactions. Both give near-identical results for SF$_5$CF$_3$. Mayhew and Kennedy [59]
measure the thermal attachment rate coefficient to be $7.7 \pm 0.6 \times 10^{-8}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, with the only
anionic product being SF$_5^-$ . Miller et al. [32] measure the rate coefficient at 296 K to be slightly higher,
$8.6 \pm 2.2 \times 10^{-8}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, the rate constant increasing very slightly with increasing temperature
giving an Arrhenius activation barrier of 25 meV or 2.4 kJ mol$^{-1}$. Again, the only observed charged
product is SF$_5^-$ . The groups of Illenberger (Berlin, Germany) [25] and Märk (Innsbruck, Austria) [26]
operate under collision-free beam conditions and use a tunable electron beam with energies in the range
ca. 0-15 eV. Absolute cross-sections and rate coefficients cannot be determined, but the presence of low-
energy resonances leading to the production of certain anions can be measured. For SF$_5$CF$_3$, both beam
studies show a large dissociative electron attachment cross section at 0 eV to form SF$_5^-$ + CF$_3$, in
excellent agreement with the Swarm and FALP studies. The 0 eV CF$_3^-$ peak is over four orders of
magnitude weaker [25]. Of less importance to the atmospheric chemistry of SF$_5$CF$_3$, absolute cross
sections for electron scattering at much higher electron energies, 100-10,000 eV, have been measured [24].

A schematic of the Birmingham Swarm apparatus, described in detail elsewhere [60], is shown in Figure
10. The source of electrons is a radioactive $^{63}$Ni $\beta$-emitting sample which is located in front of a drift
tube. An electric potential can be applied along this drift tube, creating a uniform electric field, $E$, which
draws electrons towards a Faraday plate detector. An electron energy distribution is established,
determined by a dynamic balance between the kinetic energy gained from the electric field and energy
loss through multiple collisions with the high-pressure buffer gas (either Ar or N$_2$, number density $N$).
Anions formed by electron attachment can be distinguished from electrons by a simple timing
arrangement in which the electrons are admitted into the drift tube as a short (ca. 1 ms) pulse via an
electron gate. By monitoring the attenuation of electron pulses as a function of the reactant gas
concentration, the density-reduced electron attachment coefficient, $\alpha$, is determined as a function of $E/N$.
Multiplication of $\alpha$ by the mean electron drift speed gives the electron attachment rate constant, $k_a$, again
as a function of $E/N$. A small hole in the Faraday plate allows anions formed within the drift tube to enter
a differentially-pumped region and a quadrupole mass spectrometer. The product anions formed from
electron attachment can thus be determined, the major advantage of this instrument compared to more conventional electron swarm apparatus [61].

Using Ar or N₂ buffer gas, electrons in the Swarm apparatus have a non-thermal energy distribution. However, many detectors commercially used for the detection of electronegative molecules rely on thermal electron attachment in high-pressure buffer gas. Furthermore, if the concentration of an electron-attracting gas is to be accurately determined, thermal attachment rate coefficients are needed. Until recently, we could only obtain such rate coefficients, such as for SF₅CF₃ [59], by extrapolation of our data to zero electric field strengths (Figure 11). We have now developed a new drift tube ideally suited for thermal electron attachment in a high pressure (ca. 1 bar) of CO₂ buffer gas, and confirmed its correct mode of operation with SF₆ where α is now determined to be independent of E/N [62].

The electron attachment rate coefficient for SF₅CF₃, 7.7 ± 0.6 x 10⁻⁸ cm³ molecule⁻¹ s⁻¹, is fast, but significantly slower than that for SF₆, 2.38 ± 0.15 x 10⁻⁷ cm³ molecule⁻¹ s⁻¹ [62], which is close to the theoretical maximum for pure s-wave capture [63]. From an atmospheric point of view, however, perhaps the most important result is that the product of such low-energy electron attachment to SF₅CF₃ is not the parent anion, but a fragment anion SF₅⁻. Assuming SF₅CF₃ is not recycled by subsequent reactions of SF₅⁻, low-energy electron attachment is therefore one route to remove SF₅CF₃ from the earth’s atmosphere, and the pseudo-first-order rate constant for this process, the fourth term in the bracket on the right-hand side of equ. (V), will be significant. Energetically, this removal process is also possible. Table 5 shows that, within the error for ∆fH₀(SF₅CF₃) quoted in Section 5.3, the reaction SF₅CF₃ + e⁻ (0 eV) → SF₅⁻ + CF₃ is exothermic, using both the 0 K enthalpy of formation for SF₅CF₃ from experiment [20] and the less negative value from theory [32]. Using either value for ∆fH₀(SF₅CF₃), formation of CF₃⁻ + SF₅ is significantly endothermic, explaining the almost total absence of this anion in the zero eV peak in the electron beam studies [25,26].

### 6.3 Reactions of SF₅CF₃ with vacuum-UV photons, especially Lyman-α (121.6 nm) radiation

The VUV absorption spectrum of SF₅CF₃, including a determination of the absorption cross section at the Lyman-α wavelength of 121.6 nm, has been measured by five groups using either direct absorption and the Beer-Lambert law [4,16,17] or a double-ion chamber [15,18]. Measurements have been made using VUV laser radiation [17], fixed-energy lamp sources [18], and tunable VUV radiation from a synchrotron [4,15-17]. The synchrotron experiments have been performed with wavelengths both above [4,17] and below [15,16] the lithium fluoride window cutoff of ca. 105 nm (or 11.8 eV). The apparatus used by the Birmingham group, described in detail by Hoxha et al. [64] and used at both the Bessy-I, Berlin and SuperAco, Paris synchrotron sources, is shown in Figure 12.
In brief, radiation from the synchrotron passes through the exit slit of a VUV monochromator (a 1.5 m normal-incidence at Bessy-I), a two-stage differential pumping section, and a 1 mm thick stainless steel microchannel plate into an absorption cell of length, \(L\), 300 mm. A pressure differential of 1000:1 across the microchannel plate is possible. This plate can transmit wavelengths well below 100 nm, so this experiment can measure absorption spectra below the LiF cutoff, and yet the path length of the absorption ‘cell’ is defined. The gas pressure in the absorption cell, in the range 5-60 \(\mu\)bar, is maintained constant via a slow controlled flow of gas. The VUV radiation at the end of the cell is detected through a sodium-salicylate-coated window and a visible photomultiplier tube. Since the pressure of gas and optical path length are known, measurement of the ratio of transmitted intensity observed for background (no gas) and sample spectra (with gas) can yield, via the Beer Lambert law \(\ln(I_o/I) = \sigma c L\), absolute absorption cross sections, \(\sigma\), in units of \(\text{cm}^2\text{ molecule}^{-1}\); \(c\) is the number density of the gas in units of molecules \(\text{cm}^{-3}\), and \(L\) has units of cm. In the calculation of \(I_o/I\) at every value of the VUV energy, allowance is made for the natural decay of the VUV flux over the time of an experiment. No allowance is made for the small pressure gradient within the absorption cell due to gas leakage through the microchannel plate, and the small effects of second-order radiation from the VUV monochromator are ignored. We estimate that cross sections are accurate to \(ca.\) 15-20 \%, and the ignorance of second-order effects may make this error greater at wavelengths close to twice the blaze wavelength of the grating in the VUV monochromator.

Absorption spectra in the range \(ca.\) 50-200 nm at a resolution of better than 0.1 nm can routinely be measured. The range of cross sections that can be determined span \(ca.\) \(10^{-16}\) to \(10^{-20}\) \(\text{cm}^2\). The absorption spectrum of SF\(_5\)CF\(_3\) measured with this apparatus is shown in Figure 13 [16], and the cross section at 121.6 nm is determined to be \(1.3 \pm 0.2 \times 10^{-17}\) \(\text{cm}^2\) (or \(13 \pm 2\) Mb); note that, following a re-analysis of the data, this value is 2 Mb lower than that reported in [16]. This value is significantly greater than that for SF\(_6\) where \(\sigma_{121.6}\) is \(1.76 \times 10^{-18}\) \(\text{cm}^2\), and orders of magnitude greater than for CF\(_4\) which shows negligible absorption at this wavelength (\(\sigma_{121.6} < 8 \times 10^{-22}\) \(\text{cm}^2\)) [65]. As commented in Section 5.4, the evidence from the assignment of the absorption spectrum by Holland et al. [15] is that SF\(_5\)CF\(_3\) behaves more like SF\(_6\) than CF\(_4\) in its VUV photoabsorption properties. The value of \(\sigma_{121.6}\) from Chim et al. [16] is slightly higher than that determined by others. Holland et al. [15], Takahashi et al. [17] and Limao-Vieira [4] obtain 6.4, 7.8 and 6.5 Mb, respectively, and the result of Chim et al. may be anomalously high because higher-order effects are ignored. That said, using the known value of \(J_{121.6}\) [66], the absolute value of this cross section makes negligible difference to the lifetime of SF\(_5\)CF\(_3\) in the mesosphere, because VUV photodissociation only contributes \(ca.\) 1 \% to the removal of SF\(_5\)CF\(_3\) by processes that occur in the mesosphere (Section 7).
As with the states of SF₅CF₃− excited by low-energy electrons, VUV photoabsorption will only contribute to the removal of SF₅CF₃ from the atmosphere if the valence or Rydberg electronic states photoexcited at 121.6 nm are dissociative. The bottom half of Table 5 shows that whilst dissociation to neutral SF₅ and CF₃ is clearly energetically possible since the SF₅−CF₃ bond strength is only 3.86 eV (Section 5.3), dissociation to the ion pair CF₃⁺ + SF₅− (but not to SF₅⁺ + CF₃−) can also occur with 10.2 eV photon excitation. In determining the energetics of these reactions, we use values for the electron affinity of the CF₃ and SF₅ radicals of 1.82 and 3.8 eV, respectively [67,68]. Although the quantum yields for dissociation into ion pairs are generally unknown, their formation following VUV photoexcitation of polyatomic molecules should be prevalent, but this process for SF₅CF₃ has not yet been observed.

Table 6 summarises the electron attachment rate coefficient and Lyman-α absorption cross section for SF₅CF₃, SF₆ and CF₄; reactions of these molecules with small cations and anions are not included as these processes make negligible contribution to their removal from the mesosphere. It is clear that SF₅CF₃ behaves much more like a modified SF₆ than a modified CF₄ molecule. The electron attachment rate coefficient for SF₅CF₃ is ca. 3 times slower than that for SF₆, σ₁₂₁.₆ is ca. 7 times larger, but ke and σ₁₂₁.₆ for CF₄ are orders of magnitude smaller. The only aspect of the photochemistry of SF₅CF₃ where it behaves like a modified CF₄ molecule is its vacuum-UV fluorescence properties. Following VUV photoexcitation in the range 8−25 eV, weak UV/visible emission is observed from CF₃, CF₂ and possibly the parent ion in the energy ranges ca. 10-12, 15-17 and hν > 20 eV [70]. In this aspect, SF₅CF₃ appears to behave like the group of molecules CF₃X (X = H,F,Cl,Br) [71]. This work does not relate to the atmospheric chemistry of SF₅CF₃, but will be described in detail in a future publication [70].

7. Lifetime of SF₅CF₃ in the earth’s atmosphere and conclusions

The lifetime of a pollutant or greenhouse gas is a term that can mean different things to different people, and confusion can sometimes arise [72]. To a laboratory-based physical chemist, the lifetime usually means the inverse of the pseudo-first-order rate constant of the dominant chemical or photolytic process that removes the pollutant from the atmosphere. Using CH₄ as an example, it is predominantly removed in the troposphere via oxidation by the OH free radical, OH + CH₄ → H₂O + CH₃. The rate coefficient for this reaction at 298 K is 6.39 x 10⁻¹⁵ cm³ molecule⁻¹ s⁻¹ [73], so the lifetime is approximately equal to (k₂₉₈[OH])⁻¹. Using an average value for the tropospheric OH concentration of 10⁶ molecules cm⁻³, the atmospheric lifetime of CH₄ is calculated to be ca. 5 years. This is within a factor of 2.4 of the accepted value of 12 years (Table 1). The difference arises because CH₄ is not emitted uniformly from the earth’s surface, a finite time is needed to transport CH₄ via convection and diffusion into the troposphere, and oxidation occurs at different altitudes in the troposphere where the concentration of OH will vary from its average value of 10⁶ molecules cm⁻³. In simple terms, this is an example of a two-step kinetic process, A
B → C with first-order rate constants $k_1$ and $k_2$. The first step, A → B, defines the transport of the pollutant into the atmosphere, whilst the second step, B → C, defines the chemical or photolytic process (e.g. reaction with an OH radical in the troposphere, electron attachment in the mesosphere) that removes the pollutant from the atmosphere. In general, the overall rate of the process (whose inverse is called the *lifetime*) will be a function of both $k_1$ and $k_2$, but its value will be dominated by the *slower* of the two steps. Thus, in writing the lifetime of CH$_4$ simply as $(k_{298}[OH])^{-1}$, we are assuming that the first step, transport into the region of the atmosphere where chemical reactions occurs, is infinitely fast.

SF$_5$CF$_3$, perfluorocarbons such as CF$_4$, and SF$_6$ behave in the opposite sense. The slow, rate-determining process is now the first step, transport of the greenhouse gas from the surface of the earth into the mesosphere, and the chemical or photolytic processes that remove SF$_5$CF$_3$ *etc.* in the mesosphere will have very little influence on the lifetime. We can define a chemical lifetime, $\tau_{\text{chemical}}$, as $(k_e[e^-] + \sigma_{121.6}J_{121.6})^{-1}$, assuming the quantum yield for dissociation at 121.6 nm is unity, but the value will be a function of position, particularly altitude, in the atmosphere. In the troposphere, $\tau_{\text{chemical}}$ will be infinite because both the concentration of electrons and $J_{121.6}$ are effectively zero at low altitude, but in the mesosphere $\tau_{\text{chemical}}$ will be much less. In other words, multiplication of $k_e$ for SF$_5$CF$_3$ *etc.* by a typical electron density in the mesosphere, ca. $10^4$ cm$^{-3}$ [58], yields a chemical lifetime which is far too small and bears no relation to the true atmospheric lifetime, simply because most of the SF$_5$CF$_3$ *etc.* does not reside in the mesosphere.

The global atmospheric lifetime is obtained from globally-averaged loss frequencies. In forming the average, the psuedo-first-order destruction rate coefficient for each region of the atmosphere is weighted according to the number of molecules of compound in that region,

$$<k>_{\text{global}} = \frac{\sum_i k_i V_i n_i}{\sum_i V_i n_i}$$  \hspace{1cm} \text{(IX)}$$

where $i$ is a region, $k_i$ is the pseudo-first-order rate coefficient for region $i$, $V_i$ is the volume of region $i$, and $n_i$ is the number density of the greenhouse gas under study in region $i$. The averaging process thus needs input from a 2-D or 3-D model of the atmosphere in order to supply the values of $n_i$. This is essentially a meterological problem, and may explain why such scientists and physical chemists sometimes have different interpretations of what the lifetime of a greenhouse gas actually means. The only sensible definition is the inverse of $<k>_{\text{global}}$. Many such studies have been made for SF$_6$ [65,74,75], and differences in the kinetic model ($k_i$) and the atmospheric distributions ($n_i$) from different climate or
transport models account for the variety of atmospheric lifetimes that have been reported. For molecules such as SF₆ (and SF₅CF₃) which are only destroyed in the mesosphere above 60 km, the importance of both these factors has been explored by Hall and Waugh [76]. Their results show that because the fraction of the total number of SF₆ molecules in the mesosphere is very small, the global atmospheric lifetime is very much longer than the mesospheric, chemical lifetime. Thus, they quote that if the mesospheric loss frequency is 9 x 10⁻⁸ s⁻¹, corresponding to a local lifetime of 129 days, then the global lifetime ranges between 1424 and 1975 years, according to which climate or transport model is used. In conclusion, therefore, the lifetime of SF₆, SF₅CF₃ and CF₄ is dominated by the meteorology that transports these pollutants into the mesosphere, but the fate of each molecule when it reacts with low-energy electrons and Lyman-α radiation may make a small contribution to <k> global.

To the author’s knowledge, no full meteorological analysis of SF₅CF₃ in the atmosphere has been performed. SF₆ and SF₅CF₃ both attach electrons with a fast rate coefficient (> 10⁻⁸ cm³ molecule⁻¹ s⁻¹) and show significant absorption with Lyman-α radiation. Furthermore, the height profiles of SF₆ and SF₅CF₃ are similar in the atmosphere [1]. It should therefore be appropriate to make quantitative comparisons between these two molecules, expecting them to have similar lifetimes in the atmosphere. By contrast, the almost infinite value for the lifetime of CF₄, greater than 50,000 years [65,77], arises because this molecule shows no absorption at 121.6 nm [65] and attaches electrons at a negligibly slow rate [69]. kₑ for SF₅CF₃ at 298 K is 3.1 times smaller than for SF₆, whereas the absorption cross-section is 7.4 times larger (Table 6). Therefore, the pseudo-first-order rate constant for removal of SF₅CF₃ by electrons divided by that due to photons at the same altitude of the mesosphere, kₑ[e⁻] / σ₁₂₁.₆J₁₂₁.₆, is 22.9 times smaller than this ratio for SF₆. (This calculation assumes that electron attachment to both SF₆ and SF₅CF₃ leads only to the formation of a fragment anion, SF₅⁻, which is not recycled.) Assuming an incorrect value for kₑ of 1 x 10⁻⁹ cm³ molecule⁻¹ s⁻¹, Ravishankara et al. [65] initially determined that this ratio of pseudo-first-order rate constants for SF₆ was 3.2, and obtained an atmospheric lifetime of 3200 years. Using the much larger and correct value for kₑ (SF₆) of 2.3 x 10⁻⁷ cm³ molecule⁻¹ s⁻¹, Morris et al. [74] showed that this ratio of pseudo-first-order rate constants was 1998, and determined a lower limit to the atmospheric lifetime of 800 years. The lower limit arises because they assumed that every electron attachment event led to the permanent removal of SF₆ from the atmosphere. Reddmann et al. [75] considered a number of scenarios for the destruction of SF₆⁻. They found that if less than 100 % was destroyed, the lifetime, not surprisingly, increased. They calculated values spanning 400 to 10000 years, depending on the loss mechanism and the value for the electron density in the upper stratosphere / mesosphere.

Assuming that the analysis of Morris et al. [74] for SF₆ is correct, then the predominant removal process for SF₅CF₃ remains electron attachment and not VUV photolysis, since the ratio of the pseudo-first-order
rate constants for SF$_5$CF$_3$ is 1998 / 22.9 or 87.2, i.e. still much greater than unity. Without a full meterological study, however, as explained above it is very difficult to convert these first-order rate constants into a global lifetime for SF$_5$CF$_3$. What can be said is that, per molecule, the quantum yield for electron attachment to SF$_5$CF$_3$ in the mesosphere is 87.2/88.2 or 0.989, and for Lyman-α photodissociation 1/88.2 or 0.011. The atmospheric lifetime is dominated by the meterology that transports SF$_5$CF$_3$ to the upper altitudes of the atmosphere and is likely to be ca. 1000 years, with an uncertainty in this number possibly as large as ± 300 years.

In conclusion, it is quite clear that, whilst the current concentrations of SF$_5$CF$_3$ in the atmosphere are still very low, mankind is stuck with this problem for a very long time. Since the original paper by Sturges et al. was published five years ago [1], a huge amount of fundamental research has meant that the atmospheric properties and reactions of SF$_5$CF$_3$ are now well understood, and further research effort will only contribute at the edges to understand the environmental impact of this greenhouse gas. It is suggested that reliable monitors, probably using the IR absorption properties of this molecule, are installed to determine the concentration of SF$_5$CF$_3$ both at the earth’s surface and in the atmosphere. It is also suggested that policymakers should be concerned at the high annual growth rate of SF$_5$CF$_3$, and measures should be put in place world-wide to control its emissions.

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Table Captions

Table 1  Examples of greenhouse gases, and their contribution to global warming [35-37].

Table 2  Appearance energies of fragment ions from SF₅CF₃ over the photoexcitation energy range 12.7-26.4 eV.

Table 3  Total mean translational kinetic energy release, <KE>ₜ, for the two-body fragmentation of the valence states of SF₅CF₃⁺.

Table 4  Rate coefficients, cation product ions and their branching ratios for the reactions of cations with possible atmospheric importance with SF₅CF₃. The recombination energies of the reagent cations are listed in eV in brackets under the cations. The capture rate coefficients, calculated using Langevin theory [57], are presented in square brackets under the experimental values. 0 K enthalpies of reaction for these channels are given in column 5. For formation of SF₃⁺ and SF₄⁺, two values for Δ_rH° are given. The first value is calculated using the data set of Fisher et al. [43], whereas the second corresponds to that of Irikura [47]. The two values given for Δ_rH° for formation of SF₅⁺ by dissociative charge transfer result from taking the IE (SF₅) to be either 9.60 eV [43] or 9.71 eV [47].

Table 5  Possible products from e⁻ attachment and Lyman-α excitation of SF₅CF₃.

Table 6  Thermal electron attachment rate constants, absorption cross-sections at 121.6 nm, and atmospheric lifetimes for CF₄, SF₆ and SF₅CF₃.
Table 1. Examples of greenhouse gases, and their contribution to global warming [35-37].

<table>
<thead>
<tr>
<th>Greenhouse Gas</th>
<th>CO₂</th>
<th>CH₄</th>
<th>N₂O</th>
<th>CF₂Cl₂</th>
<th>SF₅CF₃</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentration in 1998 / ppmv</td>
<td>365</td>
<td>1.75</td>
<td>0.31</td>
<td>0.0005</td>
<td>1.2 x 10⁻⁷</td>
</tr>
<tr>
<td>ΔConcentration (1750–1998) / ppmv</td>
<td>87</td>
<td>1.045</td>
<td>0.044</td>
<td>0.0005</td>
<td>1.2 x 10⁻⁷</td>
</tr>
<tr>
<td>ΔConcentration (ca. 2000) / % pa</td>
<td>0.45</td>
<td>0.60</td>
<td>0.25</td>
<td>ca. 5.0</td>
<td>ca. 6.3</td>
</tr>
<tr>
<td>Radiative forcing $a_o / W m^{-2} ppbv^{-1}$</td>
<td>1.68 x 10⁻³</td>
<td>4.59 x 10⁻⁴</td>
<td>3.41 x 10⁻³</td>
<td>0.32</td>
<td>0.60</td>
</tr>
<tr>
<td>Total radiative forcing $^a / W m^{-2}$</td>
<td>1.46</td>
<td>0.48</td>
<td>0.15</td>
<td>0.16</td>
<td>7.2 x 10⁻³</td>
</tr>
<tr>
<td>Lifetime $^b / years$</td>
<td>50-200 $^c$</td>
<td>12</td>
<td>120</td>
<td>102</td>
<td>ca. 1000</td>
</tr>
<tr>
<td>Global warming potential $^d$</td>
<td>1</td>
<td>23</td>
<td>296</td>
<td>10600</td>
<td>&gt; 17500</td>
</tr>
<tr>
<td>Contribution to greenhouse effect / %</td>
<td>52</td>
<td>17</td>
<td>5</td>
<td>ca. 15 $^e$</td>
<td>&lt;&lt; 0.01</td>
</tr>
</tbody>
</table>

$^a$ Due to change in concentration of greenhouse gas from the pre-Industrial era to the present time.

$^b$ Assumes a single-exponential decay for removal of greenhouse gas from the atmosphere.

$^c$ CO₂ does not show a single-exponential decay [37,38].

$^d$ Calculated for a time period of 100 years.

$^e$ Cumulative effect of all chlorofluorocarbons.
<table>
<thead>
<tr>
<th>Fragment ion</th>
<th>( AE_{298} / \text{eV} )</th>
<th>Neutral partner</th>
<th>( \Delta r H^0 ) / \text{eV}</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Major ions</strong>(^a)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \text{CF}_3^+ )</td>
<td>12.92 ± 0.18</td>
<td>( \text{SF}_5 )</td>
<td>12.90</td>
</tr>
<tr>
<td>( \text{SF}_5^+ )</td>
<td>13.9 ± 1.2</td>
<td>( \text{CF}_3 )</td>
<td>13.67</td>
</tr>
<tr>
<td><strong>Minor ions</strong>(^b)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \text{SF}_4^+ )</td>
<td>13.5 ± 1.3</td>
<td>( \text{CF}_4 )</td>
<td>12.42</td>
</tr>
<tr>
<td>( \text{SF}_3^+ )</td>
<td>14.94 ± 0.13</td>
<td>( \text{CF}_4 + \text{F} )</td>
<td>12.36</td>
</tr>
<tr>
<td>( \text{CF}_2^+ )</td>
<td>16.0 ± 2.0</td>
<td>( \text{SF}_6 )</td>
<td>15.22</td>
</tr>
</tbody>
</table>

\(^a\) A *major* ion is defined as one formed by a single bond cleavage; formation of \( \text{CF}_3^+ \) and \( \text{SF}_5^+ \) involves cleavage of the S-C bond.

\(^b\) A *minor* ion is defined as one in which intramolecular rearrangement must occur. Note that the formation of \( \text{SF}_4^+ \), \( \text{SF}_3^+ \) and \( \text{CF}_2^+ \) is only possible at their observed appearance energies given in Column 2 if migration of \( \text{F}^- \) across the S-C bond occurs in the transition state to form \( \text{CF}_4 \) and \( \text{SF}_6 \), respectively, as accompanying neutrals.
Table 3  Total mean translational kinetic energy release, \( <\text{KE}>_T \), for the two-body fragmentation of the valence states of SF\(_5\)CF\(_3^+\)

<table>
<thead>
<tr>
<th>Fragment</th>
<th>( E ) / eV</th>
<th>( E_{\text{avail}} ) / eV (^a)</th>
<th>( &lt;\text{KE}&gt;_T ) / eV</th>
<th>( f_{\text{expt}} )</th>
<th>( f_{\text{statistical}} ) (^b)</th>
<th>( f_{\text{pure impulsive}} ) (^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CF(_3^+) (^d)</td>
<td>14.25</td>
<td>1.52</td>
<td>0.32 (5)</td>
<td>0.21</td>
<td>0.04</td>
<td>0.20</td>
</tr>
<tr>
<td></td>
<td>15.69</td>
<td>2.96</td>
<td>0.29 (2)</td>
<td>0.10</td>
<td>0.04</td>
<td>0.20</td>
</tr>
<tr>
<td></td>
<td>16.98</td>
<td>4.25</td>
<td>0.38 (1)</td>
<td>0.09</td>
<td>0.04</td>
<td>0.20</td>
</tr>
<tr>
<td></td>
<td>17.97</td>
<td>5.24</td>
<td>0.40 (1)</td>
<td>0.08</td>
<td>0.04</td>
<td>0.20</td>
</tr>
<tr>
<td></td>
<td>19.07</td>
<td>6.34</td>
<td>0.37 (1)</td>
<td>0.06</td>
<td>0.04</td>
<td>0.20</td>
</tr>
</tbody>
</table>

\(^a\) \( E_{\text{avail}} \) is defined in equ. (VII). The average thermal energy of SF\(_5\)CF\(_3\) is 0.17 eV.

\(^b\) From Klots [50].

\(^c\) From Holdy, Klots and Wilson [42].

\(^d\) Assumes the neutral fragment is SF\(_5\).
Table 4  Rate coefficients, cation product ions and their branching ratios for the reactions of cations with possible atmospheric importance with SF₅CF₃. The recombination energies of the reagent cations are listed in eV in brackets under the cations. The capture rate coefficients, calculated using Langevin theory [57], are presented in square brackets under the experimental values. 0 K enthalpies of reaction for these channels are given in column 5. For formation of SF₃⁺ and SF₄⁺, two values for ∆rH₀ are given. The first value is calculated using the data set of Fisher et al. [43], whereas the second corresponds to that of Irikura [47]. The two values given for ∆rH₀ for formation of SF₅⁺ by dissociative charge transfer result from taking the IE (SF₅) to be either 9.60 eV [43] or 9.71 eV [47].

<table>
<thead>
<tr>
<th>Reagent Ion</th>
<th>Rate Coefficient / x 10⁻⁹ cm³ s⁻¹</th>
<th>Product Ion / %</th>
<th>Proposed Neutral Products</th>
<th>∆rH₀ / kJ mol⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>N₂⁺ (15.58)</td>
<td>1.6 [1.4]</td>
<td>CF₅⁺ (65)</td>
<td>SF₅ + N₂</td>
<td>-258</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (28)</td>
<td>CF₄ + F + N₂</td>
<td>-248, -289</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₄⁺ (trace)</td>
<td>CF₄ + N₂</td>
<td>-283, -345</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₄⁺ CF₃ (2)</td>
<td>F + N₂</td>
<td>?</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (5)</td>
<td>CF₃ + N₂</td>
<td>-185, -175</td>
</tr>
<tr>
<td>N⁺ (14.53)</td>
<td>2.2 [1.9]</td>
<td>CF₅⁺ (80)</td>
<td>SF₅ + N</td>
<td>-157</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (17)</td>
<td>CF₄ + F + N</td>
<td>-147, -187</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₄⁺ (trace)</td>
<td>CF₄ + N</td>
<td>-181, -243</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (3)</td>
<td>CF₃ + N</td>
<td>-84, -73</td>
</tr>
<tr>
<td>CO⁺ (14.01)</td>
<td>1.6 [1.4]</td>
<td>CF₅⁺ (75)</td>
<td>SF₅ + CO</td>
<td>-107</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (22)</td>
<td>CF₄ + F + CO</td>
<td>-96, -137</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₄⁺ (trace)</td>
<td>CF₄ + CO</td>
<td>-131, -193</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (3)</td>
<td>CF₃ + CO</td>
<td>-34 (± 43), -23 (± 46)</td>
</tr>
<tr>
<td>CO₂⁺ (13.77)</td>
<td>1.2 [1.1]</td>
<td>CF₅⁺ (76)</td>
<td>SF₅ + CO₂</td>
<td>-84</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (14)</td>
<td>CF₄ + F + CO₂</td>
<td>-73, -114</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₄⁺ (8)</td>
<td>CF₄ + CO₂</td>
<td>-108, -170</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (2)</td>
<td>CF₃ + CO₂</td>
<td>-11 (± 43), 0 (± 46)</td>
</tr>
<tr>
<td>O⁺ (13.62)</td>
<td>1.9 [1.8]</td>
<td>CF₅⁺ (83)</td>
<td>SF₅ + O</td>
<td>-69</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (16)</td>
<td>CF₄ + F + O</td>
<td>-59, -100</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₄⁺ (trace)</td>
<td>CF₄ + O</td>
<td>-93, -156</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (1)</td>
<td>CF₃ + O</td>
<td>4 (± 43), 14 (± 46)</td>
</tr>
<tr>
<td>N₂O⁺ (12.89)</td>
<td>1.1 [1.1]</td>
<td>CF₅⁺ (75)</td>
<td>SF₅ + N₂O</td>
<td>1 (± 39)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (19)</td>
<td>CF₄ + F + N₂O</td>
<td>12 (± 45), -29 (± 47)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₄⁺ (5)</td>
<td>CF₄ + N₂O</td>
<td>-23 (± 45), -85</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (1)</td>
<td>CF₃ + N₂O</td>
<td>74, 84</td>
</tr>
<tr>
<td>H₂O⁺ (12.61)</td>
<td>1.6 [1.7]</td>
<td>CF₅⁺ (8)</td>
<td>SF₅ + H₂O</td>
<td>28 (± 39)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (92)</td>
<td>CF₄ + F + H₂O</td>
<td>39 (± 45), -2 (± 47)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₄⁺ (trace)</td>
<td>CF₄ + H₂O</td>
<td>4 (± 45), -58</td>
</tr>
<tr>
<td>O₂⁺ (12.07)</td>
<td>0.01 [1.3]</td>
<td>CF₅⁺ (63)</td>
<td>SF₅ + O₂</td>
<td>80, 62, 43 (±39)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (31)</td>
<td>CF₄ + F + O2</td>
<td>91, 50 (± 47)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₄⁺ (2)</td>
<td>CF₄ + O₂</td>
<td>56, -6 (± 46)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SF₅⁺ (4)</td>
<td>CF₃ + O₂</td>
<td>153, 164</td>
</tr>
</tbody>
</table>

a The three values given are for the ν = 0, 1 and 2 vibrational levels of O₂⁺, respectively.
<table>
<thead>
<tr>
<th>Reaction</th>
<th>$\Delta_fH^0_0$ / kJ mol(^{-1}) (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Scheme A</td>
</tr>
<tr>
<td>SF(_5)CF(_3) + e(^-) (0 eV) $\rightarrow$ SF(_5)^- (-1282) (^c) + CF(_3) (-463) (^c)</td>
<td>$-122$ (-1.26)</td>
</tr>
<tr>
<td>(−1623 or −1750)</td>
<td></td>
</tr>
<tr>
<td>$\rightarrow$ CF(_3)^- (-639) + SF(_5) (-915)</td>
<td>+69 (+0.72)</td>
</tr>
<tr>
<td>SF(_5)CF(_3) + h(\nu) (10.2 eV) $\rightarrow$ SF(_5) (-915) + CF(_3) (-463)</td>
<td>$-739$ (-7.66)</td>
</tr>
<tr>
<td>(−1623 or −1750)</td>
<td></td>
</tr>
<tr>
<td>$\rightarrow$ CF(_3)^+ (+409) + SF(_5)^- (-1282)</td>
<td>$-234$ (-2.43)</td>
</tr>
<tr>
<td>$\rightarrow$ SF(_5)^+ (+29) + CF(_3)^- (-639)</td>
<td>+29 (+0.30)</td>
</tr>
</tbody>
</table>

\(^a\) Scheme A assumes $\Delta_fH^0_0$ (SF\(_5\)CF\(_3\)) = −1623 kJ mol\(^{-1}\) (theory [32]).

\(^b\) Scheme B assumes $\Delta_fH^0_0$ (SF\(_5\)CF\(_3\)) = −1750 kJ mol\(^{-1}\) (experiment: updated value from [20]).

\(^c\) Values in brackets are $\Delta_fH^0_0$ in kJ mol\(^{-1}\).
Table 6  Thermal electron attachment rate constants, absorption cross-sections at 121.6 nm, and atmospheric lifetimes for CF₄, SF₆ and SF₅CF₃.

<table>
<thead>
<tr>
<th>Perfluoro compound</th>
<th>$k_e$ (298 K) / cm³ s⁻¹</th>
<th>$\sigma_{121.6}$ / cm²</th>
<th>lifetime / years</th>
</tr>
</thead>
<tbody>
<tr>
<td>CF₄</td>
<td>$&lt; 10^{-16}$&lt;sup&gt;a&lt;/sup&gt;</td>
<td>$&lt; 8 \times 10^{-22}$&lt;sup&gt;b&lt;/sup&gt;</td>
<td>$&gt; 50000$</td>
</tr>
<tr>
<td>SF₆</td>
<td>$2.38 \times 10^{-7}$&lt;sup&gt;c&lt;/sup&gt;</td>
<td>$1.76 \times 10^{-18}$&lt;sup&gt;b&lt;/sup&gt;</td>
<td>$&gt; 800$</td>
</tr>
<tr>
<td>SF₅CF₃</td>
<td>$7.7 \times 10^{-8}$&lt;sup&gt;d&lt;/sup&gt;</td>
<td>$1.3 \times 10^{-17}$&lt;sup&gt;e&lt;/sup&gt;</td>
<td>$ca. 1000$</td>
</tr>
</tbody>
</table>

<sup>a</sup> Schumacher et al. [69].
<sup>b</sup> Ravishankara et al. [65].
<sup>c</sup> Mayhew et al. [62].
<sup>d</sup> Kennedy and Mayhew [59].
<sup>e</sup> Re-analysis of the data of Chim et al. [16], resulting in a reduction of $\sigma_{121.6}$ by 2 Mb.
Figure 1: Electron density map of the highest occupied molecular orbital of SF₅–CF₃. The FSF and FCF bond angles are all either 90° or 109.3°. The S–C bond length is 0.186 nm, the three C–F lengths all 0.130 nm, the three S–F_equatorial lengths are 0.158 nm, and the two S–F_axial lengths are 0.157 nm. The dipole moment is calculated to be 0.95 Debye.

Figure 2: Definition of the dissociative ionisation energy (DIE) of a generalised polyatomic molecule AB–CD. In the particular application to SF₅CF₃, AB represents CF₃, CD represents SF₅.

Figure 3: Extrapolation method to determine the first dissociative ionisation energy of AB–CD.

Figure 4: Schematic of the threshold photoelectron photoion coincidence apparatus. The apparatus can also study coincidences between threshold photoelectrons or mass-resolved cations with fluorescence photons, but these modes of operation are not used in this work.

Figure 5: Results of the extrapolation experiment for CF₃⁺ + SF₅ from SF₅CF₃. Only datapoints that define the Franck-Condon region of the ground state of SF₅CF₃⁺ are used to define the slope and intercept of the graph [17]. The slope is 0.19, and the first dissociative ionisation energy of SF₅CF₃ is determined to be 12.9 ± 0.4 eV. (Reproduced by permission from J. Phys. Chem. A., 2001, 105, 8403-8412.)

Figure 6: TPEPICO-TOF spectrum (closed circles) for CF₃⁺ / SF₅CF₃ recorded at a photon energy of 14.09 eV. Shown as a solid line, the data fit to a mean translational KE release, <KE>_T, of 0.24 ± 0.05 eV. (Reproduced by permission from J. Phys. Chem. A., 2001, 105, 8403-8412.)

Figure 7: Threshold photoelectron spectrum of SF₅CF₃ recorded at a resolution of 0.3 nm. Assignment of the bands to the ground state (X) and excited electronic states (A, B, C, etc.) of SF₅CF₃⁺ is shown. (Reproduced by permission from J. Phys. Chem. A., 2001, 105, 8403-8412.)

Figure 8: Time-of-flight spectrum of fragment ions from SF₅CF₃ integrated over the photoexcitation energies 12.7-26.4 eV. (Reproduced by permission from J. Phys. Chem. A., 2001, 105, 8403-8412.)

Figure 9: Schematic of the selected ion flow tube used to determine rate coefficients and product ion yields of ion-molecule reactions. (Reproduced by permission of Dr C R Howle, PhD thesis, University of Birmingham, 2004.)

Figure 10: Schematic of the electron swarm drift tube used in Birmingham for the SF₅CF₃ measurements. A small pinhole at the right-hand end of the drift tube leads via differential pumping, to a quadrupole mass spectrometer for detection of the anionic products. (Reproduced by permission from Int. J. Mass Spectrom., 2001, 205, 253-270.)

Figure 11: Graph of electron attachment rate coefficient, k_a, as a function of mean electron energy for SF₅CF₃ in atmospheric pressure of N₂ (<ε> less than 0.5 eV) and Ar (<ε> greater than 0.5 eV) buffer gas. (Reproduced by permission from Int. J. Mass Spectrom., 2001, 206, i-iv.)

Figure 12: Apparatus used to record the vacuum-UV absorption spectrum of SF₅CF₃. (Reproduced by permission from Chem. Phys., 2000, 260, 237-247.)

Figure 13: Absorption spectrum of SF₅CF₃ recorded with a photon resolution of 0.08 nm. The absorption cross section at 121.6 nm is 1.3 ± 0.2 x 10⁻¹⁷ cm² or 13 ± 2 Mb. (Reproduced by permission from Chem. Phys. Letts., 2003, 367, 697-703.)
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