Database for the kinetics of the gas-phase atmospheric reactions of organic compounds

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Abstract. We present a digital, freely available, searchable and evaluated compilation of rate coefficients for the gas-phase reactions of organic compounds with OH, Cl and NO₃ radicals and with O₃. Although other compilations of much of these data exist, many are out-of-date, most have limited scope, and all are difficult to search and to load completely into a digitized form. This compilation uses results of previous reviews, though many recommendations are updated to incorporate new or omitted data or address errors, and includes recommendations on many reactions that have not been reviewed previously. The database, which incorporates over 50 years of measurements, consists of a total of 2765 recommended bimolecular rate coefficients for the reactions of 1357 organic substances with OH, 709 with Cl, 310 with O₃, and 389 with NO₃, and is much larger than previous compilations. A large variety of functional groups is present in this database, including naturally occurring chemicals formed in or emitted to the atmosphere and anthropogenic compounds such as halocarbons and their degradation products. Recommendations were made for rate coefficients at 298K and, where possible, the temperature dependences over the entire range of the available data. The primary motivation behind this project has been to provide a large and thoroughly evaluated training dataset for the development of structure-activity relationships (SARs), whose reliability depends fundamentally upon the availability of high-quality experimental data. However, there are other potential applications of this work, such as research related to atmospheric lifetimes and fates of organic compounds, or modelling gas-phase reactions of organics in various environments. This database is freely accessible at https://doi.org/10.25326/36 (McGillen et al., 2019).
1 Introduction

The composition of realistic atmospheric and combustion chemical mixtures can be forbiddingly complex, as has recently been emphasized by the advent of automated mechanism generation software (Aumont et al., 2005; Battin-Leclerc et al., 2011; Carter, 2017; Gao et al., 2016). Such complexity presents a major challenge for chemical modellers, since the physical and chemical properties of the vast majority of oxidation products of volatile organic compounds (VOCs) have not been determined experimentally. For example, in the GECKO-A model, the number of possible products formed from a single VOC of only intermediate complexity, $\alpha$-pinene, can result in $>400,000$ different species (Aumont et al., 2005). To model the impact of these chemicals on air quality and climate change, information about their chemical and physical behaviour needs to be available. Given the time, expense and difficulty of making laboratory measurements, it is clear that with current technologies it will be necessary to estimate or compute the properties of almost all of these compounds. To help address this challenge an expert panel on structure-activity relationship (SAR) evaluation was formed in 2017. This panel has identified several current challenges in atmospheric chemical modelling, which are described in Vereecken et al. (2018). High among the priorities of this group is the assessment of structure-activity relationships for predicting the atmospheric reactivity of VOCs. In this regard, to test the performance of a SAR, it is necessary to compare estimated reaction rate coefficients with available experimental data. The compilation of such data is thus an essential first step in this process, and is the focus of the current work.

Compendia of kinetic data already exist. Notable among these are the thoroughly evaluated datasets provided by the IUPAC Task Group on Atmospheric Chemical Kinetic Data Evaluation (Anmann et al., 2013; Atkinson et al., 2004, 2006, 2007, 2008; Crowley et al., 2010, 2013; IUPAC, 2019), the JPL Panel for Data Evaluation (Burkholder et al., 2015), and the Calvert et al. reviews (Calvert et al., 2000, 2002, 2008, 2011, 2015). The NIST Chemical Kinetics database is an extensive compilation of kinetic data (Manion et al., 2015), which, although it is not evaluated, possesses an extremely large scope. Despite the many useful aspects of these resources, they have several drawbacks such as:

1.) The frequency with which kinetic data are published is much faster than that with which these data are compiled and reviewed. In the case of IUPAC and JPL, review cycles tend to be ~3–4 years.

2.) The number of reaction rates that have been evaluated is considerably smaller than the number that have been determined. In the case of IUPAC and JPL, vastly so. It is therefore inevitable that such evaluations do not currently capture the full chemical diversity available in the experimental literature.

3.) The data within these reviews are currently not downloadable in a digital, searchable format.

4.) Non-downloadable databases that cannot be accessed offline are subject to downtime (e.g. NIST was recently out of commission for 38 days as a consequence of the 2018–19 US federal government shutdown). Also, changes can be made to such databases, which are not necessarily traceable.

The database described here (McGillen et al., 2019) aims to overcome these drawbacks by accounting for the information contained within these previous evaluations, and simultaneously, to augment them by making a new and thorough survey of the chemical kinetics literature. As can be seen in Figure 1, the size of the database is considerably larger than previous
evaluation projects, which is a consequence of merging each of the available evaluations; new measurements becoming available; and the inclusion of measurements that were overlooked previously.

In addition to this increase in scope, we plan to periodically update this database as new data become available. This is being carried out as part of the activities of the abovementioned SAR evaluation panel (Vereecken et al., 2018). It is intended that this new database will be more agile than these previous efforts, and will adopt the Earth System Science Data “living data” approach to incorporate new kinetic data that become available, and to improve the treatment and description of data herein, where necessary. The database can be downloaded in its entirety in the form of an Excel spreadsheet. Our goal is to provide a comprehensive database to serve both as a useful reference source for the kinetics community, and as a sound basis upon which to develop SARs for use in atmospheric chemistry and other models.

2 Scientific background

The reactions of oxidants with organic compounds considered in this compilation can be either bimolecular or termolecular. Bimolecular reactions involve the interaction of two molecules, or an atom and a molecule in the case of the chlorine atom reactions. In termolecular reactions, an excited intermediate is formed, which can be stabilized by collisions with a third body, otherwise decomposition may occur, re-forming reactants. In practice, almost all of the reactions in this compilation are in their high pressure limit within the pressure and temperature range of interest to atmospheric chemistry, and these reaction rates are therefore readily described as bimolecular reactions.

The rate of a bimolecular chemical reaction is defined in terms of the rate of change of the concentration of reactants or products. The rate coefficient (sometimes referred to as a rate constant) is denoted by the symbol “k” and is the constant of proportionality relating the rate of the reaction to the concentration of the reactants. As an example, for the unit stoichiometry reaction \( \text{OH} + \text{CH}_4 \rightarrow \text{CH}_3 + \text{H}_2\text{O} \), the rate of reaction is the rate of loss of OH radicals or CH\(_4\), or the rate of formation of CH\(_3\) radicals or H\(_2\)O according to Eq. (1):

\[
\text{Rate} = k[\text{OH}][\text{CH}_4] = -\frac{d[\text{OH}]}{dt} = \frac{d[\text{CH}_3]}{dt} = \frac{d[\text{CH}_4]}{dt} = \frac{d[\text{H}_2\text{O}]}{dt},
\]

(1)

The units of bimolecular rate coefficients in this database are cm\(^3\) molecule\(^{-1}\) s\(^{-1}\), which are those preferred by the atmospheric chemical kinetics community [e.g. (Finlayson-Pitts and Pitts, 2000; Seinfeld and Pandis, 2016)]. Note that “molecule” is not a unit, but is typically included for clarity. Where rate coefficients have been reported in units of dm\(^3\) mol\(^{-1}\) s\(^{-1}\) these have been multiplied by 1.66 x 10\(^{-21}\) to convert to units of cm\(^3\) molecule\(^{-1}\) s\(^{-1}\).

Rate coefficients are measured using either absolute or relative rate methods. In absolute measurements, the rate coefficient is determined directly by monitoring the change in concentration of, at least, one of the reactants as a function of time. Typically, experiments are conducted where the pseudo-first-order decay of one reactant is measured under conditions where the other reactant is in excess, such that the concentration of the excess reactant does not change appreciably over time. As an example, flash photolysis can be used to generate OH radicals in the presence of a large excess of CH\(_4\). The pseudo-first-order loss of
OH can be probed using a variety of techniques such as resonance fluorescence, resonance absorption, or laser-induced fluorescence. The *pseudo*-first-order loss rate, $k' = -d\langle n[OH]\rangle/dt$, is related to the bimolecular rate coefficient $k$ by the expression $k' = k[CH_4]$. Experiments are conducted using different $CH_4$ concentrations and a plot of $k'$ versus $[CH_4]$ has a slope equal to $k$.

In relative rate studies, the rate of the reaction of interest is measured relative to that of a reference reaction whose rate coefficient has been determined by an absolute technique. Having established the rate coefficient of $OH + CH_4$ by an absolute method, this reaction can be used as a reference to measure the rate coefficients for reactions of $OH$ radicals with other organic compounds. As an example, the reactions of $OH$ radicals with a VOC can be studied by exposing mixtures containing the VOC and a reference to $OH$ radicals. The reactant and reference compounds are monitored using conventional analytical techniques such as gas chromatography or FTIR spectroscopy. A plot of $\ln([VOC]_t/[VOC]_0)$ versus $\ln([reference]_t/[reference]_0)$ has a slope equal to the rate coefficient ratio $k_{VOC}/k_{reference}$, where $t_0$ and $t$ refer to initial concentrations and concentrations at time $t$ respectively.

In absolute studies, the reaction time must be measured accurately, otherwise systematic errors will be introduced. Furthermore, careful attention must be paid to the reactant purity, where, depending on the relative reactivity of an impurity, even a small fraction ($<0.001$) in a sample can affect the retrieved rate coefficient adversely. In some systems, the absolute method is sensitive to regeneration of reactants (e.g. $OH$ recycling), and it is necessary to perform tests to establish that this is not affecting the phenomenological rate coefficient. Conversely, in relative studies, conditions must be selected such that the reactant and reference are lost only by the reaction of interest and that neither reactant, nor reference, are re-formed in any process. One of the difficulties associated with many relative rate studies is that they are conducted under chamber conditions, where reactive intermediates from either the VOC of interest or the reference may be present. Therefore, it is often desirable to conduct several measurements in the presence of different reference compounds. It is also better that reaction rates between reference and compounds of interest are not too dissimilar, such that a sufficient amount of chemical conversion is achieved for each in an overlapping timeframe. Absolute rate techniques are generally capable of higher accuracy than relative rate methods, once the uncertainties in the reference reaction are considered. Relative rate techniques are generally simpler to implement and capable of higher precision than absolute rate techniques. Absolute and relative rate methods are complementary and have been used together to provide the wealth of kinetic information documented in this compilation.

The temperature dependence of bimolecular reactions over limited temperature ranges can usually be described by the Arrhenius equation, Eq. (2):

$$k(T) = A\exp\left(-\frac{B}{T}\right),$$

where $B = E/R$. 

The pre-exponential $A$-factor represents the rate of molecular collisions with the correct orientation for reaction and the exponential term is the fraction of those collisions with sufficient energy for reaction to occur. Over extended temperature ranges many reactions exhibit curved Arrhenius plots because of the importance of multiple reaction pathways each with different temperature dependencies, formation of pre-reactive complexes, and quantum tunnelling at low temperatures, among
several other reasons why curvature is expected (Gardiner, 1977). Where rate coefficients have been determined over large temperature ranges and curvature has been observed in Arrhenius-space, we have expressed temperature dependences using a 3-parameter equation that is sometimes referred to as Kooij’s equation (Laidler, 1984), and is referred to here as the “extended” Arrhenius expression shown in Eq. (3):

$$k(T) = A \exp\left(-\frac{B}{T}\right)\left(\frac{T}{300}\right)^n.$$  

Here, an additional term, \((T/300)^n\), is added to account for the curvature, where "n" is an additional parameter adjusted to fit the data, along with "A" and "B". Note that this is the same as the standard Arrhenius expression when \(n = 0\). We use the \((T/300)^n\) parameterization rather than the simpler \(T^n\) because this allows "n" to be dimensionless and the units of \(A\) to be independent of \(n\), and of comparable magnitude to the \(A\) parameter in the standard expression (see Eq. 2). Although this parameterization is arbitrary and \(A\), \(A\), and \(B\) do not have any clear physical or chemical meaning (Carvalho-Silva et al., 2019), it works well in fitting kinetic data for most compounds over wide temperature ranges with only one additional parameter. This is shown, for example, in Figure 2, which gives an Arrhenius plot for rate coefficient measurements for the reaction of OH with dimethyl ether over a temperature range of 195–1470 K. The dashed blue line is the standard Arrhenius expression derived from the data for 230–300K, \(k(T)=5.7\times10^{-12}\ \exp(-215/T)\ \text{cm}^3\ \text{molecule}^{-1}\ \text{s}^{-1}\), while the solid line shows the extended expression, \(k(T)=1.02\times10^{-12}\ \left(T/300\right)^{0.09}\ \exp(308/T)\ \text{cm}^3\ \text{molecule}^{-1}\ \text{s}^{-1}\), which fits the data over the full range of 195–1470 K. It can be observed that, whether the measurement is absolute or relative, that the vast majority of kinetic studies provide total phenomenological rate coefficients rather than site-specific data. In principle, any rate coefficient that contains several non-degenerate reactive site can be expressed in the following form, for \(i\) number of reactive sites, see Eq. (4):

$$k_{\text{total}}(T) = \sum_i A_i \exp\left(-\frac{B_i}{T}\right)\left(\frac{T}{300}\right)^{n_i},$$  

It follows that where data on the branching ratios between these reactive sites are absent, then the kinetic information encoded within the total rate coefficient is incomplete. Unfortunately, this is the general state of affairs for the vast majority of reactions contained within the literature, and hence no attempt is made within the framework of the current version of the database to describe branching ratios.

There are 800 rate coefficients in the current database where we have recommended extended Arrhenius parameters, based either on acceptance of recommendations from previous evaluations, or our own refitting of the data. For practical reasons, we did not re-fit data unless we judged that the existing recommendations did not encompass all measurements (which may include recently published data) or had other problems. This was done for ~50 rate coefficients, all of which were for OH reactions. For the remaining ~2000 reactions, recommendations were provided for rate coefficients around 298K, because of lack of data at other temperatures, or (in a few very cases) because the temperature dependence around 298K could not be described by the extended Arrhenius expression above. The current database does not contain rate coefficients at other temperatures, other than what could be computed by our recommended temperature-dependent expressions within the stated temperature range.
Reactions occurring on essentially every collision have a rate coefficient known as the gas kinetic limit, which is approximately $5.0 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ at 298 K, although its precise value will vary with the structure of the reactants. The recommended rate coefficients at 298 K in the present compilation span the range from the gas kinetic limit for the reactions of chlorine atoms with long-chain hydrocarbons, to less than approximately $10^{-22}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ for the reaction of ozone with halogenated alkenes which corresponds to reaction in approximately 1 out of $10^{11}$–$10^{12}$ collisions.

3 Methods

The reviews of IUPAC, JPL and Calvert et al. (Ammann et al., 2013; Atkinson et al., 2004, 2006, 2007; Burkholder et al., 2015; Calvert et al., 2000, 2002, 2008, 2011, 2015; Crowley et al., 2010, 2013; IUPAC, 2019) constituted the starting point of our data compilation effort. Each recommendation provided in these reviews was transcribed into our own database. Where overlap existed between reviews, recommendations were generally consistent, which provided an opportunity to check for errors in transcription or errors in the reviews. Errors identified in published reviews were excluded from our dataset. Following this initial phase of compilation, kinetic data published between (2015 and present) were compiled by searching keywords in Google Scholar over these years and transcribing data from the original publications. Whereas any data published after 2015 cannot be contained in JPL Evaluation Number 18 and Calvert et al. (2015), IUPAC can be more up-to-date owing to the more localized update cycle of this review body. The NIST kinetic database was also interrogated to find data that is contained within their extensive database that is not contained within the reviews that we considered, although, at the time of writing, it is noted that this database also does not appear to contain updates since 2015. Following this review of available literature, further, more general searches of kinetic publications were made for all years, which would be able to locate data that had been overlooked by the extensive reviews of Calvert et al. or the large NIST database.

Once all data known to this study were compiled, reviews for individual reactions were made. There are several possible outcomes from entering data into the database and these are described in Figure 3.

Some of the decisions in this review process are easy to arrive at objectively, such as whether or not all measurements are consistent, which can be determined by a simple comparison. However, other decisions are more nuanced, such as whether or not a measurement is trustworthy. In this instance many factors can influence this decision, including:

- Is the measurement technically difficult?
- How well was the measurement performed?
- Were appropriate tests made?
- Is the apparatus suited to measuring this reaction?
- Is the measurement generally consistent with analogous reactions?

Because each of these questions requires considerable experience and judgement to answer, the review process was conducted in duplicate and occasionally triplicate, such that if discrepancies between individual reviewers emerged, these discrepancies were discussed and resolved prior to a final review being accepted by the panel.
Where temperature dependence was available for a reaction, the review process can require further decisions to be made. Firstly, if temperature dependence was determined in some measurements but not others, then A-factors were normalized to all available data and the activation energy, $E$, was taken from an individual study, or an average of several studies if more than one determination was available. In some instances, where general agreement was observed in A-factors, but major differences in activation energies were reported, we chose not to recommend temperature-dependent parameters. Secondly, where there are several temperature-dependent studies that span a large range in temperature, and where the temperature dependence can be described adequately by the extended Arrhenius equation (see Eq. 3), an error-weighted linear least squares fit was performed on the entire dataset, and the resultant expression constitutes our recommendation, as shown in Figure 2, for example. Finally, if temperature dependence information is available but all data are at temperatures higher than 298K, then extrapolation is necessary to estimate the rate coefficient at 298K. If the extrapolation is sufficiently close, i.e., causes the rate coefficient to change by less than a factor of 2 compared to that calculated for the lowest temperature in the measurement range, then the extrapolated 298K rate coefficient is recommended but with an increased uncertainty assignment. We give no 298K rate coefficient recommendation if the change is greater than that, though the extrapolated rate coefficient is provided as an estimate in the comments. This approach is pragmatic, and more exhaustive treatments are possible, we therefore list this as one of the items of ongoing work listed in Section 6.

The structure of the current database, and the information it contains, is summarized in the instruction manual and supplementary information document that is provided with the database file. The database file is an Excel spreadsheet with tables containing the data and also with worksheets giving information about the database, worksheets and macros to search and extract information from the database. The database itself consists of tables giving information about the compounds used, the kinetic data, the references cited, and a table of compound names and other identifiers. The "Compounds" table gives structural information about the compounds, codes indicating the types of compounds that may be useful for lookup purposes, and the recommended kinetic parameters for the four types of reactions that are currently considered (OH, NO$_3$, O$_3$, and Cl). Most of the compounds have more than one name or identifier that can be used for lookup purposes, and those that can be used for this application are given in the "Names DB" table. The kinetics data are given in two tables: the "k-Data" table gives the 298K rate coefficient and temperature dependence parameters from the various reviews or primary studies, while the "kT-Data" table gives the temperatures and rate coefficients that were used for manual fitting. In both cases, codes giving the references used are provided alongside the kinetic data. The reference citations and (where available) URLs for the various references are given in the "References" table.

The chemical identifiers used included commonly used names that were taken from the original publications or the NCI database (National Cancer Institute, 2010), and other identifiers such as CAS registry numbers when available. Unique textual identifiers (canonical SMILES, InChI and InChiKey) were also included, making this database easily searchable, such that kinetic information can be obtained rapidly without knowledge of how the molecule is named within the database. There are many chemical informatics software packages and resources that are available for generating SMILES and InChI codes, both freeware e.g. (ACD/ChemSketch, 2018) and commercial software packages e.g. ChemDraw and online services such as the
NCI/CADD Online SMILES Translator (National Cancer Institute, 2017). Note that SMILES strings are not strictly unique and may be dependent upon the algorithm used in a given software implementation, therefore all SMILES were provided in their canonical form as output using the open-source Open Babel software program (O’Boyle et al., 2011). Furthermore, other user-specified differences to SMILES output can still occur, even in canonical form, an important example being the representation of nitrogen–oxygen bonds, where we prefer to represent these bonds as dative. SMILES strings can be easily converted to canonical SMILES using this package, and InChI and InChIKey using this and other open-source/proprietary resources. Less specific identifiers, including molecular formula, molecular mass and types of compounds and functional groups are also provided, and these can be used to make broader searches to the database possible.

### 4 Results

Table 1 is a summary of the number of compounds, reactions, and rate coefficient recommendations in our database, together with the number of functional groups contained within each molecule. As shown above, Figure 1 provides an overview of the size of the current database in relation to existing compilations of data, and Figure 4 shows the temperature range covered by our data. From Figure 4 and Table 1 it is clear that there is a large majority of data that is available at room temperature only or within the range of 250–370 K, which coincides with the general temperature limitations of ambient chamber measurements and jacketed flow reactors respectively. It is also notable that the OH radical dataset possesses the largest number of reactions and the largest fraction of temperature-dependent measurements. By contrast, the number of compounds measured for the NO$_3$ radical is much smaller, and the fraction of temperature-dependent measurements is also much less than for OH.

Regarding the functional form that is used to describe temperature dependences, where obvious curvature can be observed, as shown in Figure 2 for example, the extended Arrhenius expression is preferred, since this describes the data more faithfully. Ultimately, although most reactions in the OH dataset are expected to be non-Arrhenius, most reactions have yet to be studied over a sufficient temperature range with enough precision to require the third ("n") parameter of the modified Arrhenius expressions to fit the data, so Arrhenius equations constitute the majority of temperature-dependent expressions in the dataset. By contrast, for reactions such as alkene ozonolysis, where quantum tunnelling is not expected to be feasible, any curvature in Arrhenius-space is likely to be small and so far no ozonolysis reactions known to this study have been shown to exhibit non-Arrhenius behaviour.

As shown in Table 1, of the 1644 compounds studied so far, most reactions have been measured for species that contain two or fewer functional groups. Generally, as the number of functional groups increases in a molecule, the boiling point increases and the saturation vapour pressure decreases, making measurements more challenging in the gas phase, which explains why there are very few measurements on compounds with 5 or more functional groups. Conversely, for those compounds with no functional groups – defined as a compound that possesses no atom type besides carbon and hydrogen and no higher-order bonds (i.e., alkanes) – the relatively small number of these compounds relates to the fact that there are fewer possible isomers available within the range of volatility that is convenient for experimentation.
5 Discussion

As shown in Figure 1, the database presented in this work is substantially larger than previous compendia, and reflects our attempts to compile all available data concerning gas-phase reactions of organic compounds with selected atmospheric oxidants under atmospheric conditions. The current database provides recommendations for the reactions of VOCs with OH and NO$_3$ radicals, O$_3$ and Cl atoms, the major oxidants that react with organic compounds in the atmosphere. Rate coefficients for the reactions of VOCs with other oxidants can be added in later versions of the dataset if there is sufficient interest. However, the focus of the development of the current database is to support the needs of assessing and modelling the impacts of organic compounds in the atmosphere.

For this objective, the ideal is to present rate coefficients for every compound that is emitted into the atmosphere, and for every oxidized organic compound that is formed and reacts in the atmosphere. Knowledge of rate coefficients of emitted compounds is necessary to assess their atmospheric lifetimes and the impacts of their atmospheric reactions on air quality. A total of ~1700 individual compounds have been identified or estimated to be present in the various chemical categories used in U.S. emissions profiles, of which ~1000 compounds are present in VOC mixtures derived to represent total U.S., California and Texas anthropogenic emissions (Carter, 2015). This database provides rate coefficient assignments for at least the OH reaction for ~90% of the mass, though only ~40% by number of compounds with non-zero emissions. The high coverage in terms of mass emissions is expected, since such compounds are most likely to be a priority for research. However, a very large number of other species are emitted, and although individually these may be insignificant, they may become important in the aggregate, and should be of interest at least to those who use or emit such compounds. Therefore, it is reasonable to expect that more of these substances will be studied in the future.

Knowledge of the rate coefficients of the oxidized products formed when VOCs react in the atmosphere is necessary to determining the ultimate environmental fates of the emitted compounds and modelling their overall impacts on air quality. However, there are large numbers of possible reactions that many organic compounds and their reactive intermediates can undergo, and the use of automated mechanism generation systems such as GECKO-A (Aumont et al., 2005) is necessary to derive complete mechanisms. Complete coverage of experimental rate coefficients of such a large number of oxidation products is currently unfeasible, and the best that can be hoped for in this regard is to provide rate coefficients for compounds with a variety of representative structures, chemical functionalities, and combinations of functionalities, which may serve as a basis for developing SARs or other methods to estimate rate coefficients for this large array of species.

To obtain an approximate indication of the types of compounds predicted to be formed by mechanism generation systems, and to assess the coverage of this database concerning their rate coefficients, we used GECKO-A to derive complete mechanisms for the atmospheric reactions of the representative compounds n-octane and α-pinene, which are associated with anthropogenic and biogenic activities respectively, and which are expected to yield distinctly different product distributions. The results of this comparison are shown in Figures 5 and 6.
Figure 5 shows the overlap between the current database and the \( n \)-octane and \( \alpha \)-pinene products predicted by GECKO-A in terms of individual organic product species. The area of each curve in this diagram is proportional to the number of species contained within it. It is clear from this comparison that the number of species that have been studied so far is very small compared with the total number of species produced in the oxidation of these quite structurally simple primary emissions. Furthermore, the overlap between the species studied and the GECKO-A output is vanishingly small. Under the proviso that the GECKO-A mechanism is representative of the state-of-the-knowledge in atmospheric chemistry, species that occupy this overlap region can be regarded as the “known knowns” of atmospheric chemistry, i.e. those species that are known to be produced and have known rate coefficients. When known primary emissions are subjected to the rules of atmospheric chemistry known to GECKO-A, those species that don’t overlap with our database are considered as “known unknowns”. The area that falls outside these curves is expected to be vast, and relates to all species that are formed from all primary emissions that do not overlap with the product distribution of \( \alpha \)-pinene, \( n \)-octane or our database. We consider this area to represent “unknown unknowns” in atmospheric chemistry. By this definition, the size of this area cannot be known, but it is anticipated that it is very large, especially when all known and unknown primary emissions are included, and when it is acknowledged that there may be many unusual or exceptional product formation pathways that are currently unknown to the GECKO-A model. It is possible that this representation of the state-of-the-knowledge in atmospheric chemistry may be unduly pessimistic, in that these model runs present information on the total number of species, but do not account for product fluxes, which could be very small for any species that are produced in rare events.

Notwithstanding, even if the overall flux to the atmosphere was low for a large number of these species, it appears reasonable to expect that given the sheer number of species present in the atmosphere (Goldstein and Galbally, 2007), primarily emitted or produced through oxidation, that the fraction of species for which kinetic measurements are available will remain miniscule. This observation is underscored by the fraction of species for which measurements are available over the complete atmospherically relevant temperature range, the bulk of which will be consumed in troposphere, which experiences temperatures between 220 and 300 K (see Figure 4). This means that almost all the organic product rate coefficients used by mechanism generation systems like GECKO-A are dependent upon estimation techniques.

From Figure 6, it is evident that certain functional groups that are relatively uncommon among atmospheric oxidation products of hydrocarbons (e.g. ethers and esters), are well represented in our database, and yet there are many functional groups that are expected to be commonplace that are very much underrepresented (e.g. nitrates, hydroperoxides, peroxyacids, carboxylic acids and PANs). Furthermore, the number of functional groups contained within a molecule is generally smaller in our database (typically between 2 and 3 functional groups per molecule) compared with the molecules produced in GECKO-A, where the modal distribution ranges between ~3 and 7 functional groups per molecule. The reasons for these disparities are easily rationalized. For example, many of the functional groups that are poorly represented are thermally unstable, and compounds with these functional groups are difficult to purchase, synthesize, store, and handle in experimental studies. Other functional groups such as the carboxylic acids, are stable but they suppress vapour pressure to such an extent that only the most volatile members of this family have rate coefficient measurements. Similarly, it is well known that increasing the number
of oxygenated functional groups within a molecule reduces the vapour pressure profoundly, and it is therefore often impractical to perform measurements upon highly functionalized species in the gas phase with current technologies and experimental approaches.

As shown in Figure 6, the situation is more optimistic regarding primary emissions from anthropogenic sources, where industrially important compounds such as ethers, esters and alcohols are reasonably well represented. Furthermore, the distribution of the number of functional groups per molecule also suggests good overlap. However, it is generally the case that oxidation in the atmosphere will be the predominant fate of each of these primary emissions, and such oxidation will lead to further functionalization. Therefore, as with the example of \( n \)-octane and \( \alpha \)-pinene oxidation in GECKO-A, it is expected that these primary emissions will generate an immense number of daughter products under atmospheric conditions.

With such a large number of unknown rate coefficients, it is vital that accurate and computationally inexpensive methods such as SARs for estimating rate coefficients are available so that explicit models such as GECKO-A can be employed to make accurate representations of atmospheric chemistry. Although it is anticipated that in-depth analyses of SAR performance will be forthcoming from our expert panel in the future, one well-established method of estimating rate coefficients that arises naturally from the compilation of data presented in our database is that of the free-energy relationships exhibited by rate coefficients of VOCs between different oxidants. In Figure 7, several such relationships are presented. It is clear that some of these relationships are stronger than others. For example, the correlations of ozone against both hydroxyl and chlorine are relatively high, which has been observed previously in the case of \( O_3 \) and OH (McGillen et al., 2011). In this example, the mechanism of all reactions in these relationships is electrophilic addition. Conversely, other relationships within this diagram involve a combination of addition and abstraction reactions (e.g. any correlations between OH, Cl and NO\(_3\)). Furthermore, some reactions may be more affected by steric hindrance (e.g. ozonolysis) than others (McGillen et al., 2008). Consequently, several trends arise depending on the relative efficiencies with which an oxidant participates in a given mechanism. Therefore, when taken as a whole, such correlations appear surprisingly scattered, although it is noted that individual subsets of these correlations may have good predictive power. As has been observed in the OH–Cl free-energy relationship for halocarbons and ethers for example (Sulbæk Andersen et al., 2005).

6 Ongoing work and outlook

The work contained in the present database represents clear progress in terms of its comprehensive coverage, availability and accuracy, and the fact that it can be downloaded and readily searched. However, limitations remain and the following future improvements can be envisioned:

1.) There are several oxidants that are of importance to combustion chemistry and there are some atmospheric or laboratory conditions that are not currently included, e.g. \( O(3P) \), \( O(1D) \), carbonyl oxides, H and Br atoms, low-temperature OH reactions.
2.) Quantitative information on branching ratios for sites of attack is available for certain reactions, which is not yet implemented in the current database.

3.) There is at present only a limited amount of metadata based on experimental conditions, but no information on technique/ reactor details/ pressure/ bath gas/ reference compounds in relative rate experiments.

4.) There is a wealth of information published on kinetics in the solution phase that is beyond the scope of the current database, which focuses purely on gas-phase reactions.

5.) The current approach to extrapolation of rate coefficients using temperature-dependent data outside 298K is not statistically rigorous. Improvements will require further data analysis such as that outlined in Hites (2017).

6.) Similarly, uncertainty estimates that are ≥100% are not physically meaningful. Improvements upon this may require the asymmetrical distribution of errors afforded by the approach of the IUPAC task group. Again, further statistical analyses will be necessary.

The timescales over which such improvements can be made is likely to be determinant upon external factors such as funding, the continued participation of members of the expert panel and the possible participation of other experts. However, work will continue on several of these aspects in anticipation of future versions of this database.

7 Data availability
The current version of this database, together with instructions on how to use it are freely available at the following URL: https://doi.org/10.25326/36

8 Conclusions
We present a digital, freely available, searchable and evaluated compilation of chemical kinetic information with a current focus on gas-phase bimolecular reactions. This database responds to a perceived need within the atmospheric chemistry community and elsewhere for an up-to-date, reviewed database that captures the chemical diversity that is found within the kinetics literature. It is intended that this will be a valuable resource for research into SARs, among other applications, where the quality of training sets will impact accuracy and predictiveness directly. Experimentalists will also be able to use this database to compare their measurements with previous data and analogous compounds, and will also be able to easily locate evaluated reference rate coefficients. Although the current version of this database is the largest database of its kind, there remains much kinetic data that are currently not included in this project, including reactions with several important oxidants; reaction branching ratios; and reactions in other phases besides the gas phase. This, together with the fact that new rate coefficients are published each year, means that further work will be necessary to both improve, extend and maintain this database.
Acknowledgements

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References


Figure 1: Area-proportional Venn diagrams constructed using the eulerAPE software (Micallef and Rodgers, 2014). The size and position of each curve is proportional to the number of species studied with respect to each oxidant, the number of compounds available in each review, and the overlap between these reviews.
Figure 2: Temperature dependence of the reaction of OH with dimethyl ether. As with many other reactions, curvature in Arrhenius-space is observed over sufficiently large temperature ranges, especially in systems where quantum tunnelling, pre-reactive complexes and multiple reaction channels are active. This highlights the need to use the modified Arrhenius expression for some of the reactions in this database.
Figure 3: Decision tree describing the data review process for our database.
Figure 4: Histograms showing the number of rate coefficients at each temperature, separated by oxidant. A large number of reactions have been studied only at room temperature, especially for some oxidants such as NO$_3$, where temperature dependent data is lacking.
Figure 5: Area-proportional Venn diagram showing the overlap between species formed in GECKO-A for n-octane and α-pinene oxidation, and species present in our database. Here, “known knowns” reflect compounds that are formed in GECKO-A for which measurements are available. “Known unknowns” represent chemicals that are formed in GECKO-A for which no measurements are available. “Unknown unknowns” represent species that could be formed from the oxidation of other primary emissions besides n-octane and α-pinene that are not considered in this diagram, and may also represent compounds that are formed through mechanisms that are currently unknown to/ not considered in GECKO-A.
Figure 6: Frequency plots comparing the functionalization of compounds within the GECKO-A mechanism of $n$-octane and $\alpha$-pinene oxidation; a database of anthropogenic emissions (Carter, 2015); and the compounds present in our kinetic database. (Left) Functional groups such as ethers and esters are overrepresented within this database compared with GECKO-A, whereas other functional groups (e.g. nitrates, PANs and hydroperoxides) are very poorly represented. (Right) a mismatch is demonstrated between the number of functional groups per molecule in GECKO-A and that of the compounds found in this database. Better agreement is observed in both cases compared with primary emissions profiles.
Figure 7: Free-energy relationships between the reaction rates of the various oxidants within this database. Since it is possible that a compound may belong to several of the groupings shown in the legend, categorization of these compounds has been prioritized by reactivity (e.g. an alkene that is also an oxygenate is described as an alkene, since this is likely to be the dominant reactive site).
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<th>$k(\text{O}_3)$</th>
<th>$k(\text{NO}_3)$</th>
<th>$k(\text{Cl})$</th>
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<tr>
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<tr>
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<td>389</td>
<td>707</td>
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<td>389</td>
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Table 1: Summary of current contents of the experimental atmospheric rate coefficient database for VOCs.

[a] Numbers of compounds with rate coefficient entries for this reaction. Note that there may be more than one entry per compound because some compounds have more than one type of structural group.

[b] Each entry represents a different reference or source for a rate coefficient, but with no more than one recommended for assessments or SAR development.

[c] Where \( f = 0 \), compounds contain no substitutions besides C and H, and contain no higher order bonds, the only compounds that fit this definition are alkanes and cycloalkanes. Halocarbons which may contain many halogen substitutions are treated as one functional group, \( f = 1 \), because they tend not to behave uniformly, unlike complex multifunctional compounds.