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Products and mechanism of the OH-initiated photo-oxidation of perfluoro ethyl vinyl ether, $C_2F_5OCF=CF_2$

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The OH-initiated photo-oxidation of perfluoro ethyl vinyl ether ($C_2F_5OCF=CF_2$, PEVE) in air (298 K, 50 and 750 Torr total pressure) was studied in a photochemical reactor using *in situ* detection of PEVE and its products by Fourier transform IR absorption spectroscopy. The relative rate technique was used to derive the rate coefficient, k_1 , for the reaction of PEVE with OH as $k_1 = (2.8 \pm 0.3) \times 10^{-12}$ cm³ molecule⁻¹ s⁻¹. The photo-oxidation of PEVE in the presence of NO_x at 1 bar results in formation of C_2F_5OCFO , FC(O)C(O)F and CF_2O in molar yields of 0.50 \pm 0.07, 0.46 \pm 0.07 and 1.50 \pm 0.22, respectively. FC(O)C(O)F and CF_2O are formed partially in secondary, most likely heterogeneous processes. At a reduced pressure of 50 Torr, the product distribution is shifted towards formation of FC(O)C(O)F, indicating the important role of collisional quenching of initially formed association complexes, and enabling details of the reaction mechanism to be elucidated. An atmospheric photo-oxidation mechanism for PEVE is presented and the environmental implications of PEVE release and degradation are discussed.

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1 Introduction

The thermal and chemical resistance of fluoropolymers has led to their use in many industrial processes and to their production in large quantities^{1,2} and thus to their release to the environment. We recently presented³ the first detailed kinetic study on the reaction of perfluoro ethyl vinyl ether (C₂F₅OCF=CF₂, henceforth referred to as PEVE) with OH radicals (R1), determining the rate coefficient, k_1 , using absolute and relative-rate techniques. k_1 was found to be independent of pressure but it has a significant negative temperature dependence, indicating that the first step is addition of OH to the C=C double bond. A room temperature value of k_1 close to $3 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ implies an atmospheric lifetime of a few days, indicating that PEVE is not an important greenhouse gas. The main environmental concern about fluorinated vinyl ethers is whether their photo-oxidation leads to formation of persistent, fluorinated pollutants in the environment.

The products formed in the atmospheric photo-oxidation of organic trace gases depends on the fate of peroxy-radicals formed in the reaction between the initially generated organic radical fragments and O_2 . The chemistry is typically described as either "high NO_x " where the peroxy radicals react with NO to form oxy radicals, which then propagate the radical chain, or "low NO_x " where the dominating fate of the peroxy radical is reaction with HO_2 or other peroxy radicals. Most short lived anthropogenic pollutants are oxidized under high NO_x conditions, but as the two limiting cases have different mechanisms, differences in their product distribution provide valuable information about key reaction steps.

Extending our kinetics study, this work is a detailed examination of the products formed in the OH-initiated atmospheric degradation of PEVE in air, with the aim of deriving a detailed reaction mechanism applicable to atmospheric conditions (both high and low NO_x). We also present a new measurement of the rate coefficient, k_1 , for the title reaction.

$$OH + PEVE \rightarrow products$$
 (R1)

2 Experimental methods

The experimental set-up used to study the OH-initiated photooxidation of PEVE has been described in detail previously⁴ and only a brief summary of its central components is given here. The reaction volume is a 44 l cylindrical quartz chamber equipped with a White-type multiple reflection mirror system resulting in a 43.7 m optical path length. Infrared spectra of the

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cell contents were recorded using a Bruker Vector FTIR instrument with a liquid N2-cooled MCT detector. The spectrometer was operated at a resolution of 0.5 cm⁻¹ using boxcar apodization.

The quartz reactor was at room temperature (298 \pm 2 K) and for most experiments 1 bar total pressure. Six external, radially mounted, UV photolysis lamps provided a homogeneous light flux within the reactor for radical generation.

For the determination of k_1 and for the product study in high-NO_x conditions, OH radicals were generated by photolysis of methyl nitrite $(2-10 \times 10^{14} \text{ molecule cm}^{-3})^5$ in the presence of up to 4×10^{14} molecule cm⁻³ NO using six Philips TL08 lamps ($\lambda_{\rm max} \sim 350$ nm):

$$CH_3ONO + h\nu \rightarrow CH_3O + NO$$
 (R2)

$$CH_3O + O_2 \rightarrow CH_2O + HO_2$$
 (R3)

$$HO_2 + NO \rightarrow OH + NO_2$$
 (R4)

Under these conditions, the dominant fate of the organic peroxy radicals formed will be reaction with NO.

As the methyl nitrite based OH generation scheme is unsuitable for NO_x free photo-oxidation, we also generated OH radicals by photolysis of ozone $(1-2 \times 10^{14} \text{ molecule cm}^{-3})$ at 254 nm in a large excess of hydrogen (up to 2×10^{17} molecule cm⁻³). The large excess of H₂ ensures that O(1D) does not react with PEVE. As desribed previously, variations in [H₂] in this range did not lead to changes in the observed kinetics in a related system, confirming that unwanted O(1D) reactions were not important.

$$O_3 + h\nu \to O(^1D) + O_2$$
 (R5)

$$O(^{1}D) + H_{2} \rightarrow OH + H \tag{R6}$$

$$OH + H_2 \rightarrow H_2O + H \tag{R7}$$

$$H + O_2 + M \rightarrow HO_2 + M \tag{R8}$$

$$H + O_3 \rightarrow OH + O_2$$
 (R9)

$$HO_2 + O_3 \rightarrow OH + 2O_2$$
 (R10)

This OH radical generation scheme generates both OH and HO₂ radicals, and under these conditions, the organic peroxy radicals formed in the PEVE photo-oxidation will either react with HO2 radicals or undergo self-reaction.

Chemicals

Synthetic air (Westfalen), hydrogen (Westfalen 99.999%), ethylene (Matheson, Res. Pur.) and PEVE (Merck, 99%) were used as supplied without further purification. Ozone was generated by flowing air over a low-pressure Hg "Penray" lamp (thereby dissociating O2 to ground state O-atoms, which recombine with O_2 to form O_3) before entering the reaction cell.

Methyl nitrite was prepared by the drop-wise addition of 50% sulphuric acid (Sigma-Aldrich) to a saturated solution of sodium nitrite in 50% methanol (Merck) in a three-necked round flask with a stirring magnet at 0 °C. To collect the resulting methyl nitrite, a slow flow of nitrogen was passed over the reaction mixture and into a trap at -78 °C. The resulting yellow liquid

was then purified by vacuum distillation. Finally, a gas-phase sample of CH₃ONO, diluted in nitrogen was prepared and stored in a blackened glass bulb at 298 K. The IR spectrum of a dilute gas-phase sample of methyl nitrite revealed minor impurities of methanol only (<0.3%).

3 Results

3.1 Infrared spectrum of PEVE

The IR spectrum of PEVE is characterized by a progression of strong absorption bands in the range 1050-1400 cm⁻¹ (shown in Fig. 1A), a weak C=C stretching mode at 1842 cm⁻¹ and weaker absorption features at 749 to 840 cm⁻¹. The spectrum closely resembles that of perfluoro methyl vinyl ether (CF₃OCF=CF₂, henceforth referred to as PMVE) previously reported and assigned.6

The reference spectrum of PEVE was obtained by recording spectra at different pressures of accurately diluted mixtures of PEVE (0.3 to 3.0 \times 10¹³ molecule cm⁻³) and converting to absorption cross-sections after baseline correction and checking for Beer-Lambert behaviour.

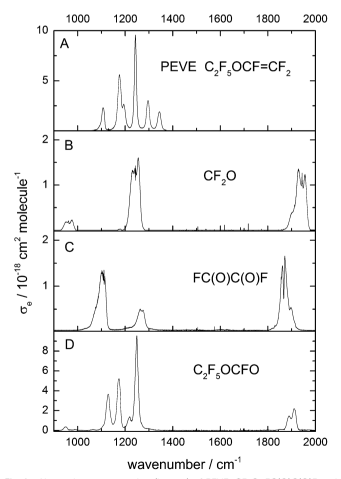


Fig. 1 Absorption cross section (base e) of PEVE, CF₂O, FC(O)C(O)F and C_2F_5OCFO in the C-F and C-O stretching region. The PEVE and C_2F_5OCFO spectra were obtained in this study, those of CF₂O and CF(O)CFO have been reported previously.7

3.2 Infrared spectra of photo-oxidation products

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Using previously reported reference spectra (Fig. 1) 7 CF₂O and FC(O)C(O)F were readily identified as products of the OH-initiated oxidation of PEVE. The product spectrum obtained following PEVE oxidation showed strong absorption features at 1100–1300 cm $^{-1}$ in addition to a progression of carbonyl absorption features at 1800–2000 cm $^{-1}$.

After subtracting features due to CF2O and FC(O)C(O)F (Fig. 1 panels B and C) the resulting spectrum is characterised by three strong absorptions at 1128, 1173 and 1249 cm⁻¹, one at 1899 cm⁻¹ and some weaker absorptions at 720, 757, 853, 950 and 1218 cm⁻¹ (Fig. 1D). The spectrum has some similar spectral features to PEVE, but without the two vinylic C-F stretching modes at 1345 and 1296 cm⁻¹. The other features in the range 1000-1300 cm⁻¹ are somewhat blue-shifted compared to the PEVE spectrum and the weak C=C-stretching absorption band at 1842 cm⁻¹ is replaced by a much stronger absorption at 1899 cm⁻¹. The initial assignment of this spectrum to C₂F₅OCFO was based on similarities to the spectra of perfluoromethyl fluoroformate.8 This assignment was independently confirmed by experiments in which the oxidation of PEVE was initiated by reaction with Cl-atoms rather than OH. In this case, only CF₂O and C₂F₅OCFO are observed as products (see Fig. 2). This is in agreement with the results of Mashino et al. who observed CF₃OCFO as a photo-oxidation product of the Cl initiated oxidation of PMVE with a yield of 100%. The reasons for the simplified product distribution when using Cl rather than OH to initiate the oxidation of PEVE is discussed later when we examine details of the reaction mechanism.

PEVE +
$$Cl(O_2) \rightarrow C_2F_5OCFO + CF_2O$$
 (R11)

As each PEVE lost by reaction with Cl forms one molecule of C_2F_5OCFO , we were able to put the C_2F_5OCFO spectrum on an absolute basis. Once spectral features corresponding to water and CF_2O had been subtracted, the calibrated C_2F_5OCFO absorption spectrum was obtained, as displayed in Fig. 1D.

3.3 Relative-rate measurements of k_1

In our previous paper³ on the kinetics of the reaction between OH and PEVE we reported a rate coefficient under NO_x free conditions in 50 and 100 Torr N_2 and in 1 bar air. In this work we have measured the rate coefficient of PEVE with OH radicals relative to that of ethylene with NO_x present.

In the absence of loss processes other than reaction with OH, the depletion factors, $\ln(\text{initial concentration/concentration})$, for reactant and reference are described by:

$$\ln \left(\frac{[\text{PEVE}]_0}{[\text{PEVE}]_t} \right) = \frac{k_1}{k_{12}} \ln \left(\frac{[\text{ethylene}]_0}{[\text{ethylene}]_t} \right)$$
 (i)

[PEVE]₀, [ethylene]₀, [PEVE]_t and [ethylene]_t are the concentrations of PEVE and ethylene at times t_0 and t respectively, and k_1 and k_{12} are the rate coefficients of reactions (1) and (12), respectively.

$$PEVE + OH \rightarrow Products$$
 (R1)

$$C_2H_4 + OH \rightarrow Products$$
 (R12)

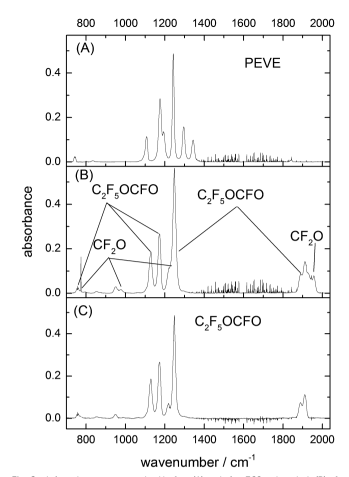


Fig. 2 Infrared spectrum acquired before (A) and after 360 s photolysis (B) of a mixture of 3.3 $\times~10^{13}$ molecule cm $^{-3}$ PEVE and 2.6 $\times~10^{14}$ molecule cm $^{-3}$ Cl₂. Panel (C) shows the spectrum of C₂F₅OCFO obtained by subtracting water and CF₂O.

The initial concentration ranges used for PEVE and ethylene were (1.3–2.9) \times 10^{13} molecule cm^{-3} and (0.8–2.9) \times 10^{14} molecule cm^{-3} , respectively. A potential interference in relative rate measurements of fluorocarbons is the reaction of CF $_3$ O radicals with organics like ethylene. 9

$$CF_3O + C_2H_4 \rightarrow Products$$
 (R13)

To minimise the impact of (R13), high concentrations ((1–4) \times 10¹⁴ molecule cm⁻³) of NO were added to the mixture. NO reacts rapidly with CF₃O (k_{14} (298 K) = 5.4 \times 10⁻¹¹ cm³ molecule⁻¹ s⁻¹) forming CF₂O and FNO.^{10,11}

$$CF_3O + NO \rightarrow CF_2O + FNO$$
 (R14)

The relative concentrations of NO and C_2H_4 used, and the rate coefficients for their reactions with CF_3O , ensure that CF_3O reacts predominantly (to 80%) with NO and variation of [NO] by a factor of four did not result in any significant changes in the measured rate coefficients. We thus conclude that our derivation of k_1 is not significantly influenced by undesired reactions of CF_3O radicals.

A typical experiment lasted around one hour during which the photolysis lamps were switched on for 5–10 periods

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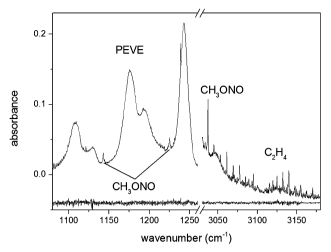


Fig. 3 FTIR spectrum from a relative rate experiment in the spectral ranges used in the analysis with residuals from the fitting of reference spectra.

of 10-300 s. The intermittent acquisition of FTIR spectra (128 scans at 0.5 cm⁻¹ resolution) took ≈ 5 min. The time dependent depletion factors for PEVE and C2H4 were obtained by least squares fitting to reference spectra in the range 1080–1260 cm⁻¹ for PEVE and 3030-3180 cm⁻¹ for ethylene. Absorption by reaction products were accounted for by including reference spectra of CF₂O, FC(O)C(O)F and C₂F₅OCFO as well as CH₃ONO and its degradation products in the fit procedure (DOASIS¹²). An example spectrum with fit residuals is shown in Fig. 3.

A total of six relative rate experiments were performed under different experimental conditions, which are summarised in Table 1 and Fig. 4. A linear, least squares fit to all data (solid line in Fig. 4) gives $k_1/k_{12} = 0.361 \pm 0.006$ (2 σ). From this k_1 can be determined using an evaluated literature value¹³ for k_{11} $(7.8 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at 298 K and 1 bar pressure})$ to give: $k_1 = (2.8 \pm 0.3) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ where the}$ uncertainty includes assessment of the accuracy of the reference rate coefficient.¹³ The rate coefficient is in good agreement with the value $(k_1 = (3.0 \pm 0.3) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$ from our previous kinetic study on the title reaction³ and for other perfluoro vinyl ethers for which the OH-rate coefficients fall in the range $(2.2-3.4) \times 10^{-12}$ cm³ molecule⁻¹ s⁻¹.^{7,14,15} The rate coefficients and trends therein for different vinyl ethers are discussed in our previous publication³ and the interested reader is referred to that publication for details.

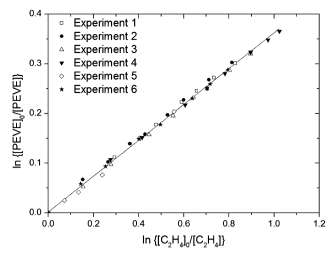


Fig. 4 Decay of PEVE versus C₂H₄ in the presence of OH radicals from six individual experiments in 750 Torr air diluent. The conditions for each experiment are listed in Table 1.

3.4 Reaction products and yields (Φ_{NO_x}) in air (with NO_x)

Products of the OH-initiated photo-oxidation of PEVE were detected and quantified in a separate series of experiments. In these experiments, NO is generated in the photolysis of CH₃ONO and in some experiments was also added to the initial reaction mixture.

As mentioned above, C₂F₅OCFO FC(O)C(O)F and CF₂O were observed as products. The experiments were conducted with several photolysis periods interrupted by spectrum acquisition so that the time dependence of PEVE loss and product formation could be analysed. Time dependent concentrations of PEVE were obtained by fitting post photolysis spectra to reference spectra between 1080 and 1360 cm⁻¹. Not only PEVE but also the fluorocarbon products (CF2O, FC(O)C(O)F and C₂F₅OCFO) absorb in this range, as do CH₃ONO, CH₃ONO₂ and HNO₃ and they were thus included in the spectral deconvolution procedure. Compared to PEVE, the fluorinated products absorb only weakly in this range (Fig. 1) and the concentrations of CF₂O, FC(O)C(O)F and C₂F₅OCFO were therefore obtained by analysing the spectrum between 1855 and 2000 cm⁻¹. In this range, the spectra of CF2O, FC(O)C(O)F and C2F5OCFO have very distinct carbonyl stretching modes (Fig. 1).

Concentration time profiles for PEVE, CF₂O, FC(O)C(O)F and C₂F₅OCFO from a single experiment are shown in Fig. 5.

Table 1 Rate coefficients and experimental conditions for the relative rate experiments

Expt no.	P (Torr)	$[PEVE]^a$	$[C_2H_4]^a$	$[\mathrm{CH_3ONO}]^a$	$[NO]^a$	$[OH]^b$	$k_{\mathrm{rel}}{}^{c}$
1	750	2.9	0.8	8.6	2.0	3.7	0.367 ± 0.010
2	750	2.8	1.5	8.2	2.1	7.4	0.368 ± 0.010
3	750	1.3	1.5	9.7	2.1	5.8	0.359 ± 0.020
4	750	2.9	1.5	9.8	1.0	11.9	0.354 ± 0.006
5	750	2.7	1.5	8.9	3.9	2.8	0.369 ± 0.016
6	750	2.8	2.9	8.6	2.0	5.6	0.358 ± 0.008

 $[^]a$ The initial concentrations of PEVE (units of 10^{13} molecule cm $^{-3}$), C_3H_8 , NO and CH_3ONO (units 10^{14} molecule cm $^{-3}$) were determined from pressure measurements. b The OH concentration (units 10^8 molecule cm $^{-3}$) was determined from the initial decay of PEVE. c 2σ statistical error.

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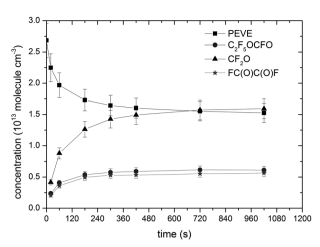


Fig. 5 Concentration—time profiles of PEVE loss and simultaneous generation of fluorinated organic products from an experiment at 1 bar air in the presence of NO_x . The error bars reflect 10% uncertinty in the cross-sections used.

A plot of $\Delta[\text{product}]$ *versus* $\Delta[\text{PEVE}]$ is displayed for a single experiment in Fig. 6. Fig. 6 illustrates that the plots of $\Delta[\text{C}_2\text{F}_5\text{OCFO}]$ and $\Delta[\text{FC}(\text{O})\text{CFO}]$ *versus* $\Delta[\text{PEVE}]$ are straight lines, from which the product yields can be directly obtained. The experimental conditions and product yields (Φ_{NO_x}) from the total of seven experiments carried out in the presence of NO_x are given in Table 2. In contrast to C₂F₅OCFO and FC(O)CFO, the plot of $\Delta[\text{CF}_2\text{O}]$ *versus* $\Delta[\text{PEVE}]$ is non-linear and has a significant induction period before a roughly constant slope is achieved (in this case after $\approx 1 \times 10^{13}$ PEVE has been reacted). This is a clear indication that CF₂O is formed in secondary reactions. The yield of a product which is formed directly (or very rapidly) will be independent of time, whereas one which is formed in slower, secondary processes will show more complex time dependence.

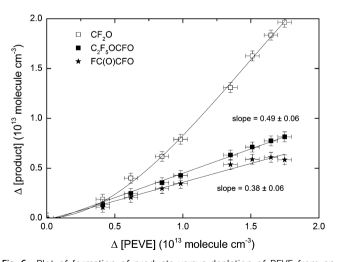


Fig. 6 Plot of formation of products *versus* depletion of PEVE from an experiment on the OH-initiated oxidation of PEVE in the presence of NO_x. The slopes of the linear fits are the yields $(\pm 2\sigma)$ of C₂F₅OCFO and FC(O)CFO from this particular experiment. The polynomial fit to the CF₂O data is added to emphasise the non-linearity at low PEVE conversions.

Time dependent yields of each of CF₂O, FC(O)C(O)F and C₂F₅OCFO are displayed in Fig. 7. Whereas the yields of C₂F₅OCFO and FC(O)C(O)F are roughly constant, that of CF₂O acquires a maximum value after several hundred seconds. The CF₂O yields reported in Table 2 are averages obtained from the plateau after ≈ 240 s. Combining an estimated 10% uncertainty in absorption cross-sections with experimental scatter results in the following, average molar yields of products obtained in the presence of NO_x at 1 bar pressure of air: $\Phi_{\text{NO}_x}(\text{FC}(\text{O})\text{C}(\text{O})\text{F}) = 0.46 \pm 0.07$, $\Phi_{\text{NO}_x}(\text{C}_2\text{F}_5\text{OCFO}) = 0.50 \pm 0.07$ and $\Phi_{\text{NO}_x}(\text{CF}_2\text{O}) = 1.50 \pm 0.22$. The experimental CF₂O yields are more variable than those of FC(O)C(O)F and C₂F₅OCFO, which is related to its production both in gas-phase and heterogeneous reactions (see below).

For FC(O)C(O)F and C₂F₅OCFO there is no significant difference in the yield between the different experiments, with or without added NO. In contrast, as illustrated in Fig. 7, the CF₂O yields obtained at 1 bar pressure of air were largest when NO was added. The yields of all products were sensitive to the total pressure with values of $\Phi_{NO_x}(FC(O)C(O)F) = 0.69 \pm 0.15$, $\Phi_{NO_x}(C_2F_5OCFO) = 0.11 \pm 0.03$ and $\Phi_{NO_x}(CF_2O) = 1.78 \pm 0.25$ obtained at 50 Torr air.

Possible reasons and mechanistic implications of the [NO] and pressure dependence are discussed below along with the overall degradation mechanism.

3.5 Reaction products and yields (Φ) in air (without NO_x)

The formation of products in the OH-initiated oxidation of PEVE in the absence of NO_x was studied in a separate series of experiments where OH radicals were generated by the photolysis of ozone in a large excess of hydrogen (see Section 2). The initial concentration range used for PEVE was $(1.1–3.7) \times 10^{13}$ molecule cm⁻³. The concentrations of ozone and hydrogen were $(0.7–2.2) \times 10^{14}$ and $(0.8–2.2) \times 10^{17}$ molecule cm⁻³, respectively. The fluorinated products observed, C_2F_5OCFO , CF_2O and FC(O)C(O)F, were the same as those found in the CH_3ONO/NO experiments. The molar yield of C_2F_5OCFO was similar, with $\Phi(C_2F_5OCFO) = (0.46 \pm 0.07)$ whereas that of CF_2O was slightly lower with $\Phi(CF_2O) = 0.9–1.3$. A further difference was the significant depletion of [FC(O)C(O)F] during these experiments at longer times, which appeared as a stable product in the presence of NO_x .

The results and experimental conditions of these experiments are summarised in Table 3, the mechanistic implications of these observations are discussed below.

The time profiles and product yields observed in the experiments with NO_x present, indicate that CF_2O , FC(O)C(O)F and CF_3CF_2OCOF are not depleted to any significant extent by reaction with OH radicals. This is consistent with the fact that perfluorinated, saturated organic trace gases do not react rapidly with OH. In the NO_x free experiments with 253.65 nm photolysis of $O_3/H_2/air$ as OH precursor, a lower yield of FC(O)C(O)F was observed and it decayed significantly during the measurement. As FC(O)C(O)F does absorb weakly at 254 nm, ¹⁶ some of this loss may be caused by photolysis. To examine whether other processes also play a role we stopped the photolysis after

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Summary of experimental conditions and results in the product experiments with CH₃ONO as OH precursor (with NO_x) Table 2

P/Torr	$[PEVE]_0^a$	$[CH_3ONO]_0^a$	$[NO]_0^{\ a}$	$[OH]_0^{\ b}$	$\Phi_{\mathrm{NO}_{\mathrm{x}}}\!(\mathrm{CF_2O})^c$	$\Phi_{\mathrm{NO}_x}\!(\mathrm{FC}(\mathrm{O})\mathrm{C}(\mathrm{O})\mathrm{F})^d$	$\Phi_{\mathrm{NO_x}}\!\!\left(\mathrm{C_2F_5OCFO}\right)^d$
750	3.7	2.2	0.0	3.9	1.21	0.38 ± 0.06	0.49 ± 0.06
750	1.2	4.5	0.0	3.5	1.52	0.38 ± 0.04	0.44 ± 0.02
750	1.5	4.5	2.9	0.3	1.67	0.48 ± 0.06	0.46 ± 0.04
750	2.8	3.3	0.0	3.2	1.38	0.45 ± 0.16	0.48 ± 0.16
50	3.6	2.2	0.0	3.9	1.78	0.69 ± 0.04	0.11 ± 0.02
750	2.0	3.2	0.0	0.7	1.45	0.45 ± 0.18	0.49 ± 0.18
750	1.4	3.1	1.7	0.5	1.67	0.47 ± 0.06	0.56 ± 0.02

^a The initial concentrations of PEVE (units of 10¹³ molecule cm⁻³), NO and CH₃ONO (units 10¹⁴ molecule cm⁻³) were determined from pressure measurements. ^b The OH concentration (units 10⁹ molecule cm⁻³) was determined from the initial decay of PEVE. ^c $\Phi_{NO_{c}}$ (CF₂O) is the value obtained at long reaction times when the yield is constant (see text for details). d Φ_{NO_o} (FC(O)C(O)F) and Φ_{NO_o} (C₂F₅OCFO) were obtained from plots of Δ product *versus* Δ PEVE as in Fig. 6. Uncertainties are 2σ .

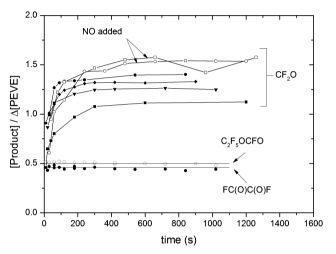


Fig. 7 Product yields from experiment with PEVE and CH₃ONO. The results of only two experiments are displayed for C₂F₅OCFO and FC(O)CFO to preserve clarity of presentation. The solid, horizontal lines represent the average yield from all experiments. The CF₂O yields represented by open symbols are from experiments in which NO was added.

generation of a significant amount of the products and left the mixture in the dark with $2-6 \times 10^{14}$ molecule cm⁻³ ozone for one hour with spectra recorded at regular intervals. Even in the dark, the concentration of FC(O)C(O)F decreased (with a decay constant of $2.5 \times 10^{-4} \text{ s}^{-1}$) while no significant loss of CF₂O or C₂F₅OCFO was observed. Potential causes of the dark loss of FC(O)C(O)F could be direct reaction with ozone in the gas phase, or on surfaces, the latter mediated by surface catalysed dissociation of ozone.¹⁷ The scatter in the yields in the experiments without NO_x most likely reflect variation in the removal rate and we therefore chose to report the product yield of FC(O)C(O)F as a lower limit with $\Phi(FC(O)C(O)F) > 0.36$. The average yields of C₂F₅OCFO and CF₂O are 0.47 ± 0.07 and 1.20 ± 0.20 , respectively.

Mechanism for PEVE photo-oxidation in the presence of NO

From the observations described above, we can develop a reaction scheme for the OH-initiated oxidation of PEVE in air in the presence of NO_x, which is summarised in Fig. 8.

The reaction of PEVE with OH radicals is initiated by addition of the OH radical to the double bond, which can in principle happen at both carbon atoms.

$$PEVE + OH \rightarrow C_2F_5OCFCF_2OH$$
 (R1a)

$$\rightarrow$$
 C₂C₅OCF(OH)CF₂ (R1b)

Computational studies on reaction of OH with PMVE18 and perfluoro propyl vinyl ether (C₃F₇OCF=CF₂, PPVE),¹⁵ indicate that addition to the terminal C-atom (R1a) dominates. The large experimental yield of FC(O)C(O)F is not commensurate with a significant contribution of (R1b) and we conclude that the OH-addition is predominantly at the terminal carbon atom (R1a). The adduct formed in (R1a) may either dissociate via C-O bond fission (R15) or react with oxygen at the neighbouring site giving a hydroxy-peroxy radical (R16).

$$C_2F_5OCFCF_2OH \rightarrow C_2F_5 + HOCF_2C(O)F$$
 (R15)

$$C_2F_5OCFCF_2OH + O_2 \rightarrow C_2F_5OCF(OO)CF_2OH$$
 (R16)

$$HOCF_2C(O)F \rightarrow FC(O)C(O)F + HF$$
 (R17)

The conversion of FC(O)CF₂OH to FC(O)C(O)F in (R17) is unlikely to be an elementary reaction, as discussed below.

Table 3 Experimental conditions and results of product study experiments using O₃/H₂ as OH precursor (without NO_x)

P/Torr	[PEVE] ₀ ^a	$[O_3]_0^a$	$[\mathrm{H_2}]_0^{~a}$	$[OH]_0^b$	$\Phi(\mathrm{CF_2O})^c$	$\Phi(FC(O)C(O)F)^d$	$\Phi(C_2F_5OCFO)^c$
600	2.4	1.3	2.0	4.3	1.37 ± 0.04	>0.36	0.49 ± 0.02
760	1.9	2.2	1.9	5.1	1.31 ± 0.02	> 0.34	0.49 ± 0.01
750	1.1	0.7	1.7	5.0	0.91 ± 0.05	> 0.30	0.44 ± 0.02

^a The initial concentrations of PEVE (units of 10¹³ molecule cm⁻³), O₃ (units 10¹⁴ molecule cm⁻³) and H₂ (units 10¹⁷ molecule cm⁻³) were determined from pressure measurements. ^b The OH concentration (units 10⁹ molecule cm⁻³) was determined from the initial decay of PEVE. $\Phi(FC(O)C(O)F)$ and $\Phi(C_2F_5OCFO)$ were obtained from plots of Δ product versus Δ PEVE, uncertainties are 2σ . $\Phi(FC(O)C(O)F)$ uses only data at short times and is a lower limit (see text for details).

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Fig. 8 Proposed mechanism for the OH-initiated photo-oxidation of PEVE in the presence of NO_x. Observed products are marked with black circles. The formation of CF2O and FC(O)C(O)F from their respective alcohols are assumed to be heterogeneous reaction and are marked with "Het"

The fate of the hydroxyl-peroxy radical formed in R16 depends on the experimental conditions. In the experiments with NO_r present, the radical will react with NO, forming an oxy-radical (R18) which may then dissociate to form either HOCF₂C(O)F and the C₂F₅O radical (R19a) or the observed product C₂F₅OC(O)F along with the CF₂OH radical (R19b).

$$C_2F_5OCF(OO)CF_2OH + NO \rightarrow C_2F_5OCF(O)CF_2OH + NO_2$$
(R18)

$$C_2F_5OCF(O)CF_2OH \rightarrow C_2F_5O + CF(O)CF_2OH$$
 (R19a)

$$\rightarrow$$
 C₂F₅OC(O)F + CF₂OH (R19b)

$$CF_2OH + O_2 \rightarrow CF_2O + HO_2$$
 (R20)

As HOCF2C(O)F may dissociate to FC(O)C(O)F and HF, both reactions fit the observations, but the calculations by Vereecken et al.18 on PMVE and PPVE indicate (R19b) to be the dominant pathway. The fate of the CF₂OH co-product of (R19b), is reaction with O₂ to form CF₂O.

As indicated above, the product yields were found to be dependent on the total pressure of air, with a substantial decrease in Φ_{NO} (C₂F₅OCFO) when going from 750 Torr to 50 Torr, accompanied by increases in Φ_{NO} (FC(O)C(O)F) and Φ_{NO} (CF₂O). These trends are consistent with the results of Mashino et al.⁷ who (for PMVE) found that the CF3OCFO yield was reduced from 0.53 \pm 0.04 at 700 Torr, to 0.08 \pm 0.02 at 10 Torr.

They attributed this pressure dependence to competition between HF elimination and collisional stabilization of the chemically activated PMVE-OH adduct. However, Vereecken et al. 18 calculated that the barrier of this reaction is too high for it to be of any significance. Instead, they suggested that (for PMVE) the OH-adduct would preferentially dissociate to CF₃ + CF(O)CF₂OH, with a barrier around 125 kJ mol⁻¹ below the entrance energy and proposed the equivalent mechanism for PPVE. 15 Our observations of pressure dependent product yields can thus be understood in terms of competition between dissociation of an activated OH-PEVE complex ((R15), most important at low pressures) and thermalisation/reaction of the complex with O_2 ((R16), most important at high pressures).

As addition of chlorine to double bonds is significantly less exothermic than addition of OH, the PEVE-Cl adduct is formed at lower energies. One may therefore expect a different product distribution if the conclusions drawn above are valid. This is indeed the case as only C₂F₅OCFO is observed, indicating that the PEVE-Cl adduct is formed below the barrier for dissociation to $C_2F_5 + CF(O)CF_2Cl$ and confirming that the product distribution is largely determined through the competition between dissociation and collisional stabilization of the initial PEVE-OH adduct.

The C₂F₅ radical formed (R15) in the dissociation of the C₂F₅OCFCF₂OH radical will react with oxygen to give the perfluoroethyl peroxy radical (C₂F₅O₂) which reacts with NO to form C₂F₅O ((R21) and (R22)). This radical is thermally unstable and will dissociate on a microsecond timescale to CF₃ and carbonyl fluoride (R23).^{19,20} The CF₃ radical reacts further with O2 and NO similar to the methyl radical giving another carbonyl fluoride molecule as final product ((R24), (R25) and (R14)).

$$C_2F_5 + O_2 + M \rightarrow C_2F_5O_2 + M$$
 (R21)

$$C_2F_5O_2 + NO \rightarrow C_2F_5O + NO_2$$
 (R22)

$$C_2F_5O \rightarrow CF_3 + CF_2O$$
 (R23)

$$CF_3 + O_2 \rightarrow CF_3OO$$
 (R24)

$$CF_3OO + NO \rightarrow CF_3O + NO_2$$
 (R25)

According to the reaction scheme ((R14)-(R25)) describing the subsequent fate of the initially formed OH-PEVE adduct, there are two main reaction routes with CF2O yields of one and two respectively, the total yield of CF₂O should approximately be equal to the FC(O)C(O)F yield plus one, which is consistent with the experimental results.

Fig. 6 and 7 indicate that the formation of CF₂O is delayed, whereas FC(O)C(O)F and C₂F₅OCFO are formed promptly. This is consistent with CF₂O being formed from slow dissociation of CF₃OH (R26).

$$CF_3OH \rightarrow CF_2O + HF$$
 (R26)

Perfluorinated primary alcohols are known to dissociate to the corresponding perfluoroacyl halide and HF.21,22 The mechanism for formation of FC(O)C(O)F in the photo-oxidation of a **Paper**

fluorinated vinyl ether has been discussed by Mashino *et al.*⁷ who suggested direct HF loss from CFOCF₂OH in the gas phase. Later, Vereecken *et al.*¹⁸ calculated the barrier of this reaction to be 190 kJ mol⁻¹ indicating that the reaction is too slow to be relevant at ambient temperatures and that FC(O)C(O)F is likely formed in a different process. The dissociation of CF₃OH to CF₂O + HF²³ has a barrier of similar magnitude to that of CFOCF₂OH, and it is reported to take place rapidly in chamber experiments with a rate depending on the history of the reaction chamber, indicating that a heterogeneous reaction is taking place.²⁴ It has also been suggested that the dissociation of CF₃OH is catalysed in the gas phase by water vapour,²³ HF,²⁵ CF₃OH, or radicals.^{26–28}

The difference in the observed CF_2O yields upon changing the NO concentration can thus be rationalized as follows: When there is enough NO available for scavenging CF_3O , CF_2O is formed directly in the gas phase via (R14). At lower NO concentrations, a fraction of the CF_3O radicals react to CF_3OH which will then convert to CF_2O and HF (R26). The time dependence of CF_2O formation is thus controlled by the rate of diffusion of CF_3OH to and from surfaces and explains the induction period of minutes. Note that neither FNO nor CF_3OH were observed, and HF only in low concentrations. This reflects their high reactivity of these molecules on the quartz and metal surfaces inside the reactor.

3.7 Mechanism for PEVE photo-oxidation in the absence of NO

In the experiments with ozone and H_2 , there is no NO to scavenge either the CF_3O radicals or the peroxy radicals. The peroxy radicals may react with HO_2 to form either the hydroperoxide (R27) or with ozone to generate the oxy radical and oxygen (R28).

$$C_2F_5OCF(OO)CF_2OH + HO_2 \rightarrow C_2F_5OCF(OOH)CF_2OH + O_2$$
(R27)

$$C_2F_5OCF(OO)CF_2OH + O_3 \rightarrow C_2F_5OCF(O)CF_2OH + 2O_2 (R28)$$

No evidence for formation of the hydroperoxide was found in the IR spectra and we assume that if it is formed it will either dissociate to the oxy radical and the OH radical (R29), be photolysed or be lost on surfaces.

$$C_2F_5OCF(OOH)CF_2OH \rightarrow C_2F_5OCF(O)CF_2OH + OH$$
 (R29)

The CF₃O radicals react with hydrogen to give CF₃OH and a hydrogen atom which rapidly react with O₂ to form an HO₂ radical

$$CF_3O + H_2 \rightarrow CF_3OH + H$$
 (R30)

$$H + O_2 + M \rightarrow HO_2 + M \tag{R8}$$

 CF_3OH will decompose to CF_2O and HF as described above, but as this most likely happens on surfaces, some of the resulting CF_2O may then hydrolyse to CO and HF, explaining the lower yield of CF_2O in the ozone experiments.

3.8 Theoretical calculations on the fate of CF_3OH and $HOCF_2C(O)F$

As discussed above, the formation of CF₂O from CF₃OH and FC(O)C(O)F from HOCF₂C(O)F, has been shown to be driven by heterogeneous reactions. While a detailed analysis of such reactions is outside the scope of the present paper, it is worthwhile to examine the intrinsic difference in reactivity between CF₃OH and HOCF₂C(O)F. We therefore examined the reactants and transition states (TS) for a set of representative reactions at the M06-2X/aug-cc-pVTZ level of theory. ^{29,30} The relative energies were refined by single point CCSD(T)-F12a/aug-cc-pVDZ calculations. ^{31,32} All calculations were performed using the Gaussian-09 and Molpro 2015.1 program suites. ^{33,34}

For both CF₃OH and HOCF₂C(O)F, reaction proceeds by conversion of the -CF₂OH moiety to a carbonyl group, with HF as a co-product. Literature data indicates that direct HF elimination has a large energy barrier, >160 kJ mol⁻¹, but that the reaction can be catalysed by many co-reactants. 23,25-28 The experimental data presented in this work (see Fig. 6 and 7) suggest that CF₃OH eliminates HF significantly slower than HOCF₂C(O)F. To study this dependence on the molecular structure, we have calculated the rate-limiting TS for HF formation both for the un-catalysed reaction, as for catalysed processes, where we examined CF₃OH, CH₃OH, H₂O, HF, HC(O)OH, HO₂, and OH as potential catalysts (see Table 4). In agreement with literature data, we find that the barrier for the un-catalysed reaction is too high to allow a significant contribution at room temperature, but that the various catalysts reduce the barrier heights, by up to 160 kJ mol⁻¹. For nearly all catalysts, the barrier height for the HOCF₂C(O)F substrate is lower than for the CF₃OH compound, up to 13 kJ mol⁻¹. The lower TS energies afford a faster rate coefficient by over an order of magnitude, affirming the experimentally observed difference in conversion rates for HOCF2C(O)F and CF₃OH. The difference in the reactivity trend is expected to remain valid for nearly all reactions including heterogeneous reactions, as the underlying cause is the difference in reaction enthalpy imposed by the -C(O)F moiety in HOCF₂C(O)F, which is absent in CF₃OH (see Table 4). Of the gas phase co-reactants considered, we find that HO2 is the most efficient catalyst, pairing

Table 4 Barrier heights^a (kJ mol^{-1}) for conversion of the $-CF_2OH$ moiety to -CFO + HF, with a series of catalysts. The pre- and post-reaction complexes are not indicated

Catalyst	CF ₃ OH	HOCF ₂ C(O)F
(None)	186	186
ČF₃OĤ	97	93
OH	93	86
HF	89	86
H_2O	75	64
CH ₃ OH	52	39
HO_2	45	38
HC(O)OH	28	22
Product energy	$CF_2O + HF: 30$	CF(O)C(O)F + HF: 24

 $[^]a$ Calculated at the ZPE-corrected CCSD(T)-F12/aug-cc-pVDZ//M06-2X/aug-cc-pVTZ level of theory.

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a radical reaction site with an -OO- molecular span that reduces the TS ring strain compared to the shorter -O- or -F- moieties in the other gas phase reactants. Intriguingly, we also find that CH₃OH is a more efficient catalyst than CF₃OH. This is related to the electron density on the -OH moiety; this aspect is not investigated in detail here. Larger catalysts consisting of complexes (e.g. HO2 with water molecules) have been shown to be even more effective, but we do not expect significant concentrations of such gas-phase complexes under our reaction conditions. Formic acid, HC(O)OH, expected to be present on the reactor walls, affords the most drastic reduction in barrier height of the catalysts considered, essentially removing any energetic barrier beyond the intrinsic reaction endoergicity. The catalysis mechanism involves donating an H-atom from the acidic hydroxy group and receiving an H-atom with the carbonyl group while shifting the double bond, thus combining a mobile H-atom with a large -O-C=Omolecular span reducing TS ring strain, and strong H-bonding in the pre- and post-reaction complexes; this mechanism and its concomitant low energy barriers are likely to be applicable to all oxoacids including HNO3 and H2SO4. An in-depth study of heterogeneous reactions, be it with chemical contaminants or exposed metal surfaces, is beyond the scope of this manuscript.

While we cannot conclude that the reactions in Table 4 are those responsible for our observations of CF_2O and FC(O)C(O)F formation, the selected catalysts are all expected to be present in our system, and combined allow for very rapid conversion of the fluoro-hydroxy moiety to a carbonyl moiety + HF, where especially low-volatility, surface-bound catalysts such as carboxylic acids or aqueous complexes have high catalytic activity. The –CFO moiety in $HOCF_2C(O)F$ lends itself to additional interactions (e.g. H-bonding), which could again enhance the interaction with a reactive surface relative to CF_3OH .

Finally, we examined the reaction enthalpy of some other processes that could lead to carbonyl compounds indirectly, *e.g.* by formation of an –C*FOH radical moiety that can react with O₂ to form –CFO + HO₂. However, the available co-reactants NO, NO₂, FNO and OH do not afford accessible pathways for CF₃OH forming *CF₂OH, with endoergicities exceeding 175 kJ mol⁻¹ in all cases. Note that such reactions would also not readily yield HF, observed as co-product in the reactions of fluorinated esters.

3.9 Persistence of the products of PEVE photo-oxidation in the atmosphere

As for other fluorinated VOCs the atmospheric photo-oxidation products of PEVE are fluorinated carbonyl compounds. Unlike their non-fluorinated analogues, fluorinated carbonyl compounds react too slowly with OH radicals for this reaction to constitute a significant atmospheric sink and their lifetime is limited by uptake on water droplets followed by hydrolysis.³⁵

Glyoxal, the non-fluorinated analogue of FC(O)C(O)F, has a very short atmospheric lifetime with respect to photolysis.³⁶ In contrast, the first electronic excitation in FC(O)C(O)F appears at 334 nm and is very weak.¹⁶ Photolysis does therefore not constitute a significant sink of FC(O)C(O)F in the atmosphere.

To further investigate the fate of the C_2F_5OCFO , CF_2O and FC(O)C(O)F products with respect to loss to aqueous atmospheric systems, an experiment was performed in which, following PEVE oxidation, the contents of the photochemical reactor was passed (using a chemically inert membrane pump at a flow of 5 SLM) through a wash bottle (volume = 300 cm³) containing distilled water. The gas flow did not bubble through the volume of the water, but was directed \emph{via} a glass tube (velocity 4 m s $^{-1}$ in the tube) towards its surface (area 50 cm²). The residence time of the gas mixture in the head-space of the wash-bottle was ≈ 2.4 s and the flow/geometry induced turbulent flow conditions in the head-space should ensure efficient gas–liquid contact.

Infrared spectra were recorded at 10 min intervals directly after the initial, gas-phase degradation of PEVE, then over a 60 minute period in which the contents of the cell were circulated through the membrane pump without exposure to water and then a further sixty minutes in which the water surface was available for reaction. During the first hour (no interaction with water), no significant loss of any products was observed, indicating no significant losses in the membrane pump. In contrast, the concentration of all three products decreased significantly upon contact with the water surface. Whereas CF₂O was almost completely removed after ≈240 s (decay constant $6 \times 10^{-4} \text{ s}^{-1}$), the decay of C_2F_5OCFO (4 × 10⁻⁵ s⁻¹) was an order of magnitude slower. FC(O)C(O)F decayed on a similar time scale to CF₂O, but as the only absorptions not overlapping with the saturated water lines are very weak, we were unable to determine a decay constant.

The lifetime with respect to hydrolysis of trace gases in contact with aqueous surfaces is determined by their Henry's law solubility (H) and hydrolysis rate coefficient $(k_{H,O})$ as well as their rate of transport to the surface and diffusion in the bulk. George et al.³⁷ studied the uptake of CF₂O by aqueous surfaces and report $H(k_{H,O})^{\frac{1}{2}} = 350 \text{ M atm}^{-1} \text{ s}^{-1/2}$ at 273 K. A substantially lower values (at 2 78 K) of 4.3 M atm $^{-1}$ s $^{-1/2}$ was reported by de Bruyn et al. 38 Subsequently, Kanakidou et al. 39 modelled the atmospheric fate of CF₂O based on the data of George et al.³⁷ and reported a lifetime with respect to in-cloud losses of the order of one week, allowing for a considerable uncertainty in the Henry law coefficients and hydrolysis rates. Given that the rate of turbulent transport to the water surface is expected to be identical for all the trace gases we can reasonably assume that the similar removal rates of FC(O)C(O)F and CF2O are due to similar values for $H(k_{H_2O})^{\frac{1}{2}}$ and that they have a similar timescales for removal to aqueous surfaces in the atmosphere. In contrast, the much lower rate of removal of C₂F₅OCFO compared to CF_2O implies a large (factor of ≈ 10) reduction in the $H(k_{H,O})^{\frac{1}{2}}$ term and thus potentially longer atmospheric lifetime.

The hydrolysis of FC(O)C(O)F is likely to lead to formation of HF and CO, whereas the potential hydrolysis products of C_2F_5OCFO are HF, CO and $CF_3C(O)OH$. As both HF and $CF_3C(O)OH$ are highly soluble it is expected that they would remain undetected in the gas-phase but remain in the bulk water sample in our experiments.

Our crude experiments to investigate the relative loss rates of CF₂O, C₂F₅OCFO and FC(O)C(O)F to aqueous surfaces can

only give a broad indication of the fate of these trace gases in the atmosphere and dedicated experiments to investigate the atmospheric fate of C₂F₅OCFO and FC(O)C(O)F with respect to gas-phase and heterogeneous processes are necessary to fully characterise the environmental impact of PEVE.

4 Conclusions

The main products formed in the OH-initiated photo-oxidation of PEVE are C_2F_5OCFO , FC(O)C(O)F and CF_2O with respective yields (at 1 bar and in the presence of NO_x) of 0.50 ± 0.07 , 0.46 ± 0.07 and 1.50 ± 0.22 . Variation of product yields with pressure and concentration of NO enabled a detailed mechanism to be derived, in which product branching is controlled by competition between dissociation and reaction (with O_2) of the initially formed reaction complex. In the environment, CF_2O formed from PEVE degradation will hydrolyse, forming HF and CO. The fate of C_2F_5OCFO and FC(O)C(O)F in the atmosphere remains unknown, though heterogeneous processes are likely to be significant. In addition to the mechanistic studies, we also derived a rate coefficient $(2.8 \pm 0.3 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$ which is in good agreement with the values from our previous kinetic study.

Conflicts of interest

There are no conflicts to declare.

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