



Cite this: *Environ. Sci.: Atmos.*, 2023, 3, 855

A computer-based solution to the oxidation kinetics of fluorinated and oxygenated volatile organic compounds†

Luis P. Viegas *^a and Frank Jensen ^b

The OH radical is the most powerful atmospheric oxidant, being responsible for the chemical breakdown of many pollutants released into the troposphere, including saturated volatile organic compounds (VOCs). Numerous of these VOCs are strong greenhouse gases, including hydrofluorocarbons, whose industrial emissions to the environment have grown significantly over time and are gradually being phased out. The calculation of OH-initiated oxidation reaction rate constants represents an approach to determine the atmospheric impact of fluorine-containing oxygenated VOCs, which have been regarded as a good substitute for hydrofluorocarbons. Such calculations are challenging, especially when the VOCs are conformationally flexible. We have recently developed a cost-effective computational protocol for such systems based on density functional calculations and on the coupling of multiconformer transition state theory with constrained transition state randomization. The protocol has, however, only been calibrated against a small number of experiments, practically all of which involved OVOCs from the ether family. In this study, we extend the scope by applying the protocol to five families of OVOCs (C₃-fluorine containing alcohols, aldehydes, esters, ethers and ketones) and compare the calculated reaction rates with the recommended experimental values. Our calculated rate coefficients, on average, agree with the recommended experimental values to within a factor of two, with the maximum deviation being a factor of 2.8. These results validate the use of the protocol for computational tropospheric degradation studies of conformationally flexible OVOCs with an unprecedented accuracy.

Received 29th November 2022
Accepted 13th March 2023

DOI: 10.1039/d2ea00164k

rsc.li/esatmospheres

Environmental significance

The impact of hydrofluorocarbons (HFCs) on global warming can be hundreds to thousands of times greater than that of CO₂. With the rapid rise of HFC emissions and their projected catastrophic effects on the atmosphere and climate, the Kigali Amendment of 2016 seeks to drastically reduce the global production and consumption of HFCs, thus contributing to the Paris agreement target of limiting the global temperature rise below 2 degrees Celsius. For this reason, new ideas and green technologies are necessary to achieve the Kigali goals and mitigate climate change effectively. Based on the results of the present study, we demonstrate that our cost-effective computational protocol is capable of predicting with unprecedented accuracy the rate coefficients of the OH-initiated reactions of several families of saturated organic compounds that could replace HFCs. Consequently, our computational approach has the potential not only to deliver the required fundamental knowledge but also to actively take part in this critical scientific endeavour.

1 Introduction

In the wake of the realization that chlorofluorocarbon (CFC) compounds were part of the ozone depletion problem and were causing global warming,^{1–4} theoretical and experimental efforts to find environmentally friendly CFC replacements have expanded dramatically. Almost 200 nations adopted the Kigali Amendment to the Montreal Protocol⁵ in October 2016, making

this subject even more relevant. Ratifying this protocol makes countries stop using and producing hydrofluorocarbons (HFCs), which makes it important to find environmentally friendly alternatives, since these second-generation replacements have a high global warming potential.⁶ One of the most important parameters defining the global warming potential of a compound is its atmospheric lifetime.⁷ The atmospheric lifetime is often decreased by introducing oxygen-containing functional groups, and fluorine-containing oxygenated volatile organic compounds (OVOCs) are candidates for third-generation replacements in industrial and commercial applications.⁸ Relative to their non-oxygenated fluorinated counterparts, the oxygen functional groups promote chemical activation by hydrogen-atom abstraction at distant sites from

^aCoimbra Chemistry Centre-Institute of Molecular Sciences (CQC-IMS), Department of Chemistry, University of Coimbra, Coimbra 3004-535, Portugal. E-mail: lpviegas@ci.uc.pt

^bChemistry Department, Aarhus University, DK-8000 Aarhus, Denmark

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d2ea00164k>



the substituent group, and thus increase the atmospheric reactivity by OH radicals.^{9–11} There are a variety of applications⁸ for fluorinated alcohols, aldehydes, esters, ethers and ketones in industry and they can potentially replace HFCs, as mandated by the Kigali Amendment to the Montreal Protocol.

The two main degradation pathways associated with OVOCs are photolysis and reaction with OH radicals, with the former being relevant only for certain families of OVOCs, such as aldehydes and ketones:^{9,11,12} these two families of compounds contain a carbonyl group which absorbs radiation in the ultra-violet region around 280 nm due to its $n \rightarrow \pi^*$ transition.¹³ The global lifetime^{7,14} (also referred to as “lifetime”, “total lifetime” or “turnover time”¹⁵) of OVOCs based on these two degradation pathways can be estimated as follows:

$$\frac{1}{\tau_{\text{OVOC}}^{\text{global}}} = \frac{1}{\tau_{\text{OVOC}}^{\text{photo}}} + \frac{1}{\tau_{\text{OVOC}}^{\text{OH}}}, \quad (1)$$

where $\tau_{\text{OVOC}}^{\text{photo}}$ represents the OVOC atmospheric lifetime resulting from the photolytic process (which depends on the altitude^{9,11}) and $\tau_{\text{OVOC}}^{\text{OH}}$ is the atmospheric lifetime associated with a pseudo-first-order loss process with a corresponding $k_{\text{OH}}[\text{OH}]$ rate constant. Interestingly, fluorination of aldehydes and ketones leads to a bathochromic shift in the UV absorption bands of the carbonyl groups,^{13,16,17} enabling a better overlap with the actinic flux available at lower altitudes and decreasing the value of the $\tau_{\text{OVOC}}^{\text{photo}}$ atmospheric lifetime to a range between a few days and a few weeks.^{13,16–18} Consequently, and in the context of the compounds studied in this research, discarding $\tau_{\text{OVOC}}^{\text{photo}}$ from the calculation of the $\tau_{\text{OVOC}}^{\text{global}}$ global lifetimes should, in general, be a good approximation, since the $\tau_{\text{OVOC}}^{\text{photo}} \gg \tau_{\text{OVOC}}^{\text{OH}}$ relation between lifetimes will be frequent. The exception are fluorinated aldehydes and ketones, for which the much lower discrepancy between the $\tau_{\text{OVOC}}^{\text{photo}}$ and $\tau_{\text{OVOC}}^{\text{OH}}$ lifetimes can lead to large errors in $\tau_{\text{OVOC}}^{\text{global}}$ if the first term in the right-hand-side of eqn (1) is discarded. In the present case, we will focus on the bimolecular reactions between OVOCs and the OH radical (commonly known as the atmosphere’s “detergent”), from which the atmospheric lifetimes can be obtained by

$$\tau_{\text{OVOC}}^{\text{OH}} = \frac{1}{k_{\text{OH}}[\text{OH}]}. \quad (2)$$

In these bimolecular degradation reactions, the OH radical typically abstracts a hydrogen atom from the OVOC^{9,11,19–21}



producing the corresponding OVOC radical and a water molecule. A protocol for accurately calculating such k_{OH} rate constants is useful for predicting and understanding reactivity trends, especially for species that cannot easily be observed experimentally. The use of accurate theoretical rate constants can thus provide new insights and complement information from structure–activity relationships (SAR) derived from experimental information.²¹ It is thus possible for theoretical methods to play a crucial role in accelerating the discovery of new, greener^{22,23} compounds with a low global warming

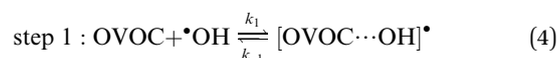
potential. Providing solutions to reduce human impact on the climate is of utmost importance within the framework of the Kigali Amendment to the Montreal Protocol.

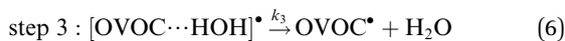
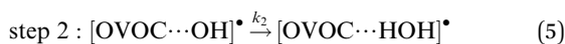
There are, however, some challenges associated with obtaining accurate rate constants by computational methods. For reactions where the dynamical bottleneck is difficult to locate, robust approaches^{24–29} based on variational transition state theory (VTST) are necessary. However, these become unmanageable for molecules exhibiting rich conformational complexity, such as in molecules of moderate/large size. Fortunately, for reactions involving the OH radical and fluorine-containing saturated oxygenated VOCs such as the ones explored in this work, the use of VTST is not necessary, since the transition states are well approximated by following the conventional TST approach of placing the dividing surfaces that separate reactants from products at the saddle-points of the potential energy surfaces (PESs), instead of having to variationally optimize them. Within such a framework, we have recently developed,^{30,31} tested^{32–36} and improved³⁷ a computational protocol for a cost-effective calculation of bimolecular rate constants for the reaction between $\bullet\text{OH}$ and VOCs of moderate/large size. This protocol is based on multiconformer transition state theory (MC-TST)^{38–44} and a method for performing transition state sampling called constrained transition state randomization (CTSR).³⁷ The MC-TST/CTSR protocol has provided promising results for fluorinated ethers, but it clearly needs to be tested for a wider range of fluorinated OVOC compounds before it can be routinely applied. In the present case we extend the calibration study to compounds belonging to the hydrofluoroalcohol, hydrofluoroaldehyde, hydrofluoroester, hydrofluoroether and hydrofluoroketone families. The study aims to assess the effectiveness of the MC-TST/CTSR protocol in obtaining accurate rate constants for the reaction between the OH radical and the five fluorinated OVOC families listed above. We have considered two examples of C₃-fluorinated OVOCs for each of the five families of compounds, for a total of 10 systems. Our main focus will be a comparison of the MC-TST/CTSR predicted rate constants with the recommended experimental values at 298.15 K, but where experimental information is available, we also compare branching ratios and Arrhenius parameters.

2 Theory and methods

2.1 Theoretical background

A typical example of a radical–molecule reaction of atmospheric interest⁴⁵ is the reaction between OVOCs and $\bullet\text{OH}$,^{9,19,20} where both reactant and product channels exhibit loose complexes that are connected by hydrogen-abstraction transition states (TSS). The reaction profile features three steps: (1) formation of a prereaction complex (PRC) from the isolated OVOC + OH radical, (2) formation of a product complex by hydrogen abstraction from the OVOC, and (3) formation of the separated products (OVOC radical and water) from the product complex (PC):





If a constant prereaction complex concentration is assumed (steady-state assumption,⁴⁶ with $k_{-1} + k_2 \gg k_1$, indicating^{46,47} that the removal process of the PRC is much faster than its formation and leading to $d[\text{PRC}]/dt = 0$), the effective rate constant for reaction (3) is given by^{48–51}

$$k_{\text{OH}} = \frac{k_1 k_2}{k_{-1} + k_2} \quad (7)$$

It is generally found that k_{-1} reverse rate constants for step 1 in tropospheric reactions are much larger than k_2 rate constants for the irreversible unimolecular step 2 (the conditions for which $k_{-1} \gg k_2$ are discussed in detail elsewhere^{30,31,47–51}), and $k_{\text{OH}} = k_1 k_2 / k_{-1}$ is therefore the overall rate constant for the total reaction. The underlying reason is that the system is fully equilibrated and thermalized at the high-pressure limit (equilibrium-state model is applicable,⁴⁷ where $k_1, k_{-1} \gg k_2$), and thus the Boltzmann equilibrium distribution can be used to describe the reactants, the PRC, and the TS in step 2. In order to check whether the steady- and equilibrium-state conditions are met, particularly if the hydrogen-abstraction TSs are energetically below the reactants, the Gibbs free energies associated with the reaction path should be calculated and their relative values verified before carrying out the calculations. For tropospheric reactions at 298.15 K, a typical Gibbs free energy profile along the reaction coordinate usually supports the following inequalities:⁴⁷ $\Delta G_{\text{R}} < \Delta G_{\text{PRC}} < \Delta G_{\text{TS}}$ and $\Delta G_{\text{R}} \gg \Delta G_{\text{PC}}$, where G_{R} is the Gibbs free energy of the reactants, against which the remaining free energies are calculated ($\Delta G_{\text{R}} = 0 \text{ kcal mol}^{-1}$). For example, for the 10 reactions studied in this investigation the following average values are obtained: $\overline{\Delta G_{\text{PRC}}} = 4.3 \text{ kcal mol}^{-1}$, $\overline{\Delta G_{\text{TS}}} = 10.4 \text{ kcal mol}^{-1}$ and $\overline{\Delta G_{\text{PC}}} = -19.9 \text{ kcal mol}^{-1}$. From these values we infer that both the steady-state and equilibrium-state assumptions are satisfied⁴⁷ for the reactions under study. We can consequently calculate k_1/k_{-1} as the K_{eq} thermal equilibrium constant for the fast pre-equilibrium between the isolated reactants and the PRC, which can be obtained using all of the usual statistical thermodynamic methods.^{48–51} We can calculate k_2 by resorting to thermal TST and simplify the TST-based rate coefficient into $k_{\text{OH}} = K_{\text{eq}} k_2$. We will utilize this approach, which has been (and still is) widely used to describe reactions between OVOCs and $\cdot\text{OH}$ in the past. Within the framework of MC-TST, this approach offers the advantage of avoiding the need to consider the sampling of PRCs, as the sum of the Boltzmann averaged partition functions of the PRC conformers cancel out in the $K_{\text{eq}} k_2$ product, with the same occurring to the energy values present in both exponential factors. Such a simplification is very important in terms of the applicability of the computational protocol, since the sampling procedures for prereaction complexes involving flexible VOCs and $\cdot\text{OH}$ can be very complex.⁴³

Our latest version of the MC-TST/CTSR protocol^{31,35,37} incorporates the approach used by Petit and Harvey,⁴⁰ where both the numerator and denominator of eqn (7) are multiplied by K_{eq}

$$k_{\text{OH}} = \frac{k_1 k_2 K_{\text{eq}}}{k_{-1} K_{\text{eq}} + k_2 K_{\text{eq}}} = \frac{k_1 k_{\text{OH}}^{\text{MC-TST}}}{k_{-1} + k_{\text{OH}}^{\text{MC-TST}}} \equiv k_{\text{OH}}(\text{calc}) \quad (8)$$

Here k_1 is the rate coefficient for the formation of the pre-reaction complex and $k_{\text{OH}}^{\text{MC-TST}}$ is identified as the calculated MC-TST rate constants. The k_1 rate constant is unique for each specific reaction, but since it is almost independent of the species reacting with the OH radical, we used⁴⁰ a value of k_1 for collision with acetone of ($k_1 = (2.67 \pm 0.15) \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$) as determined by experimental rate constants^{52–55} for quenching of vibrationally excited $\cdot\text{OH}$.

The branching ratio concerning hydrogen abstraction at a generic $-\text{RH}$ abstraction site will be calculated according to

$$I_{\text{RH}} = \frac{k_{\text{OH}}^{\text{RH}}(\text{calc})}{k_{\text{OH}}(\text{calc})} \quad (9)$$

where $k_{\text{OH}}^{\text{RH}}(\text{calc})$ will be calculated using only TS conformers involving the attack of the OH radical to the specific RH group.

2.2 Computational methods

The rate coefficient in our MC-TST/CTSR protocol is calculated from the following expression,^{30,31,37} which stems from the derivation presented in the ESI† of ref. 42:

$$k_{\text{OH}}^{\text{MC-TST}} = \kappa(T) \frac{k_{\text{B}} T}{n_0 h Q_{\text{OH}}} \frac{\sum_i^M \alpha_i \omega_{\text{TS}_i} Q_{\text{TS}_i}}{\sum_j^N \alpha_j \omega_{\text{OVOC}_j} Q_{\text{OVOC}_j}} e^{-V^\ddagger/k_{\text{B}} T} \quad (10)$$

Here, N is the number of OVOC reactant conformers, while M represents the number of associated TS conformers obtained with the CTSR approach. The Q 's are total partition functions for $\cdot\text{OH}$, the i th OVOC reactant and j th TS, calculated as $Q = q_{\text{t}} q_{\text{e}} q_{\text{v}} q_{\text{r}}$ under the rigid rotor, harmonic oscillator approximation. Each OVOC or TS conformer Q is associated with a Boltzmann weight factor ω_{OVOC_j} or ω_{TS_i} , which depends on E_{OVOC_j} and E_{TS_i} zero-point corrected relative energies of conformations OVOC_j and TS_i , respectively:

$$\omega_{\text{OVOC}_j} = e^{-(E_{\text{OVOC}_j} - E_{\text{OVOC}_0})/k_{\text{B}} T} = e^{-\Delta E_{\text{OVOC}_j}/k_{\text{B}} T} \quad (11)$$

$$\omega_{\text{TS}_i} = e^{-(E_{\text{TS}_i} - E_{\text{TS}_0})/k_{\text{B}} T} = e^{-\Delta E_{\text{TS}_i}/k_{\text{B}} T} \quad (12)$$

The $\sigma_{\text{rot,OVOC}_j}$, $\sigma_{\text{rot,TS}_i}$, α_j and α_i are the rotational symmetry numbers and reaction path degeneracy parameters of conformations OVOC_j and TS_i , respectively. The quantity V^\ddagger is calculated^{30,39,41,42} as $V^\ddagger = E_{\text{TS}_0} - (E_{\text{OVOC}_0} + E_{\text{OH}})$ where the subscript 0 indicates the lowest energy conformer. Also, $n_0 = p/k_{\text{B}} T$ is the Loschmidt number at a temperature of 298.15 K and a pressure of one atmosphere and k_{B} and h are the Boltzmann and Planck's constants, respectively. Finally, $\kappa(T)$ represents the tunneling correction which is frequently calculated at a very low computational cost^{30,41,42} from the asymmetric Eckart expression,⁵⁶ where the forward (V_1) and reverse (V_2) barrier heights are



obtained by performing intrinsic reaction coordinate calculations from TS_0 and optimizing the respective PRC and PC at the endpoints. This approach, which we have been using in this MC-TST protocol,³¹ is often able to reproduce expensive and robust multidimensional transmission coefficients, such as the small curvature tunneling correction.⁵⁷ This is mainly due to favorable error cancellation effects in the Eckart approximation, whereby the underestimation of the tunneling probability (neglect of corner-cutting^{58–60}) is compensated by an overestimation of the tunneling probability (rigidity of the asymmetrical Eckart function produces a potential energy barrier which is too narrow^{59–63}). In the case of lower temperatures, however, the overestimation of the Eckart tunneling probability may be too drastic, resulting in an overestimation of $\kappa(T)$ and of the rate coefficient itself. As a more robust but still affordable alternative to the Eckart model, we have performed extra calculations of $\kappa(T)$ through the use of semiclassical TST (SCTST).^{64–67} In particular, we have used an improved one-dimensional model⁶⁸ (iSCTST) that besides including the anharmonicity of the reaction mode it also correctly includes the energetic information about the PRC and PC, thus allowing a qualitatively correct representation of deep tunneling which was absent in the original formulation of SCTST. For a molecule with N atoms and with $F = 3N - 6$ vibrational degrees of freedom, the x_{kl} anharmonic constant for the reaction mode in this one-dimensional model can be obtained by recognizing that the transition state frequency for the reaction coordinate (usually taken as mode F) is imaginary ($\omega_F = i\bar{\omega}_F$) and that the reaction mode is assumed not to be coupled with the remaining vibrational (spectator) modes in the TS, with the latter being treated harmonically. Considering these assumptions, one can obtain the x_{FF} value for the anharmonic constant of the reaction mode as⁶⁶

$$x_{FF} = -\frac{\hbar^2}{16\bar{\omega}_F^2} \left(f_{FFFF} + \frac{5}{3} \frac{f_{FFF}^2}{\bar{\omega}_F^2} \right). \quad (13)$$

Similarly, a barrier height correction parameter (G_0) should be included in the calculation of V_1 and V_2 .⁶⁹

$$G_0 = -\frac{\hbar^2}{64\bar{\omega}_F^2} \left(f_{FFFF} + \frac{7}{9} \frac{f_{FFF}^2}{\bar{\omega}_F^2} \right), \quad (14)$$

where f_{FFF} and f_{FFFF} are the third and fourth order derivatives of the potential with respect to the reaction mode coordinate. In order to apply Wagner's model,⁶⁸ one also needs to calculate the D parameter

$$D = -\frac{(\hbar\bar{\omega}_F)^2}{4x_{FF}} \quad (15)$$

and to save the value of the reduced mass associated with the motion along the reaction path at the saddle point. Besides $\bar{\omega}_F$, V_1 and V_2 (which are also required by the asymmetric Eckart model), Wagner's approach then requires the calculation of f_{FFF} and f_{FFFF} in order to obtain x_{FF} , G_0 and D . In SCTST, the anharmonic data is calculated through second-order vibrational perturbation theory (VPT2), which can be routinely⁷⁰

employed in Gaussian⁷¹ for extraction of f_{FFF} and f_{FFFF} . Although we have performed full VPT2 calculations for each TS_0 in Gaussian, the data required for a one-dimensional case can be obtained with less computational effort with just a few single point energy calculations.^{72,73}

To determine the remaining parameters of eqn (10), electronic structure calculations were performed with the GAMESS package.⁷⁴ The calculations were carried out using our reformulated protocol,³¹ which includes optimizing minima and TSs at the M08-HX/pcseg-2 level,^{75,76} with the value of V^\ddagger being refined with single-point energy calculations at the M08-HX/pcseg-2 level.

A crucial issue in solving eqn (10) is related to the reactant and TS conformer sampling procedure. The reactant conformers are calculated *via* the open source CONFAB,⁷⁷ the best performing freely available knowledge-based conformer generation tool.⁷⁸ However, and because CONFAB is based on force fields, there may be occasions when some low- and high-energy conformers⁷⁹ may be overlooked. Such a problem could be circumvented by targeting a near-complete description of the rotameric space with more robust and computationally demanding screening methods,^{42,43,80,81} but an increase in molecular size quickly renders this solution computationally ineffective, which is the opposite of what this computational protocol tries to achieve.

It should also be added that the CTSR procedure does not guarantee that the lowest energy TS is found. Nevertheless, the probability of doing so can be augmented by increasing the number of generated geometries and/or by replacing the intermediate PM3 step by optimizations at higher levels of theory. An estimation of the success rate of the CTSR sampling procedure would require a benchmark study on a chosen set of molecules, which would imply a very thorough and extensive mapping of the PESs associated to such OH abstraction reactions. This is an extremely difficult and laborious task which, to our knowledge, has not yet been done. However, we can state that (1) we are not aware of theoretical atmospheric studies concerning OH-initiated oxidation reactions that report such a high number of TSs in molecules having a similar size to the ones studied in this work and (2) the CTSR procedure is a considerable improvement³⁷ to the simplistic method that we were using in previous versions of the MC-TST protocol.^{30,31}

Naturally, one way to evaluate the impact of these approximations is to verify the quality of the rate constants calculated by the MC-TST/CTSR protocol, which can be considered an improved version of our initial protocol.^{30,31} The CTSR approach is still relatively new³⁷ and only two MC-TST/CTSR calculated rate constants have been compared with experimental results, both agreeing with the experimental rate coefficient^{36,37} within a factor of two. In addition, it is interesting to note, since error cancellation is an important and sometimes unpredictable factor in TST calculations,^{40,42,82–84} that the MC-TST/CTSR rate constant of the reaction between 'OH and 2-butanethiol³⁷ considering a full description of the reactant conformers is actually farther from the recommended experimental value than the MC-TST/CTSR rate constant based on the CONFAB approach. Considering the fact that it is difficult to predict the



variability in error cancellation effects in a typical TST calculation, we make an effort to address this issue by introducing error bars as part of our approach.³¹

More details about the theoretical background behind eqn (10) are described in previous publications,^{30,31,37–44} and these also discuss the necessary steps to calculate the parameters described above.^{30,31,35,37} The present work focuses on the quality of the MC-TST/CTSR theoretical results for the 10 case studies for reaction (3), and the interested reader can consult the previous references for technical details.

3 Results and discussion

Experimental information for all 10 rate constants and respective Arrhenius fits of the form $k(T) = A \exp(-B/T)$ were gathered from a recent database⁸⁵ which compiles rate coefficients for the gas-phase reactions of organic compounds with OH, Cl, NO₃ radicals and O₃.

To assess the deviation between the experimental and calculated rate constants, we will use⁸⁶ a factor ρ defined by

$$\rho = \begin{cases} \frac{k_{\text{OH}}(\text{calc})}{k_{\text{exp}}} & k_{\text{OH}}(\text{calc}) > k_{\text{exp}}, \\ \frac{k_{\text{exp}}}{k_{\text{OH}}(\text{calc})} & k_{\text{exp}} > k_{\text{OH}}(\text{calc}) \end{cases} \quad (16)$$

This is always larger than one and allows a quantification of the average $\langle \rho \rangle$ deviation.

For each compound we have extended our MC-TST calculations to a temperature interval relevant for atmospheric conditions in the troposphere,^{12,85,87} namely $230 \leq T/K \leq 320$, and fitted the corresponding data to an Arrhenius-Kooij equation^{85,88,89} of the form $k(T) = A \exp(-B/T)(T/298)^n$. This provides the necessary flexibility to model the curvature of the data points caused by the quantum tunneling factor in eqn (10) and the kinetic model that includes the formation of the prereactant complex^{85,90} in eqn (8). The rate constants have been assigned error bars by propagating the error associated with the barrier heights,³⁴ which are frequently the largest source of uncertainty in a rate constant calculation.³¹ In a recent work,³⁴ the error bars were estimated by assessing the behavior of a M08-HX-based model chemistry against three specific OH hydrogen transfer reactions of the HTBH38/08 database⁹¹ that bared a resemblance to the reactions under analysis.^{31,34} Such a procedure lead to a mean unsigned error (MUE) of 0.6 kcal mol⁻¹, which was taken⁹² as the uncertainty (σ_V^\ddagger) associated to V^\ddagger and propagated accordingly. However, in the present work, we have benchmarked our M08-HX/apcseg-2//M08-HX/pcseg-2 barrier heights against corresponding high-level coupled-cluster calculations, namely CCSD(T)/cc-pVTZ//M08-HX/pcseg-2. Such a comparison between model chemistries yielded a MUE of approximately 0.3 kcal mol⁻¹, which was used⁹² as σ_V^\ddagger for the present 10 barrier heights: $V^\ddagger \pm 0.3$ kcal mol⁻¹. This value is in line with the well known fact that the M08-HX functional performs well in describing hydrogen abstraction barrier heights, yielding MUE values of 1.00 and 0.73 kcal mol⁻¹ for the

DBH76 and HTBH38 data sets involving this particular type of barrier heights.⁷⁵ The coupled-cluster calculations were performed with Gaussian⁷¹ and the details of this analysis can be found in Section 3.6.

Furthermore, the temperature interval used for plotting the experimental Arrhenius curves corresponds to the intersection between our selected $T_{\text{calc}} = [230, 320]$ temperature interval and the temperature interval identified in ref. 85 for each particular OVOC, $T_{\text{exp}} = [T_{\text{min}}, T_{\text{max}}]$: $T_{\text{exp}}^{\text{plot}} = T_{\text{calc}} \cap T_{\text{exp}} = [T_1, T_2]$.

For each OVOC with experimental Arrhenius parameters we can define an s^{OVOC} similarity index based on T_1 and T_2 temperatures:

$$s^{\text{OVOC}} = |\Delta(T_1) - \Delta(T_2)|^{-1} \quad (17)$$

where

$$\Delta(T_x) = \log k_{\text{exp}}(T_x) - \log k_{\text{AK}}(T_x), \quad (18)$$

k_{exp} is defined by the experimental Arrhenius parameters and k_{AK} is the corresponding Arrhenius-Kooij quantity obtained with our procedure. Eqn (17) provides a measure of the similarity of the temperature variation between k_{exp} and k_{AK} in the $T_{\text{exp}}^{\text{plot}}$ temperature interval, and s^{OVOC} thus evaluates the quality of our theoretical model in reproducing the observed experimental behaviour in a given temperature interval. Note that for two parallel straight lines we have maximum similarity with $\Delta(T_1) - \Delta(T_2) = 0$, causing s^{OVOC} to be undefined. However, in this work we will be comparing straight lines (k_{exp}) with curved ones (k_{AK}), which means that the higher the value of s^{OVOC} is, the greater the similarity between the trends given by both equations. We recognize that there are a multitude of mathematical possibilities that could yield very high (or even undefined) values for s^{OVOC} while representing very little similarity between trends, but these possibilities are unlikely for the present systems, and we will use s^{OVOC} to compare the quality of our Arrhenius-Kooij curves. For each compound we will show three different Arrhenius-Kooij curves which will differ in the way V^\ddagger and $\kappa(T)$ are calculated. Two of them will use the protocol's recipe for V^\ddagger with $\kappa(T)$ calculated through the Eckart and iSCTST methods (M08-HX-Eckart and M08-HX-iSCTST, respectively), while a third one will be generated by calculating V^\ddagger with our coupled-cluster approach and with the $\kappa(T)$ tunneling correction obtained with the iSCTST method (CCSD(T)-iSCTST).

Our calculated branching ratios will be compared with the results given from ref. 13, which were derived using the SARs from Kwok and Atkinson.⁹³ The SAR approach is “necessarily subject to significant uncertainty” and is used¹³ “to provide a qualitative measure of the reactivity at various sites”. While the SARs from Kwok and Atkinson perform well for simple hydrocarbons over large temperature ranges, they are much less accurate for OVOCs, where the “agreement between experimental rate data and values calculated using the SAR method for reactions of OH with compounds containing oxygenated functional groups is in general rather poor”.⁹ One of the reasons for such a poor agreement is suggested to lie in the difficulty of



incorporating long-range activating effects of oxygenated functional groups in the SAR models.⁹

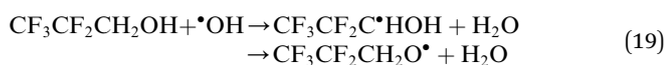
Note that the results for both the rate constants at 298.15 K and respective branching ratios will be presented with the protocol's usual procedure of calculating V^\ddagger at the M08-HX/apcseg-2//M08-HX/pcseg-2 level of theory along with $\kappa(T)$ being obtained with the asymmetric Eckart model.³¹

3.1 Hydrofluoroalcohols

Of the five families of studied OVOCs, hydrofluoroalcohols were the only one that required adjustments to the CTSR procedure. The CTSR code first randomly generates 'OH-molecule structures which are then subjected to constraints targeting an optimization of TSs at the PM3 level of theory. The three constraints used throughout this work were: (1) the distance between the 'OH oxygen atom and the hydrogen atom under attack must lie in the [1.2, 1.4] Å interval; (2) the distance between the 'OH oxygen atom and all reactants' non-hydrogen atoms must be greater than 1.70 Å and (3) the distance between the 'OH hydrogen atom and all reactants' hydrogen atoms must be greater than 0.70 Å. Structures are then generated by the CTSR code and subsequently used as input geometry in PM3 saddle-point optimizations. Next, the converged PM3 calculations are used as starting structures for the saddle-point optimizations at the M08-HX/pcseg-2 level. This procedure, however, did not lead to the discovery of any TSs regarding the 'OH attack on the -OH group at the PM3 level, even if the numerical value of the constraints were changed. Consequently, we then employed a more robust and computationally more demanding procedure that used MP2/cc-pVDZ saddle-point optimizations instead of the very fast PM3 step. The only altered constraint was the first, where the distance between the 'OH oxygen atom and the hydrogen atom under attack was limited to the interval [1.05, 1.35] Å. This modification in the procedure led to the finding of several TSs involving an attack on the -OH group.

We should emphasize that the choice of using the MP2/cc-pVDZ level was preceded by an initial test with the more obvious M08-HX/pcseg-1 level of theory. However, the latter approach yielded less TSs, with the optimization procedure often showing convergence difficulties. It should be emphasized that the use of the MP2/cc-pVDZ procedure represents an *ad hoc* fix for this particular system and there may be more cases where the original CTSR procedure³⁷ needs to be modified.

3.1.1 CF₃CF₂CH₂OH. The reaction between CF₃CF₂CH₂OH and 'OH can proceed through two different abstraction sites, with the OH radical abstracting a hydrogen atom from either the -CH₂ or the -OH group:



The calculated branching ratios for the two abstraction sites are $I_{\text{CH}_2} = 99\%$ and $I_{\text{OH}} = 1\%$, which are significantly different from the SAR predictions of $I_{\text{CH}_2} = 58\%$ and $I_{\text{OH}} = 42\%$. The low fraction of reactions leading to abstraction from the -OH

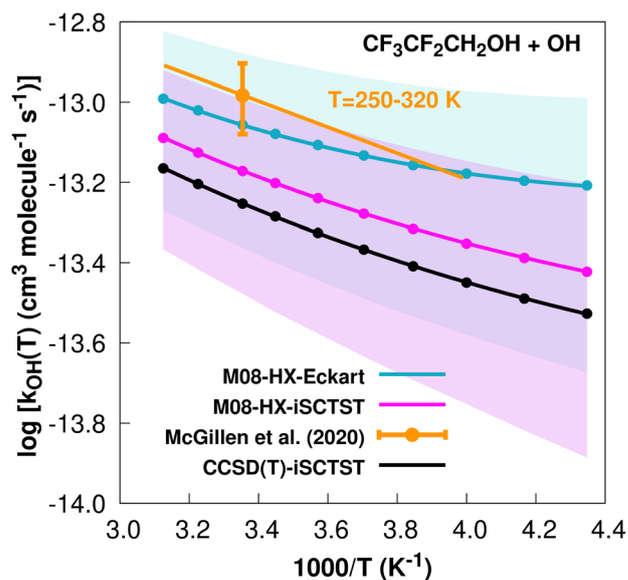
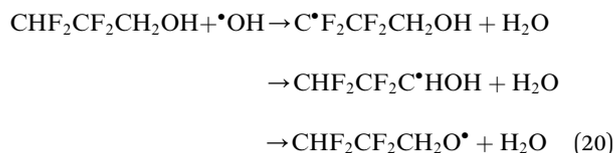


Fig. 1 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the CF₃CF₂CH₂OH + 'OH reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Standard Arrhenius fit (solid orange line) based on experimental data⁸⁵ in the $T_{\text{exp}}^{\text{plot}} = [250, 320]$ K temperature interval together with the recommended experimental rate coefficient at 298.15 K (color online).

group is due to the fact that the lowest energy TS corresponding to an abstraction from the -OH group is 1.87 kcal mol⁻¹ above TS₀, which corresponds an hydrogen abstraction from the -CH₂ group. The corresponding V^\ddagger barrier height is 1.82 kcal mol⁻¹.

The fit to the Arrhenius–Kooij equation provides $k(298.15 \text{ K}) = (8.8 \pm 3.3) \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, while the experimental results^{85,94} yield an Arrhenius expression of $k(T) = 1.28 \times 10^{-12} \exp(-748/T) \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ valid in the [250, 430] K temperature interval. The experimental recommended rate constant at 298.15 K is $k(298.15 \text{ K}) = (1.04 \pm 0.21) \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 1.2$. The $k_{\text{OH}}(\text{calc})$ points in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 1, together with the Arrhenius–Kooij fit to those points and the experimental Arrhenius fit given above.

3.1.2 CHF₂CF₂CH₂OH. For the reaction between CHF₂CF₂CH₂OH and 'OH there are three different abstraction sites, since the OH radical can attack the -CHF₂, -CH₂ and -OH groups:



The calculated branching ratios for the three abstraction sites are: $I_{\text{CHF}_2} = 5\%$, $I_{\text{CH}_2} = 94\%$ and $I_{\text{OH}} = 1\%$, which can be compared with the SAR predictions: $I_{\text{CHF}_2} = 3\%$, $I_{\text{CH}_2} = 25\%$ and $I_{\text{OH}} = 72\%$. Note that the SAR predictions of $I_{\text{CHF}_2} = 7\%$, $I_{\text{CH}_2} = 30\%$ and $I_{\text{OH}} = 69\%$ given in ref. 13 are incorrect, since



they do not sum up to 100%. The large MC-TST/CTSR result of $\Gamma_{\text{CH}_2} = 94\%$ is explained by the fact that the vast majority of the lowest energy TSs correspond to the $\cdot\text{OH}$ attack at the $-\text{CH}_2$ group. Of the 54 TSs obtained *via* the CTSR procedure, the 22 lowest energy ones correspond to an attack of the OH radical to the $-\text{CH}_2$ and $-\text{CHF}_2$ groups. The lowest energy TSs corresponding to the $\cdot\text{OH}$ attack at the $-\text{CHF}_2$ and $-\text{OH}$ groups are 0.52 and 2.13 kcal mol $^{-1}$ above TS_0 , respectively. The V^\ddagger barrier height is 0.80 kcal mol $^{-1}$ and concerns an hydrogen abstraction from the $-\text{CH}_2$ group.

The fit to the Arrhenius–Kooij equation yields $k(298.15 \text{ K}) = (4.4 \pm 1.7) \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, while the experimental results^{85,95} gives an Arrhenius expression with $k(T) = 1.31 \times 10^{-12} \exp(-605/T) \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ valid in the [263, 358] K temperature interval. The experimental recommended rate constant at 298.15 K is $k(298.15 \text{ K}) = (1.85 \pm 0.70) \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 2.4$. The $k_{\text{OH}}(\text{calc})$ points calculated in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 2, together with the Arrhenius–Kooij fit to those points and the experimental Arrhenius fit given above.

Fig. 1 and 2 show that the $\text{CHF}_2\text{CF}_2\text{CH}_2\text{OH} + \cdot\text{OH}$ reaction is significantly slower than the $\text{CF}_3\text{CF}_2\text{CH}_2\text{OH} + \cdot\text{OH}$ reaction. Antiñolo *et al.*⁹⁵ suggested that the difference between the rate constants may be due to an important role of hydrogen abstraction at the $-\text{CHF}_2$ site or a long-range inductive effect of the extra F atom in $\text{CF}_3\text{CF}_2\text{CH}_2\text{OH}$ responsible for deactivating the hydrogen abstraction in the $-\text{CH}_2$ group. Our results clearly point to the latter hypothesis for two reasons. First, our calculated value of $\Gamma_{\text{CHF}_2} = 5\%$ indicates that the hydrogen abstraction at the $-\text{CHF}_2$ group has only a minor contribution to the final rate constant. Second, our calculations show that the V^\ddagger

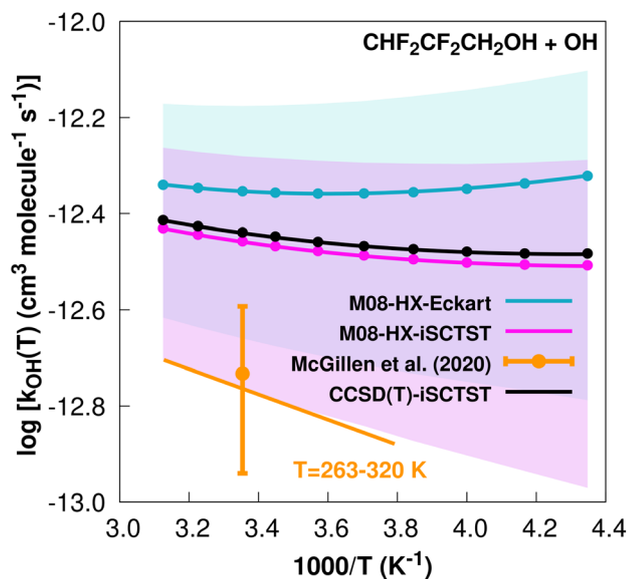


Fig. 2 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the $\text{CHF}_2\text{CF}_2\text{CH}_2\text{OH} + \cdot\text{OH}$ reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Standard Arrhenius fit (solid orange line) based on experimental data⁸⁵ in the $T_{\text{exp}}^{\text{plot}} = [263, 320]$ K temperature interval together with the recommended experimental rate coefficient at 298.15 K (color online).

barrier height, which is a major factor determining the reactivity,^{30,32–34} is approximately 1 kcal mol $^{-1}$ lower for the $\text{CHF}_2\text{CF}_2\text{CH}_2\text{OH} + \cdot\text{OH}$ reaction. This strongly suggests that the chemical environment of the $-\text{CH}_2$ group for both sets of reactants and TSs may in fact be different, for example involving intramolecular hydrogen bonds for reactants and TSs, where a bond critical point⁹⁶ is likely to be found:^{97,98} $\text{C}-\text{H}\cdots\text{F}$ (reactants) and $\text{C}-\text{H}\cdots\text{O}$, $\text{O}-\text{H}\cdots\text{O}$ and $\text{O}-\text{H}\cdots\text{F}$ (TSs). One can argue that the extra F atom in $\text{CF}_3\text{CF}_2\text{CH}_2\text{OH}$ increases the possibility of forming intramolecular hydrogen bonds which could lead to a larger difference in covalency (λ descriptor of ref. 34, linearly correlated with the size of the barrier heights) between these possible intramolecular hydrogen bonds in TS_0 and OVOC_0 of the $\text{CF}_3\text{CF}_2\text{CH}_2\text{OH} + \cdot\text{OH}$ reaction with respect to the $\text{CHF}_2\text{CF}_2\text{CH}_2\text{OH} + \cdot\text{OH}$ reaction, reflecting an increasing difficulty in breaking these bonds.

3.2 Hydrofluoroaldehydes

3.2.1 $\text{CF}_3\text{CF}_2\text{CHO}$. The $\text{CF}_3\text{CF}_2\text{CHO} + \text{OH}$ reaction can only proceed with the $\cdot\text{OH}$ hydrogen abstraction of the $-\text{CHO}$ aldehydic hydrogen



which presents a V^\ddagger barrier height of 0.55 kcal mol $^{-1}$.

The fit to the Arrhenius–Kooij equation yields $k(298.15 \text{ K}) = (6.8 \pm 0.7) \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, with the experimental results^{13,85} yielding an Arrhenius expression of $k(T) = 2.05 \times 10^{-12} \exp(-378/T) \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ valid in the [252, 373] K temperature interval. The experimental recommended rate constant at 298.15 K is $k(298.15 \text{ K}) = (5.8 \pm 0.9) \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 1.2$. The $k_{\text{OH}}(\text{calc})$ points calculated in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 3, together with the Arrhenius–Kooij fit to those points and the experimental Arrhenius fit given above.

3.2.2 $\text{CF}_3\text{CH}_2\text{CHO}$. In the case of the $\text{CF}_3\text{CH}_2\text{CHO} + \text{OH}$ reaction the OH radical can attack the $-\text{CH}_2$ and the $-\text{CHO}$ groups:



The calculated branching ratios for the two abstraction sites are: $\Gamma_{\text{CH}_2} = 0\%$ and $\Gamma_{\text{CHO}} = 100\%$, which agree with the SAR predictions. Our MC-TST/CTSR results for these branching ratios arise from the fact that the six lowest TSs refer to hydrogen abstractions at the $-\text{CHO}$ group within the energy interval of $0 \leq \Delta E_{\text{TS}_i} \leq 0.54 \text{ kcal mol}^{-1}$, while the remaining six TSs correspond to hydrogen abstractions at the $-\text{CH}_2$ group in the energy interval of $4.56 \leq \Delta E_{\text{TS}_i} \leq 5.46 \text{ kcal mol}^{-1}$. The V^\ddagger barrier height is $-0.38 \text{ kcal mol}^{-1}$, but the fact that it is slightly submerged below the reactants should not constitute a problem, since this is a well defined TS with ZPE-corrected energies of 1.5 and 30 kcal mol $^{-1}$ above the PRC and PC, respectively.

The fit to the Arrhenius–Kooij equation yields $k(298.15 \text{ K}) = (3.5 \pm 0.4) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, while the experimental



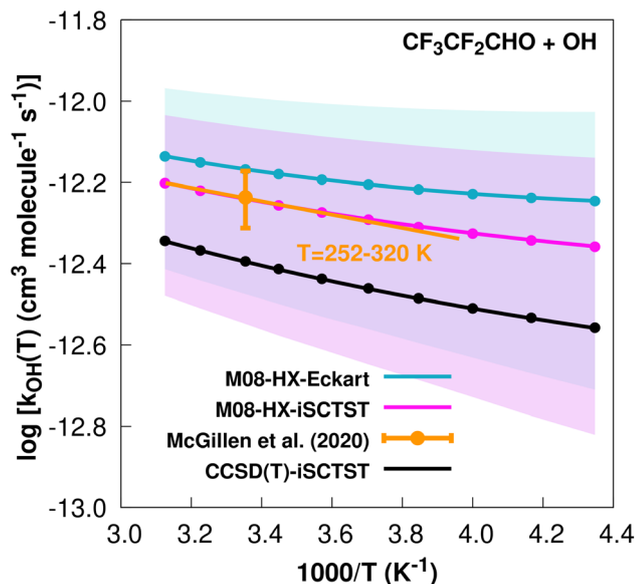


Fig. 3 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the $\text{CF}_3\text{CF}_2\text{CHO} + \cdot\text{OH}$ reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Standard Arrhenius fit (solid orange line) based on experimental data⁸⁵ in the $T_{\text{exp}}^{\text{plot}} = [252, 320]$ K temperature interval together with the recommended experimental rate coefficient at 298.15 K (color online).

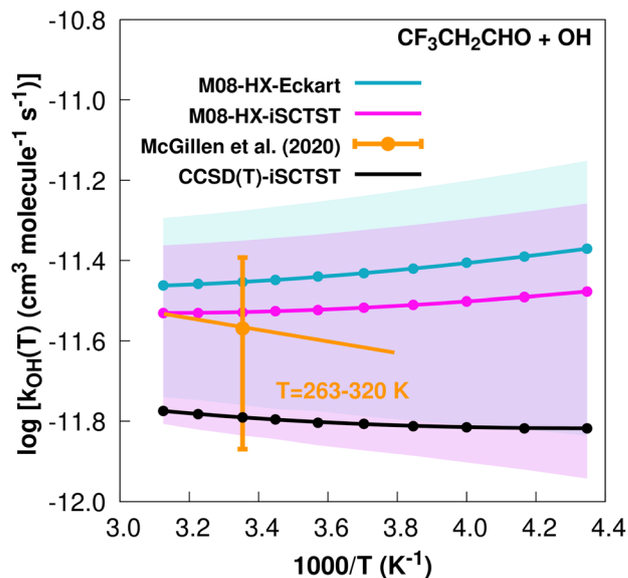


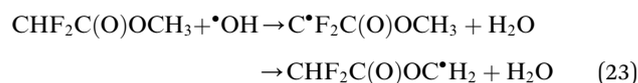
Fig. 4 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the $\text{CF}_3\text{CH}_2\text{CHO} + \cdot\text{OH}$ reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Standard Arrhenius fit (solid orange line) based on experimental data⁸⁵ in the $T_{\text{exp}}^{\text{plot}} = [263, 320]$ K temperature interval together with the recommended experimental rate coefficient at 298.15 K (color online).

results^{85,94} yield an Arrhenius expression of $k(T) = 8.34 \times 10^{-12} \exp(-334/T) \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ valid in the $[263, 371]$ K temperature interval. The experimental recommended rate constant at 298.15 K is $k(298.15 \text{ K}) = (2.7 \pm 1.4) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 1.3$. The $k_{\text{OH}}(\text{calc})$ points calculated in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 4, together with the Arrhenius–Kooij fit to those points and the experimental Arrhenius fit given above.

It is worthwhile mentioning that the higher value of the rate constant for the $\text{CF}_3\text{CH}_2\text{CHO} + \cdot\text{OH}$ reaction with respect to the $\text{CF}_3\text{CF}_2\text{CHO} + \cdot\text{OH}$ reaction originates mainly in the different values of V^\ddagger , with the former reaction having a V^\ddagger value $0.93 \text{ kcal mol}^{-1}$ lower than the latter $\text{CF}_3\text{CF}_2\text{CHO} + \cdot\text{OH}$ reaction. Similarly to the two hydrofluoroalcohols above, such behaviour suggests the existence of a different chemical environment at the corresponding $-\text{CHO}$ groups, where the common explanation considers a long range destabilizing inductive effect of the $-\text{CF}_2$ group in these TSs, thereby decreasing the reactivity towards the OH radical in the $\text{CF}_3\text{CF}_2\text{CHO}$ hydrofluoroaldehyde. More generally, and as suggested above, the two extra F atoms in this system increase the number of possible intramolecular hydrogen bonds, which could be increasing the λ descriptor³⁴ and therefore, according to this model, would explain a higher value of V^\ddagger for the $\text{CF}_3\text{CF}_2\text{CHO} + \cdot\text{OH}$ reaction.

3.3 Hydrofluoroesters

3.3.1 $\text{CHF}_2\text{C}(\text{O})\text{OCH}_3$. The $\text{CHF}_2\text{C}(\text{O})\text{OCH}_3 + \text{OH}$ reaction can proceed *via* hydrogen abstraction at two different locations, the $-\text{CHF}_2$ and the $-\text{CH}_3$ groups:



The calculated branching ratios for the two abstraction sites are: $\Gamma_{\text{CHF}_2} = 64\%$ and $\Gamma_{\text{CH}_3} = 36\%$, which are considerably different from the SAR predictions of $\Gamma_{\text{CHF}_2} = 15\%$ and $\Gamma_{\text{CH}_3} = 85\%$. This is an interesting result, considering that 13 out of the 20 TSs found with the CTSR method concern OH radical attacks at the CH_3 group and that they account for the four lowest energy TSs. The next seven TSs correspond to reactions at the CHF_2 group, with the associated higher-valued partition functions compensating for their lower Boltzmann factors, thus generating a higher value for Γ_{CHF_2} . The lowest energy TS corresponding to the $\cdot\text{OH}$ attack at the $-\text{CHF}_2$ group is $0.40 \text{ kcal mol}^{-1}$ above TS_0 . The V^\ddagger barrier height is $1.68 \text{ kcal mol}^{-1}$.

The fit to the Arrhenius–Kooij equation yields $k(298.15 \text{ K}) = (4.7 \pm 1.4) \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, but the only available experimental result^{13,85} is the rate coefficient at 298.15 K: $k(298.15 \text{ K}) = (1.33 \pm 0.33) \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 2.8$. The $k_{\text{OH}}(\text{calc})$ points calculated in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 5, together with the Arrhenius–Kooij fit to those points and the experimental rate constant at 298.15 K.

3.3.2 $\text{CF}_3\text{CFHOC}(\text{O})\text{H}$. The $\text{CF}_3\text{CFHOC}(\text{O})\text{H}$ hydrofluoroester can be attacked by the OH radical in two different locations, the $-\text{CFH}$ and the $-\text{OC}(\text{O})\text{H}$ groups:



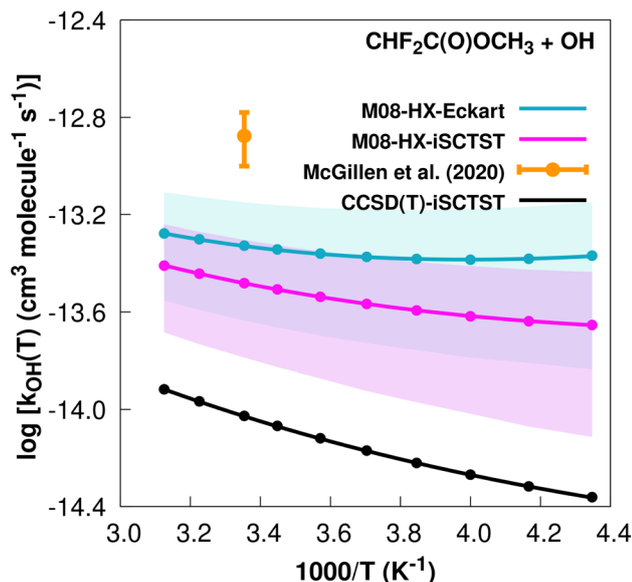


Fig. 5 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the $\text{CHF}_2\text{C}(\text{O})\text{OCH}_3 + \cdot\text{OH}$ reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Solid orange circle shows the recommended value⁸⁵ for the rate constant at 298.15 K (color online).

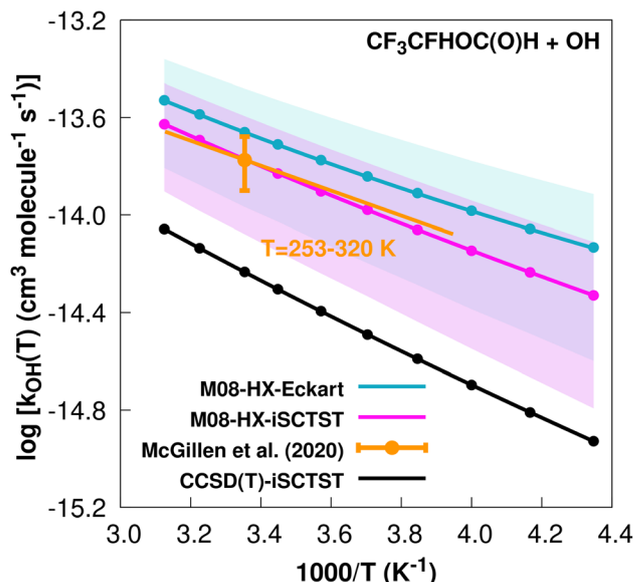


Fig. 6 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the $\text{CF}_3\text{CFHOC}(\text{O})\text{H} + \cdot\text{OH}$ reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Standard Arrhenius fit (solid orange line) based on experimental data⁸⁵ in the $T_{\text{exp}}^{\text{plot}} = [253, 320]$ K temperature interval together with the recommended experimental rate coefficient at 298.15 K (color online).

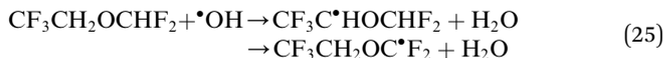


The calculated branching ratios for the two abstraction sites are $\Gamma_{\text{CFH}} = 5\%$ and $\Gamma_{\text{OC}(\text{O})\text{H}} = 95\%$ are similar to the SAR predictions of $\Gamma_{\text{CFH}} = 29\%$ and $\Gamma_{\text{OC}(\text{O})\text{H}} = 71\%$. The V^\ddagger barrier height is 2.79 kcal mol⁻¹ and corresponds to a hydrogen abstraction from the OC(O)H group. Interestingly, within the four lowest energy TSs, the second lowest corresponds to an OH radical attacking the –CFH group with its energy being only 0.04 kcal mol⁻¹ above TS₀. However, its partition function is one order of magnitude below the ones associated to the other three TSs, which is the main factor behind the difference in the branching ratios.

The fit to the Arrhenius–Kooij equation yields $k(298.15 \text{ K}) = (2.2 \pm 0.6) \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, while the experimental results^{13,85} yield an Arrhenius expression of $k(T) = 8.84 \times 10^{-13} \exp(-1181/T) \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ valid in the [253, 328] K temperature interval. The experimental recommended rate constant at 298.15 K is given by $k(298.15 \text{ K}) = (1.68 \pm 0.42) \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 1.3$. The $k_{\text{OH}}(\text{calc})$ points calculated in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 6, together with the Arrhenius–Kooij fit to those points and the experimental Arrhenius fit given above.

3.4 Hydrofluoroethers

3.4.1 $\text{CF}_3\text{CH}_2\text{OCHF}_2$. The $\text{CF}_3\text{CH}_2\text{OCHF}_2 + \text{OH}$ reaction can proceed with an attack of the OH radical on the –CH₂ and –CHF₂ groups



The calculated branching ratios for the two abstraction sites are: $\Gamma_{\text{CH}_2} = 65\%$ and $\Gamma_{\text{CHF}_2} = 35\%$, which are almost identical to the SAR predictions of $\Gamma_{\text{CH}_2} = 60\%$ and $\Gamma_{\text{CHF}_2} = 40\%$. The higher predominance of Γ_{CH_2} in this case arises from the seven lowest energy TSs corresponding to OH radical attack on the –CH₂ group, with the V^\ddagger barrier height having a value of 3.37 kcal mol⁻¹. The lowest energy TS corresponding to the $\cdot\text{OH}$ attack at the –CHF₂ group is 0.55 kcal mol⁻¹ above TS₀.

The fit to the Arrhenius–Kooij equation yields $k(298.15 \text{ K}) = (2.5 \pm 1.2) \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, while the experimental results^{85,99} yield an Arrhenius expression of $k(T) = 2.90 \times 10^{-12} \exp(-1660/T) \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ valid in the [253, 460] K temperature interval. The experimental recommended rate constant at 298.15 K is given by $k(298.15 \text{ K}) = (1.1 \pm 0.2) \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 2.3$. The $k_{\text{OH}}(\text{calc})$ points calculated in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 7, together with the Arrhenius–Kooij fit to those points and the experimental Arrhenius fit given above.

3.4.2 $\text{CF}_3\text{CHFOCHF}_2$. The $\text{CF}_3\text{CHFOCHF}_2 + \text{OH}$ reaction can occur with attack of the OH radical on two different groups, namely –CHF and –CHF₂



The calculated branching ratios for the two abstraction sites are: $\Gamma_{\text{CHF}} = 67\%$ and $\Gamma_{\text{CHF}_2} = 33\%$, which almost coincide with



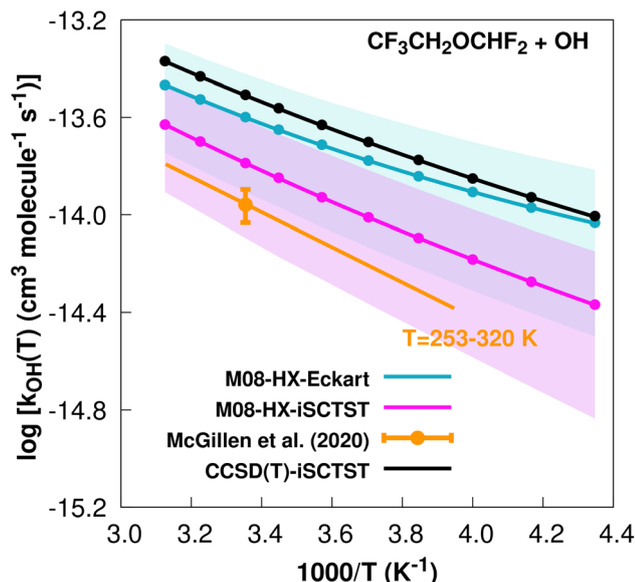


Fig. 7 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the $\text{CF}_3\text{CH}_2\text{OCHF}_2 + \cdot\text{OH}$ reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Standard Arrhenius fit (solid orange line) based on experimental data⁸⁵ in the $T_{\text{exp}}^{\text{plot}} = [253, 320]$ K temperature interval together with the recommended experimental rate coefficient at 298.15 K (color online).

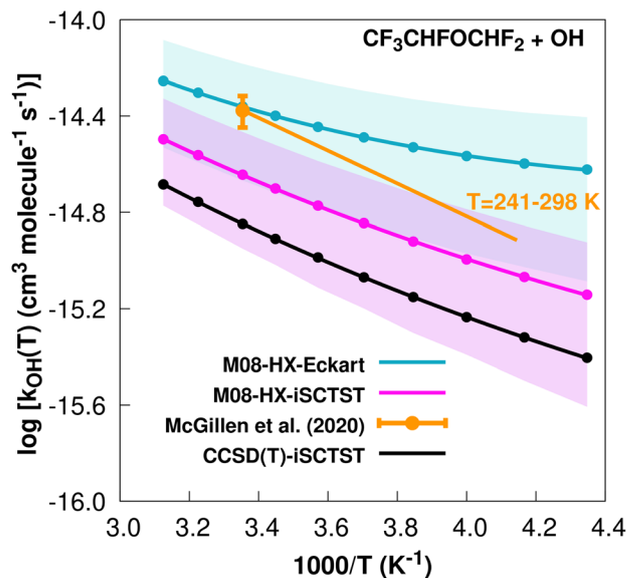


Fig. 8 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the $\text{CF}_3\text{CHFOCHF}_2 + \cdot\text{OH}$ reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Standard Arrhenius fit (solid orange line) based on experimental data⁸⁵ in the $T_{\text{exp}}^{\text{plot}} = [241, 298]$ K temperature interval together with the recommended experimental rate coefficient at 298.15 K (color online).

the SAR predictions of $\Gamma_{\text{CHF}} = 60\%$ and $\Gamma_{\text{CHF}_2} = 40\%$. A total of 13 TSs are found, of which four correspond to OH radical attack on the $-\text{CHF}_2$ group, with two of these being the highest energy TSs. Because the 13 Boltzmann-weighted partition functions have the same order of magnitude, the nine vs. four TSs tip the balance in favour of Γ_{CHF} . The V^\ddagger barrier height is $4.03 \text{ kcal mol}^{-1}$ and concerns an hydrogen abstraction from the $-\text{CHF}$ group.

The fit to the Arrhenius–Kooij equation yields $k(298.15 \text{ K}) = (4.4 \pm 1.6) \times 10^{-15} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, while the experimental results^{85,99} yield an Arrhenius expression of $k(T) = 8.15 \times 10^{-13} \exp(-1570/T) \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ valid in the $[241, 298]$ K temperature interval. The experimental recommended rate constant at 298.15 K is given by $k(298.15 \text{ K}) = (4.2 \pm 0.6) \times 10^{-15} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 1.0$. The $k_{\text{OH}}(\text{calc})$ points calculated in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 8, together with the Arrhenius–Kooij fit to those points and the experimental Arrhenius fit given above.

3.5 Hydrofluoroketones

3.5.1 $\text{CF}_3\text{C}(\text{O})\text{CH}_3$. The $\text{CF}_3\text{C}(\text{O})\text{CH}_3 + \text{OH}$ reaction can only occur *via* an attack of the OH radical on the $-\text{CH}_3$ group



having a V^\ddagger barrier height of $4.07 \text{ kcal mol}^{-1}$.

The fit to the Arrhenius–Kooij equation yields $k(298.15 \text{ K}) = (5.1 \pm 2.2) \times 10^{-15} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$. The only available experimental^{10,85} rate coefficient is $k(298.15 \text{ K}) = (1.1 \pm 0.6) \times$

$10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 2.2$. Note that at the date that this manuscript was being written, the rate coefficient given in the online database⁸⁵ was incorrect, since it was citing the value present in ref. 13, which is actually a typo. The

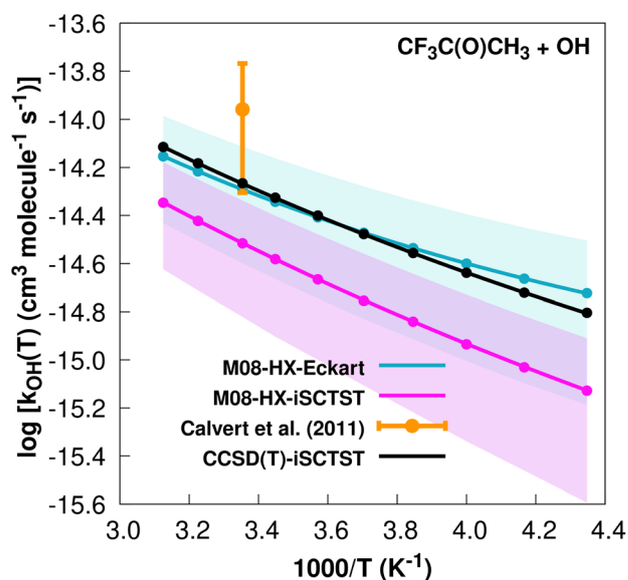


Fig. 9 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the $\text{CF}_3\text{C}(\text{O})\text{CH}_3 + \cdot\text{OH}$ reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Solid orange circle shows the recommended value¹⁰ for the rate constant at 298.15 K (color online).



correct value for the recommended rate constant is given in ref. 10. The $k_{\text{OH}}(\text{calc})$ points calculated in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 9, together with the Arrhenius–Kooij fit to those points and the experimental rate coefficient at 298.15 K.

3.5.2 $\text{CH}_2\text{FC}(\text{O})\text{CH}_3$. In the $\text{CH}_2\text{FC}(\text{O})\text{CH}_3 + \text{OH}$ reaction, the OH radical can attack the $-\text{CH}_2\text{F}$ and the $-\text{CH}_3$ groups:



The calculated branching ratios for the two abstraction sites are: $I_{\text{CH}_2\text{F}} = 80\%$ and $I_{\text{CH}_3} = 20\%$, which are quite different from the SAR predictions of $I_{\text{CH}_2\text{F}} = 39\%$ and $I_{\text{CH}_3} = 61\%$. Note that the SAR predictions of $I_{\text{CH}_2\text{F}} = 39\%$ and $I_{\text{CH}_3} = 51\%$ given in ref. 13 are incorrect, since they do not sum up to 100%. Our MC-TST/CTSR results show that of the six total found TSs, the two lowest energy ones refer to hydrogen abstractions at the $-\text{CH}_2\text{F}$ group with $\Delta E_{\text{TS}} \leq 1.2 \text{ kcal mol}^{-1}$. The V^\ddagger barrier height is $1.63 \text{ kcal mol}^{-1}$. The remaining four TSs have energies above $1.35 \text{ kcal mol}^{-1}$ and correspond to hydrogen abstractions at the $-\text{CH}_3$ group.

The fit to the Arrhenius–Kooij equation yields $k(298.15 \text{ K}) = (1.4 \pm 0.2) \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$. Again, no other experimental results^{13,85} are available besides the rate coefficient at 298.15 K: $k(298.15 \text{ K}) = (2.15 \pm 0.54) \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, which leads to $\rho \approx 1.5$. The $k_{\text{OH}}(\text{calc})$ points calculated in the $230 \leq T/\text{K} \leq 320$ temperature interval are shown in Fig. 10, together with the Arrhenius–Kooij fit to those points and the experimental rate constant at 298.15 K.

It should be noted that at 298.15 K, this rate constant is almost 30 times larger than the rate constant for the $\text{CF}_3\text{C}(\text{O})\text{CH}_3 + \text{OH}$ reaction, and this is mainly due to the V^\ddagger barriers differing by $2.44 \text{ kcal mol}^{-1}$. We refer again to the possible interpretation that because $\text{CF}_3\text{C}(\text{O})\text{CH}_3$ has two more F atoms than $\text{CH}_2\text{FC}(\text{O})\text{CH}_3$, the former molecule will have more possibilities of forming intramolecular hydrogen bonds in the TSs, with a resulting increase in their degree of covalency and therefore resulting in a larger λ descriptor.³⁴

3.6 A summary of the results

Fig. 11 summarizes the results at 298.15 K, where the MC-TST/CTSR and experimental rate constants are shown, along with the ρ value for each reaction. The rate constants span several orders of magnitude and there is an almost equal distribution (60–40%) of MC-TST/CTSR rate coefficients that overestimate or underestimate the experimental values. The average ρ factor by which our M08-HX-Eckart calculations deviate from the experimental results is calculated to be $\langle \rho \rangle = 1.7$, with six cases having $\rho < 2$ and four cases having $2 < \rho < 3$. The remaining two batches of calculations, M08-HX-iSCTST and CCSD(T)-iSCTST, yield $\langle \rho \rangle$ values of 2.0 and 3.4, respectively.

For each reaction we have also performed a comparison between the values of s^{OVOC} for the three sets of curves. Not surprisingly, the Arrhenius–Kooij curves obtained with the iSCTST tunneling factor represent the maximum value of s^{OVOC} for six (three cases for each -iSCTST type of curve) of the seven analyzed reactions, as the iSCTST tunneling factor should better describe the lower temperature region of the [230, 320] K interval in comparison with the Eckart tunneling factor. The only reaction for which an M08-HX-Eckart Arrhenius–Kooij curve represents the highest s^{OVOC} value is the $\text{CF}_3\text{CFHOC}(\text{O})\text{H} + \cdot\text{OH}$ reaction (Fig. 6), although the respective M08-HX-iSCTST curve displays an incredible agreement with the experimental Arrhenius curve, which illustrates what the s^{OVOC} metric really describes. We have therefore considered the M08-HX-iSCTST Arrhenius–Kooij curves as the better choice to describe the trends associated to the temperature dependence of these rate constants. Such a choice is supported by the values of s^{OVOC} for the M08-HX-iSCTST curves, which are always larger than the s^{OVOC} values for the M08-HX-Eckart curves, except the already mentioned exception. This improvement of the M08-HX-iSCTST curves with respect to the M08-HX-Eckart ones is also visually evident in some cases: for example in Fig. 1, 2, 8 and 10. Of the seven possible comparisons between our M08-HX-iSCTST MC-TST/CTSR Arrhenius–Kooij fits and the experimental recommended Arrhenius curves, two of them show values of s^{OVOC} clearly above the remaining five, indicating a higher similarity between k_{exp} and k_{AK} : the value for the $\text{CF}_3\text{-CF}_2\text{CH}_2\text{OH} + \cdot\text{OH}$ reaction with $s = 48$ (Fig. 1) and also for the $\text{CF}_3\text{CF}_2\text{CHO} + \cdot\text{OH}$ reaction with $s = 57$ (Fig. 3). Even though the other five cases have lower values of s^{OVOC} in the interval $8 \leq s^{\text{OVOC}} \leq 15$, they nevertheless exhibit a good similarity between k_{exp} and k_{AK} , especially if it is taken into consideration that the missing error bars from k_{exp} may help to mitigate the gap between the two curves. The exception is the $\text{CF}_3\text{CH}_2\text{CHO} + \cdot\text{OH}$

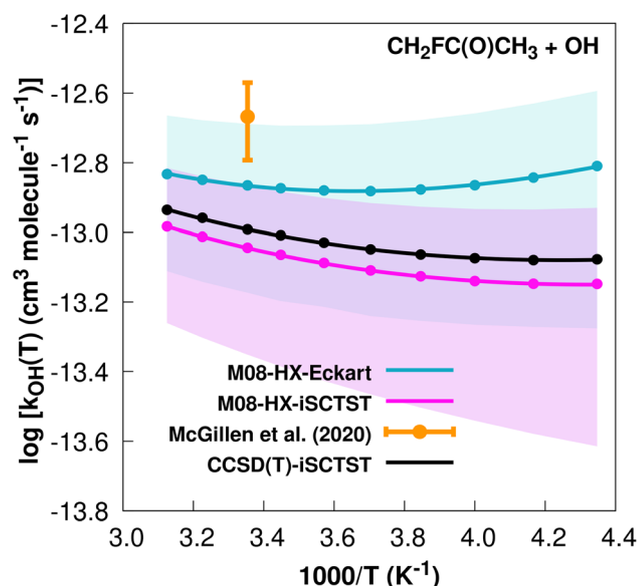


Fig. 10 $k_{\text{OH}}(\text{calc})$ data points (solid blue, purple and black circles) and respective Arrhenius–Kooij fits (solid blue, purple and black lines) with σ error bars (shaded blue and purple areas) for the $\text{CH}_2\text{FC}(\text{O})\text{CH}_3 + \cdot\text{OH}$ reaction in the $T_{\text{calc}} = [230, 320]$ K temperature interval. Solid orange circle shows the recommended value⁸⁵ for the rate constant at 298.15 K (color online).



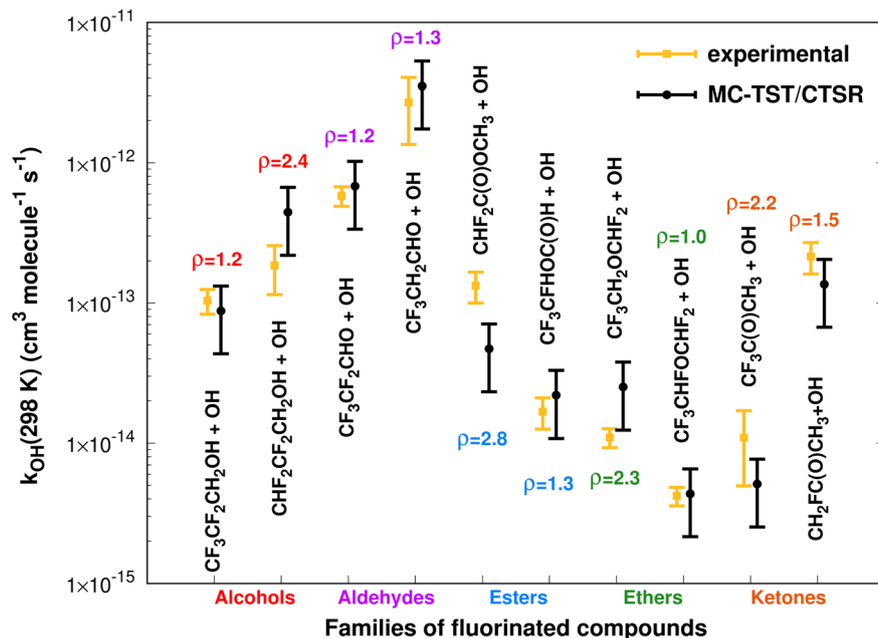


Fig. 11 Comparison between 10 (recommended) experimental and theoretical (this work, eqn (8) and (10)) rate constants at 298.15 K spanning five different families of OVOCs.

reaction, where both M08-HX curves yield an incorrect trend when compared with the experimental curve. Here, only the CCSD(T)-iSCTST curve shows an acceptable behaviour, with an Arrhenius–Kooij curve given by $k_{\text{AK}}(T) = 3.17 \times 10^{-13} \exp(486/T)(T/298)^{2.10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$. Table 1 shows the fitted parameters for all Arrhenius–Kooij fits along with the corresponding values of the relative residual standard deviation (RRSD).¹⁰⁰

Our procedure also improves the branching ratios at 298.15 K relative to those obtained *via* SARs. This is particularly evident in the case of the hydrofluoroalcohols, where the SAR values for T_{OH} are clearly overestimated, since hydrogen-atom abstraction at the –OH group is well-known to be associated with higher barrier heights than hydrogen-atom abstraction at alkyl groups.^{101–104}

Moreover, Table 1 also shows the values for the V_{\ddagger}^{\dagger} barrier heights calculated at the M08-HX and CCSD(T) levels, which were used estimate $\sigma_{V_{\ddagger}^{\dagger}}$ as the value of the MUE⁹²

$$\sigma_{V_{\ddagger}^{\dagger}} \approx \text{MUE} = \frac{\sum_{\text{OVOC}} |V_{\text{M08-HX,OVOC}}^{\ddagger} - V_{\text{CCSD(T),OVOC}}^{\ddagger}|}{10} \quad (29)$$

This equation yields a value of $\sigma_{V_{\ddagger}^{\dagger}} \approx 0.3 \text{ kcal mol}^{-1}$. However, we should stress that each value for $V_{\text{CCSD(T)}}^{\ddagger}$ is an approximation to the true CCSD(T) barrier height, since the CCSD(T)/cc-pVTZ single-point energy calculations were performed at M08-HX/pcseg-2 optimized geometries, not on geometries optimized at the coupled-cluster level of theory. The use of such a methodology is therefore likely to introduce an error of

Table 1 Calculated parameters (A , B and n) for the M08-HX-iSCTST Arrhenius–Kooij fits of the 10 studied reactions, along with the corresponding values of the relative residual standard deviation (RRSD)^a

OVOC	A	B	n	RRSD	$V_{\text{M08-HX}}^{\ddagger}$	$V_{\text{CCSD(T)}}^{\ddagger}$	$k_{\text{OH}}(\text{calc})$	$k_{\text{OH}}^{\text{MC-TST}}$	k_{exp}	$k_{\text{OH}}^{\text{LC-TST}}(\text{calc})$
CF ₃ CF ₂ CH ₂ OH	4.69×10^{-15}	–794	5.27	0.15	1.82	1.93	8.8×10^{-14}	8.8×10^{-14}	1.0×10^{-13}	2.6×10^{-14}
CHF ₂ CF ₂ CH ₂ OH	2.66×10^{-14}	–766	3.38	0.10	0.80	0.77	4.4×10^{-13}	4.5×10^{-13}	1.8×10^{-13}	1.1×10^{-13}
CF ₃ CF ₂ CHO	1.39×10^{-13}	–423	2.66	0.08	0.55	0.76	6.8×10^{-13}	7.0×10^{-13}	5.8×10^{-13}	1.7×10^{-13}
CF ₃ CH ₂ CHO	3.44×10^{-13}	–642	2.01	0.08	–0.38	0.01	3.5×10^{-12}	4.1×10^{-12}	2.7×10^{-12}	8.4×10^{-13}
CHF ₂ C(O)OCH ₃	2.48×10^{-16}	–1457	7.11	0.16	1.68	2.43	4.7×10^{-14}	4.7×10^{-14}	1.3×10^{-13}	1.8×10^{-14}
CF ₃ CFHOC(O)H	6.57×10^{-14}	405	3.40	0.30	2.79	3.42	2.2×10^{-14}	2.2×10^{-14}	1.7×10^{-14}	8.0×10^{-15}
CF ₃ CH ₂ OCHF ₂	1.52×10^{-14}	–20	5.24	0.20	3.37	2.99	2.5×10^{-14}	2.5×10^{-14}	1.1×10^{-14}	2.5×10^{-15}
CF ₃ CHFOCHF ₂	1.62×10^{-16}	–785	7.43	0.39	4.03	4.30	4.4×10^{-15}	4.4×10^{-15}	4.2×10^{-15}	6.7×10^{-16}
CF ₃ C(O)CH ₃	1.67×10^{-15}	–179	6.12	0.25	4.07	3.73	5.1×10^{-15}	5.1×10^{-15}	1.1×10^{-14}	8.1×10^{-16}
CH ₂ FC(O)CH ₃	1.80×10^{-16}	–1852	8.02	0.20	1.63	1.55	1.4×10^{-13}	1.4×10^{-13}	2.2×10^{-13}	7.9×10^{-14}

^a DFT and coupled-cluster barrier heights ($V_{\text{DFT}}^{\ddagger}$ and $V_{\text{CCSD(T)}}^{\ddagger}$) and theoretical ($k_{\text{OH}}(\text{calc})$, $k_{\text{OH}}^{\text{MC-TST}}$ and $k_{\text{OH}}^{\text{LC-TST}}(\text{calc})$, see text) and experimental (k_{exp}) rate constants. A , $k_{\text{OH}}(\text{calc})$, $k_{\text{OH}}^{\text{MC-TST}}$, k_{exp} and $k_{\text{OH}}^{\text{LC-TST}}(\text{calc})$ are in units of $\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, B is in kelvin, RRSD is in % and the V_{\ddagger}^{\dagger} barrier heights are written in kcal mol^{-1} .



unknown magnitude,¹⁰⁵ considering that the CCSD(T) calculations are performed in a topological unknown region of the coupled-cluster PES. In other words, the coupled-cluster calculations involved in obtaining V^\ddagger trade the certainty of being in the correct region of the PES with a M08-HX energy value for a higher accurate CCSD(T) energy value in an unknown topological region of the coupled-cluster PES, which may not even be a stationary point. With these limitations in mind, we then proceeded to propagate the σ_V^\ddagger uncertainty in the fitted Arrhenius–Kooij equations. The $\sigma_V^\ddagger = 0.3 \text{ kcal mol}^{-1}$ value translates into $\sigma_B = 151 \text{ K}$ and the uncertainty associated with the Arrhenius–Kooij equations is calculated as¹⁰⁶

$$\sigma = \sqrt{\left(\frac{\partial k_{AK}(T)}{\partial B}\right)^2 \sigma_B^2} = \pm \frac{k_{AK}(T)}{T} \sigma_B \quad (30)$$

Finally, Table 1 includes the values for $k_{OH}(\text{calc})$, $k_{OH}^{\text{MC-TST}}$, k_{exp} and $k_{OH}^{\text{LC-TST}}(\text{calc})$. Several observations can be made by looking at these four sets of values. The first one is that for cases where $k_{OH}^{\text{MC-TST}} \ll k_1$, $k_{OH}(\text{calc}) \approx k_{OH}^{\text{MC-TST}}$. This is expected just by analyzing eqn (8). The second observation is that when the condition $k_{OH}^{\text{MC-TST}} \ll k_1$ does not hold, $k_{OH}(\text{calc})$ is always closer to the k_{exp} experimental rate constant value than the $k_{OH}^{\text{MC-TST}}$ result. This improvement in the rate constant was already observed by Petit and Harvey⁴⁰ and it should be acknowledged even though we are making an approximation by using the same k_1 value for all studied reactions. The third one refers to the $k_{OH}(\text{calc})$ values based on lowest-conformer TST (LC-TST),^{37,42} which take into account only the lowest energy reactant and TS conformer for each reaction. These calculations therefore ignore the conformational complexity present in all reactions and show the importance of performing the necessary conformational sampling: a quick inspection of the values present in the last column in Table 1 clearly indicates that the values of $k_{OH}^{\text{LC-TST}}(\text{calc})$ are considerably less accurate than the $k_{OH}(\text{calc})$ values, which is confirmed by calculating the average ρ factor for these values: $\langle \rho \rangle = 4.9$.

4 Conclusions

In this work we have tested our cost-effective MC-TST/CTSR protocol for the calculation of bimolecular rate constants by investigating reactions between $\cdot\text{OH}$ and 10 fluorinated OVOCs belonging to five different families: hydrofluoroalcohols, hydrofluoroaldehydes, hydrofluoroesters, hydrofluoroethers and hydrofluoroketones. Our calculated M08-HX-Eckart based rate coefficients agree with the recommended experimental values, on average, to within better a factor of two ($\langle \rho \rangle = 1.7$). Given the cost-effective nature of this computational protocol and the diversity of studied systems, this is a promising set of results that builds on the successful implementation of a previous version of our protocol,³¹ where the rate constants of reactions between $\cdot\text{OH}$ and several hydrofluoropolyethers were reported to be in good agreement ($\rho < 5$).

Furthermore, the fitted Arrhenius–Kooij curves based on the iSCTST tunneling correction show a good agreement with the

recommended experimental Arrhenius fits while being a considerable improvement to the M08-HX-Eckart based curves. This implies that MC-TST/CTSR temperature-dependent data can be used to model reactions in the atmosphere in situations where experimental information is not available. The MC-TST/CTSR branching ratios can also offer insight into the underlying mechanisms for why reactions are favored at a particular site.

The present work quantifies the accuracy of the MC-TST/CTSR protocol for predicting reaction rates. Theoretically predicted rate constants may be useful for estimating lifetimes for OVOCs that have not yet been experimentally studied and pave the way for designing saturated OVOCs with short atmospheric lifetimes. The theoretical calculation of rate coefficients for the OH-oxidation of larger fluorinated OVOCs (C_n , $n > 3$) is currently underway.

Author contributions

LPV: conceptualization, formal analysis, investigation, methodology, visualization, writing. FJ: analysis, writing, resources.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

The Coimbra Chemistry Centre – Institute of Molecular Sciences (CQC-IMS) is supported by the Portuguese Foundation for Science and Technology (FCT) through projects UIDB/QUI/0313/2020, UIDP/QUI/0313/2020 and LA/P/0056/2020 (national funds). L. P. V. acknowledges FCT for an Assistant Researcher grant. The numerical results presented in this work were obtained at the Centre for Scientific Computing, Aarhus <https://phys.au.dk/forskning/faciliteter/cscaa/>.

Notes and references

- M. J. Molina and F. S. Rowland, Stratospheric Sink for Chlorofluoromethanes: Chlorine Atom-Catalysed Destruction of Ozone, *Nature*, 1974, **249**, 810–812.
- J. D. Farman, B. G. Gardiner and J. D. Shanklin, Large Losses of Total Ozone in Antarctica Reveal Seasonal ClO_x/NO_x Interaction, *Nature*, 1985, **315**, 207–210.
- United Nations, Montreal Protocol on Substances that Deplete the Ozone Layer – Adjustments and Amendment, *Int. Leg. Mater.*, 1993, **32**, 874–887.
- B. P. McGivern, Conference of the Parties to the Framework Convention on Climate Change: Kyoto Protocol, *Int. Leg. Mater.*, 1998, **37**, 22–43.
- E. A. Heath, Amendment to the Montreal Protocol on Substances that Deplete the Ozone Layer (Kigali Amendment), *Int. Leg. Mater.*, 2017, **56**, 193–205.
- UNEP, *HFCs: A Critical Link in Protecting Climate and the Ozone Layer*, United Nations Environment Programme (UNEP), 2011, p. 36, available online: <https://>



- www.unep.org/resources/report/hfcs-critical-link-protecting-climate-and-ozone-layer, accessed on 2 November 2022.
- Ø. Hodnebrog, M. Etminan, J. S. Fuglestedt, G. Marston, G. Myhre, C. J. Nielsen, K. P. Shine and T. J. Wallington, Global Warming Potentials and Radiative Efficiencies of Halocarbons and Related Compounds: A Comprehensive Review, *Rev. Geophys.*, 2013, **51**, 300–378.
 - T. J. Wallington, M. P. S. Andersen and O. J. Nielsen, in *Advances in Atmospheric Chemistry*, ch. Atmospheric Chemistry of Halogenated Organic Compounds, ed. J. R. Barker, A. L. Steiner and T. J. Wallington, World Scientific, 2017, vol. 1, pp. 305–402.
 - A. Mellouki, G. Le Bras and H. Sidebottom, Kinetics and Mechanisms of the Oxidation of Oxygenated Organic Compounds in the Gas Phase, *Chem. Rev.*, 2003, **103**, 5077–5096.
 - J. G. Calvert, A. Mellouki, J. J. Orlando, M. J. Pilling and T. J. Wallington, *The Mechanisms of Atmospheric Oxidation of the Oxygenates*, Oxford University Press, New York, New York, 1st edn, 2011.
 - A. Mellouki, T. J. Wallington and J. Chen, Atmospheric Chemistry of Oxygenated Volatile Organic Compounds: Impacts on Air Quality and Climate, *Chem. Rev.*, 2015, **115**, 3984–4014.
 - B. J. Finlayson-Pitts and J. N. Pitts Jr, *Chemistry of the Upper and Lower Atmosphere: Theory, Experiments, and Applications*, Academic Press, San Diego, California, 1st edn, 2000.
 - J. G. Calvert, J. J. Orlando, W. R. Stockwell and T. J. Wallington, *The Mechanisms of Reactions Influencing Atmospheric Ozone*, Oxford University Press, New York, New York, 1st edn, 2015.
 - M. J. Kurylo and V. L. Orkin, Determination of Atmospheric Lifetimes via the Measurement of OH Radical Kinetics, *Chem. Rev.*, 2003, **103**, 5049–5076.
 - B. Bolin and H. Rodhe, A Note on the Concepts of Age Distribution and Transit Time in Natural Reservoirs, *Tellus*, 1973, **25**, 58–62.
 - Y. Díaz-de-Mera, A. Aranda, A. Notario, A. Rodríguez, D. Rodríguez and I. Bravo, Photolysis Study of Fluorinated Ketones Under Natural Sunlight Conditions, *Phys. Chem. Chem. Phys.*, 2015, **17**, 22991–22998.
 - Y. Ren, F. Bernard, V. Daële and A. Mellouki, Atmospheric Fate and Impact of Perfluorinated Butanone and Pentanone, *Environ. Sci. Technol.*, 2019, **53**, 8862–8871.
 - M. S. Chiappero, F. E. Malanca, G. A. Argüello, S. T. Woolridge, M. D. Hurley, J. C. Ball, T. J. Wallington, R. L. Waterland and R. C. Buck, Atmospheric Chemistry of Perfluoroaldehydes ($C_xF_{2x+1}CHO$) and Fluorotelomer Aldehydes ($C_xF_{2x+1}CH_2CHO$): Quantification of the Important Role of Photolysis, *J. Phys. Chem. A*, 2006, **110**, 11944–11953.
 - L. Vereecken and J. S. Francisco, Theoretical Studies of Atmospheric Reaction Mechanisms in the Troposphere, *Chem. Soc. Rev.*, 2012, **41**, 6259–6293.
 - L. Vereecken, D. R. Glowacki and M. J. Pilling, Theoretical Chemical Kinetics in Tropospheric Chemistry: Methodologies and Applications, *Chem. Rev.*, 2015, **115**, 4063–4114.
 - L. Vereecken, B. Aumont, I. Barnes, J. W. Bozzelli, M. J. Goldman, W. H. Green, S. Madronich, M. R. McGillen, A. Mellouki, J. J. Orlando, B. Picquet-Varrault, A. R. Rickard, W. R. Stockwell, T. J. Wallington and W. P. L. Carter, Perspective on Mechanism Development and Structure-Activity Relationships for Gas-Phase Atmospheric Chemistry, *Int. J. Chem. Kinet.*, 2018, **50**, 435–469.
 - P. T. Anastas and J. C. Warner, *Green Chemistry: Theory and Practice*, Oxford University Press, Oxford, Great Britain, 1998.
 - H. C. Erythropel, J. B. Zimmerman, T. M. de Winter, L. Petitjean, F. Melnikov, C. H. Lam, A. W. Lounsbury, K. E. Mellor, N. Z. Janković, Q. Tu, L. N. Pincus, M. M. Falinski, W. Shi, P. Coish, D. L. Plata and P. T. Anastas, The Green ChemistREE: 20 Years After Taking Root With the 12 Principles, *Green Chem.*, 2018, **20**, 1929–1961.
 - T. Yu, J. Zheng and D. G. Truhlar, Multi-Structural Variational Transition State Theory. Kinetics of the 1,4-Hydrogen Shift Isomerization of the Pentyl Radical With Torsional Anharmonicity, *Chem. Sci.*, 2011, **2**, 2199–2213.
 - T. Yu, J. Zheng and D. G. Truhlar, Multipath Variational Transition State Theory: Rate Constant of the 1,4-Hydrogen Shift Isomerization of the 2-Cyclohexylethyl Radical, *J. Phys. Chem. A*, 2012, **116**, 297–308.
 - R. Meana-Pañeda and A. Fernández-Ramos, Accounting for Conformational Flexibility and Torsional Anharmonicity in the $H + CH_3CH_2OH$ Hydrogen Abstraction Reactions: A Multi-Path Variational Transition State Theory Study, *J. Chem. Phys.*, 2014, **140**, 174303.
 - J. L. Bao, R. Meana-Pañeda and D. G. Truhlar, Multi-Path Variational Transition State Theory for Chiral Molecules: the Site-Dependent Kinetics for Abstraction of Hydrogen from 2-Butanol by Hydroperoxyl Radical, Analysis of Hydrogen Bonding in the Transition State, and Dramatic Temperature Dependence of the Activation Energy, *Chem. Sci.*, 2015, **6**, 5866–5881.
 - J. L. Bao and D. G. Truhlar, Variational Transition State Theory: Theoretical Framework and Recent Developments, *Chem. Soc. Rev.*, 2017, **46**, 7548–7596.
 - D. Ferro-Costas, E. Martínez-Núñez, J. Rodríguez-Otero, E. Cabaleiro-Lago, C. M. Estévez, B. Fernández, A. Fernández-Ramos and S. A. Vázquez, Influence of Multiple Conformations and Paths on Rate Constants and Product Branching Ratios. Thermal Decomposition of 1-Propanol Radicals, *J. Phys. Chem. A*, 2018, **122**, 4790–4800.
 - L. P. Viegas, Exploring the Reactivity of Hydrofluoropolyethers toward OH Through a Cost-Effective Protocol for Calculating Multiconformer Transition State Theory Rate Constants, *J. Phys. Chem. A*, 2018, **122**, 9721–9732.



- 31 L. P. Viegas, Simplified Protocol for the Calculation of Multiconformer Transition State Theory Rate Constants Applied to Tropospheric OH-Initiated Oxidation Reactions, *J. Phys. Chem. A*, 2021, **125**, 4499–4512.
- 32 L. P. Viegas, Multiconformer Transition State Theory Rate Constants for the Reaction Between OH and α , ω -Dimethoxyfluoropolyethers, *Int. J. Chem. Kinet.*, 2019, **51**, 358–366.
- 33 L. P. Viegas, Theoretical Determination of the OH-initiated Oxidation Rate Constants of α , ω -Dialkoxyfluoropolyethers, *Theor. Chem. Acc.*, 2019, **138**, 65.
- 34 L. P. Viegas and F. Jensen, Reactivity of α , ω -Dihydrofluoropolyethers toward OH Predicted by Multiconformer Transition State Theory and the Interacting Quantum Atoms Approach, *J. Phys. Chem. A*, 2020, **124**, 3460–3470.
- 35 L. P. Viegas, Atmospheric Degradation of Two Hydrofluoroketones: Theoretical Rate Constants for the Gas-Phase OH-Oxidation of HFK-447mcc and HFK-465mc, *Atmosphere*, 2022, **13**, 1256.
- 36 L. P. Viegas, Multiconformer Transition State Theory Rate Constant and Branching Ratios for the OH-Initiated Reaction of $\text{CH}_3\text{OCF}_2\text{CHF}_2$ and its Primary Product, $\text{HC}(\text{O})\text{OCF}_2\text{CHF}_2$, *J. Phys. Org. Chem.*, 2023, **36**, e4470.
- 37 L. P. Viegas, Gas-Phase OH-Oxidation of 2-Butanethiol: Multiconformer Transition State Theory Rate Constant with Constrained Transition State Randomization, *Chem. Phys. Lett.*, 2022, **803**, 139829.
- 38 L. Vereecken and J. Peeters, The 1,5-H-Shift in 1-Butoxy: A Case Study in the Rigorous Implementation of Transition State Theory for a Multirotamer System, *J. Chem. Phys.*, 2003, **119**, 5159–5170.
- 39 A. Fernández-Ramos, B. A. Ellingson, R. Meana-Pañeda, J. M. C. Marques and D. G. Truhlar, Symmetry Numbers and Chemical Reaction Rates, *Theor. Chem. Acc.*, 2007, **118**, 813–826.
- 40 A. S. Petit and J. N. Harvey, Atmospheric Hydrocarbon Activation by the Hydroxyl Radical: A Simple yet Accurate Computational Protocol for Calculating Rate Coefficients, *Phys. Chem. Chem. Phys.*, 2012, **14**, 184–191.
- 41 M. P. Rissanen, T. Kurtén, M. Sipilä, J. A. Thornton, J. Kangasluoma, N. Sarnela, H. Junninen, S. Jørgensen, S. Schallhart, M. K. Kajos, R. Taipale, M. Springer, T. F. Mentel, T. Ruuskanen, T. Petäjä, D. R. Worsnop, H. G. Kjaergaard and M. Ehn, The Formation of Highly Oxidized Multifunctional Products in the Ozonolysis of Cyclohexene, *J. Am. Chem. Soc.*, 2014, **136**, 15596–15606.
- 42 K. H. Möller, R. V. Otkjaer, N. Hyttinen, T. Kurtén and H. G. Kjaergaard, Cost-Effective Implementation of Multiconformer Transition State Theory for Peroxy Radical Hydrogen Shift Reactions, *J. Phys. Chem. A*, 2016, **120**, 10072–10087.
- 43 A. Novelli, L. Vereecken, B. Bohn, H. Dorn, G. L. Gkatzelis, A. Hofzumahaus, F. Holland, D. Reimer, F. Rohrer, S. Rosanka, D. Taraborrelli, R. Tillmann, R. Wegener, Z. Yu, A. Kiendler-Scharr, A. Wahner and H. Fuchs, Importance of Isomerization Reactions for OH Radical Regeneration from the Photo-Oxidation of Isoprene Investigated in the Atmospheric Simulation Chamber SAPHIR, *Atmos. Chem. Phys.*, 2020, **20**, 3333–3355.
- 44 L. Vereecken and B. Nozière, H Migration in Peroxy Radicals Under Atmospheric Conditions, *Atmos. Chem. Phys.*, 2020, **20**, 7429–7458.
- 45 J. C. Hansen and J. S. Francisco, Radical-Molecule Complexes: Changing Our Perspective on the Molecular Mechanisms of Radical-Molecule Reactions and Their Impact on Atmospheric Chemistry, *ChemPhysChem*, 2002, **3**, 833–840.
- 46 R. T. Raines and D. E. Hansen, An Intuitive Approach to Steady-State Kinetics, *J. Chem. Educ.*, 1988, **65**, 757–759.
- 47 F. Xiao, X. Sun, Z. Li and X. Li, Theoretical Study of Radical-Molecule Reactions with Negative Activation Energies in Combustion: Hydroxyl Radical Addition to Alkenes, *ACS Omega*, 2020, **5**, 12777–12788.
- 48 D. L. Singleton and R. J. Cvetanović, Temperature Dependence of the Reaction of Oxygen Atoms with Olefins, *J. Am. Chem. Soc.*, 1976, **98**, 6812–6819.
- 49 J. R. Alvarez-Idaboy, N. Mora-Diez and A. Vivier-Bunge, A Quantum Chemical and Classical Transition State Theory Explanation of Negative Activation Energies in OH Addition To Substituted Ethenes, *J. Am. Chem. Soc.*, 2000, **122**, 3715–3720.
- 50 A. Galano, J. R. Alvarez-Idaboy, G. Bravo-Pérez and M. E. Ruiz-Santoyo, Gas Phase Reactions of C1 – C4 Alcohols with the OH Radical: A Quantum Mechanical Approach, *Phys. Chem. Chem. Phys.*, 2002, **4**, 4648–4662.
- 51 A. Cruz-Torres and A. Galano, On the Mechanism of Gas-Phase Reaction of C1 – C3 Aliphatic Thiols + OH Radicals, *J. Phys. Chem. A*, 2007, **111**, 1523–1529.
- 52 I. W. M. Smith, Collisional Energy Transfer, Intramolecular Vibrational Relaxation and Unimolecular Reactions, *J. Chem. Soc., Faraday Trans.*, 1997, **93**, 3741–3750.
- 53 I. W. M. Smith and A. R. Ravishankara, Role of Hydrogen-Bonded Intermediates in the Bimolecular Reactions of the Hydroxyl Radical, *J. Phys. Chem. A*, 2002, **106**, 4798–4807.
- 54 D. C. McCabe, S. S. Brown, M. K. Gilles, R. K. Talukdar, I. W. M. Smith and A. R. Ravishankara, Kinetics of the Removal of OH($v = 1$) and OD($v = 1$) by HNO_3 and DNO_3 from 253 to 383 K, *J. Phys. Chem. A*, 2003, **107**, 7762–7769.
- 55 R. K. Talukdar, T. Gierczak, D. C. McCabe and A. R. Ravishankara, Reaction of Hydroxyl Radical with Acetone. 2. Products and Reaction Mechanism, *J. Phys. Chem. A*, 2003, **107**, 5021–5032.
- 56 C. Eckart, The Penetration of a Potential Barrier by Electrons, *Phys. Rev.*, 1930, **35**, 1303–1309.
- 57 D. G. Truhlar, A. D. Isaacson, R. T. Skodje and B. C. Garret, Incorporation of Quantum Effects in Generalized-Transition-State Theory, *J. Phys. Chem.*, 1982, **86**, 2252–2261.
- 58 T. N. Truong and D. G. Truhlar, Ab initio Transition State Theory Calculations of the Reaction Rate for $\text{OH} + \text{CH}_4 \rightarrow \text{H}_2\text{O} + \text{CH}_3$, *J. Chem. Phys.*, 1990, **93**, 1761–1769.
- 59 T. N. Truong, Thermal Rates of Hydrogen Exchange of Methane with Zeolite: A Direct *ab Initio* Dynamics Study



- on the Importance of Quantum Tunneling, *Effects*, 1997, **101**, 2750–2752.
- 60 T. N. Truong, W. T. Duncan and M. Tirtowidjojo, A Reaction Class Approach for Modeling Gas Phase Reaction Rates, *Phys. Chem. Chem. Phys.*, 1999, **1**, 1061–1065.
- 61 J. Pfaendtner and L. J. Broadbelt, Elucidation of Structure-Reactivity Relationships in Hindered Phenols *via* Quantum Chemistry and Transition State Theory, *Chem. Eng. Sci.*, 2007, **62**, 5232–5239.
- 62 T. L. Nguyen, J. F. Stanton and J. R. Barker, Ab Initio Reaction Rate Constants Computed Using Semiclassical Transition-State Theory: $\text{HO} + \text{H}_2 \rightarrow \text{H}_2\text{O} + \text{H}$ and Isotopologues, *J. Phys. Chem. A*, 2011, **115**, 5118–5126.
- 63 B. Sirjean, E. Dames, H. Wang and W. Tsang, Tunneling in Hydrogen-Transfer Isomerization of *n*-Alkyl Radicals, *J. Phys. Chem. A*, 2012, **116**, 319–332.
- 64 W. H. Miller, Semiclassical Limit of Quantum Mechanical Transition State Theory for Nonseparable Systems, *J. Chem. Phys.*, 1975, **64**, 1899–1906.
- 65 W. H. Miller, Semi-Classical Theory for Non-Separable Systems: Construction of ‘Good’ Action-Angle Variables for Reaction Rate Constants, *Faraday Discuss. Chem. Soc.*, 1977, **62**, 40–46.
- 66 W. H. Miller, R. Hernandez, N. C. Handy, D. Jayatilaka and A. Willetts, Ab Initio Calculation of Anharmonic Constants for a Transition State, with Application to Semiclassical Transition State Tunneling Probabilities, *Chem. Phys. Lett.*, 1990, **172**, 62–69.
- 67 R. Hernandez and W. H. Miller, Semiclassical Transition State Theory. A New Perspective, *Chem. Phys. Lett.*, 1993, **214**, 129–136.
- 68 A. F. Wagner, Improved Multidimensional Semiclassical Tunneling Theory, *J. Phys. Chem. A*, 2014, **116**, 13089–13100.
- 69 M. J. Cohen, N. C. Handy, R. Hernandez and W. H. Miller, Cumulative Reaction Probabilities for $\text{H} + \text{H}_2 \rightarrow \text{H}_2 + \text{H}$ from a Knowledge of the Anharmonic Force Field, *Chem. Phys. Lett.*, 1992, **192**, 407–416.
- 70 V. Barone, Anharmonic Vibrational Properties by a Fully Automated Second-Order Perturbative Approach, *J. Chem. Phys.*, 2005, **122**, 014108.
- 71 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery Jr, J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, *Gaussian16 Revision B.01*, Gaussian Inc, Wallingford CT, 2016.
- 72 S. M. Greene, X. Shan and D. C. Clary, Rate Constants of Chemical Reactions from Semiclassical Transition State Theory in Full and One Dimension, *J. Chem. Phys.*, 2016, **144**, 244116.
- 73 T. A. H. Burd, X. Shan and D. C. Clary, Hydrogen Tunnelling in the Rearrangements of Carbenes: The Role of Dynamical Calculations, *Phys. Chem. Chem. Phys.*, 2020, **22**, 962–965.
- 74 M. W. Schmidt, K. K. Baldrige, J. A. Boatz, S. T. Elbert, M. S. Gordon, J. H. Jensen, S. Koseki, N. Matsunaga, K. A. Nguyen, S. Su, T. L. Windus, M. Dupuis and J. A. Montgomery Jr, General Atomic and Molecular Electronic Structure System, *J. Comput. Chem.*, 1993, **14**, 1347–1363.
- 75 Y. Zhao and D. G. Truhlar, Exploring the Limit of Accuracy of the Global Hybrid Meta Density Functional for Main-Group Thermochemistry, Kinetics, and Noncovalent Interactions, *J. Chem. Theory Comput.*, 2008, **4**, 1849–1868.
- 76 F. Jensen, Unifying General and Segmented Contracted Basis Sets. Segmented Polarization Consistent Basis Sets, *J. Chem. Theory Comput.*, 2014, **10**, 1074–1085.
- 77 N. M. O’Boyle, T. Vandermeersch, C. J. Flynn, A. R. Maguire and G. R. Hutchison, Confab – Systematic Generation of Diverse Low-Energy Conformers, *J. Cheminf.*, 2011, **3**, 8.
- 78 J.-P. Ebejer, G. M. Morris and C. M. Deane, Freely Available Conformer Generation Methods: How Good Are They?, *J. Chem. Inf. Model.*, 2012, **52**, 1146–1158.
- 79 I. Y. Kanal, J. A. Keith and G. R. Hutchison, A Sobering Assessment of Small-Molecule Force Field Methods for Low Energy Conformer Predictions, *J. Chem. Inf. Model.*, 2018, **118**, e25512.
- 80 D. Ferro-Costas and A. Fernández-Ramos, A Combined Systematic-Stochastic Algorithm for the Conformational Search in Flexible Acyclic Molecules, *Front. Chem.*, 2020, **8**, 16.
- 81 D. Ferro-Costas, I. Mosquera-Lois and A. Fernández-Ramos, TorsiFlex: An Automatic Generator of Torsional Conformers. Application to the Twenty Proteinogenic Amino Acids, *J. Cheminf.*, 2021, **13**, 100.
- 82 J. Espinosa-García, E. A. Ojalvo and J. C. Corchado, Theoretical Rate Constants: On the Error Cancellation Using Conventional Transition-State Theory and Wigner’s Tunnelling Correction, *J. Mol. Struct.: THEOCHEM*, 1994, **303**, 131–139.
- 83 P. J. Lewis, K. A. Bennet and J. N. Harvey, A Computational Study of the Atmospheric Oxidation of Nopinone, *Phys. Chem. Chem. Phys.*, 2005, **7**, 1643–1649.
- 84 G. Piccini, M. Alessio and J. Sauer, Ab Initio Calculation of Rate Constants for Molecule-Surface Reactions with Chemical Accuracy, *Angew. Chem., Int. Ed.*, 2016, **55**, 5235–5237.
- 85 M. R. McGillen, W. P. L. Carter, A. Mellouki, J. J. Orlando, B. Picquet-Varrault and T. J. Wallington, Database for the Kinetics of the Gas-Phase Atmospheric Reactions of Organic Compounds, *Earth Syst. Sci. Data*, 2020, **12**, 1203–1216.



- 86 A. G. Vandeputte, M. K. Sabbe, M.-F. Reyniers, V. V. Speybroeck, M. Waroquier and G. B. Marin, Theoretical Study of the Thermodynamics and Kinetics of Hydrogen Abstractions from Hydrocarbons, *J. Phys. Chem. A*, 2007, **111**, 11771–11786.
- 87 M. Altarawneh, A Closer Look Into the Contribution of Atmospheric Gas-Phase Pathways in the Formation of Perfluorocarboxylic Acids, *Atmospheric Pollut. Res.*, 2021, **12**, 101255.
- 88 D. M. Kooij, Über die Zersetzung des Gasförmigen Phosphorwasserstoffs, *Z. Phys. Chem.*, 1893, **12**, 155–161.
- 89 K. J. Laidler, The Development of the Arrhenius Equation, *J. Chem. Educ.*, 1984, **61**, 494–498.
- 90 W. C. Gardiner Jr, Temperature Dependence of Bimolecular Gas Reaction Rates, *Acc. Chem. Res.*, 1977, **111**, 326–331.
- 91 J. Zhen, Y. Zhao and D. G. Truhlar, The DBH24/08 Database and Its Use to Assess Electronic Structure Model Chemistries for Chemical Reaction Barrier Heights, *J. Chem. Theory Comput.*, 2009, **5**, 808.
- 92 C. J. Willmott and K. Matsuura, Advantages of the mean absolute error (MAE) over the root mean square error (RMSE) in assessing average model performance, *Clim. Res.*, 2005, **30**, 79–82.
- 93 E. S. C. Kwok and R. Atkinson, Estimation of Hydroxyl Radical Reaction Rate Constants for Gas-Phase Organic Compounds Using a Structure-Reactivity Relationship: An Update, *Atmos. Environ.*, 1995, **29**, 1685–1695.
- 94 IUPAC Task Group on Atmospheric Chemical Kinetic Data Evaluation, <http://iupac.pole-ether.fr>.
- 95 M. Antiñolo, S. González, B. Ballesteros, J. Albaladejo and E. Jiménez, Laboratory Studies of CHF₂CF₂CH₂OH and CF₃CF₂CH₂OH: UV and IR Absorption Cross Sections and OH Rate Coefficients between 263 and 358 K, *J. Phys. Chem. A*, 2012, **116**, 6041–6050.
- 96 R. F. W. Bader, *Atoms in Molecules. A Quantum Theory*, Oxford University Press, Oxford, Great Britain, 1990.
- 97 C. R. Wick and T. Clark, On Bond-Critical Points in QTAIM and Weak Interactions, *J. Mol. Model.*, 2018, **24**, 142–150.
- 98 S. Shahbazian, Why Bond Critical Points Are Not Bond Critical Points, *Chem.–Eur. J.*, 2018, **24**, 5401–5405.
- 99 J. B. Burkholder, S. P. Sander, J. Abbatt, J. R. Barker, C. Cappa, J. D. Crouse, T. S. Dibble, R. E. Huie, C. E. Kolb, M. J. Kurylo, V. L. Orkin, C. J. Percival, D. M. Wilmoth and P. H. Wine, *Chemical Kinetics and Photochemical Data for Use in Atmospheric Studies*, Evaluation No. 19, JPL Publication 19-5, Jet Propulsion Laboratory, Pasadena, 2019, <http://jpldataeval.jpl.nasa.gov>.
- 100 V. L. Orkin, V. G. Khamaganov, L. E. Martynova and M. J. Kurylo, High-Accuracy Measurements of OH' Reaction Rate Constants and IR and UV Absorption Spectra: Ethanol and Partially Fluorinated Ethyl Alcohols, *J. Phys. Chem. A*, 2011, **115**, 8656–8668.
- 101 R. J. Shannon, M. A. Blitz, A. Goddard and D. E. Heard, Accelerated Chemistry in the Reaction Between the Hydroxyl Radical and Methanol at Interstellar Temperatures Facilitated by Tunnelling, *Nat. Chem.*, 2013, **5**, 745.
- 102 P. Seal, G. A. Oyedepo and D. G. Truhlar, Kinetics of the Hydrogen Atom Abstraction Reactions from 1-Butanol by Hydroxyl Radical: Theory Matches Experiment and More, *J. Phys. Chem. A*, 2013, **117**, 275–282.
- 103 J. Zheng, R. Meana-Pañeda and D. G. Truhlar, Prediction of Experimentally Unavailable Product Branching Ratios for Biofuel Combustion: The Role of Anharmonicity in the Reaction of Isobutanol with OH, *J. Am. Chem. Soc.*, 2014, **136**, 5150–5160.
- 104 J. Zheng, G. A. Oyedepo and D. G. Truhlar, Kinetics of the Hydrogen Abstraction Reaction From 2-Butanol by OH Radical, *J. Phys. Chem. A*, 2015, **119**, 12182–12192.
- 105 L. P. Viegas and A. J. C. Varandas, Coupled-Cluster Reaction Barriers of HO₂ + H₂O + O₃: An Application of the Coupled-Cluster//Kohn-Sham Density Functional Theory Model Chemistry, *J. Comput. Chem.*, 2014, **35**, 507–517.
- 106 J. Tellinghuisen, Statistical Error Propagation, *J. Phys. Chem. A*, 2001, **105**, 3917–3921.

