Low-temperature combustion chemistry of novel biofuels: resonance-stabilized QOOH in the oxidation of diethyl ketone
Low-temperature combustion chemistry of novel biofuels: resonance-stabilized QOOH in the oxidation of diethyl ketone†

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The Cl* initiated oxidation reactions of diethyl ketone (DEK; 3-pentanone; (CH₃CH₂)₂C═O), 2,2,4,4-d,6-d,4-d plead ketone (d₄-DEK; (CH₃CD₂)₂C═O) and 1,1,5,5,5-d,6-d,6-d plead ketone (d₄-DEK; (CD₂CHOH)₂C═O) are studied at 8 Torr and 550–650 K using Cl₂ as a source for the pulsed-photolytic generation of Cl*. Products are monitored as a function of reaction time, mass, and photoionization energy using multiplexed photoionization mass spectrometry with tunable synchrotron radiation. Adding a large excess of O₂ to the reacting flow allows determination of products resulting from oxidation of the initial primary (Rₛ) and secondary (Rₛ) radicals formed via the Cl* + DEK reaction. Because of resonance stabilization, the secondary DEK radical (3-oxopentan-2-yl) reaction with O₂ has a shallow alkyl peroxy radical (RₛO₂) well and no energetically low-lying product channels. This leads to preferential back dissociation of RₛO₂ and a greater likelihood of consumption of Rₛ by competing radical–radical reactions. On the other hand, reaction of the primary DEK radical (3-oxopentan-1-yl) with O₂ has several accessible bimolecular product channels. Vinyl ethyl ketone is observed from HO₂-elimination from the DEK alkylperoxy radicals, and small-molecule products are identified from β-scission reactions and decomposition reactions of oxy radical secondary products. Although channels yielding OH + 3-, 4-, 5- and 6-membered ring cyclic ether products are possible in the oxidation of DEK, at the conditions of this study (8 Torr, 550–650 K) only the 5-membered ring, 2-methyltetrahydrofurran-3-one, is observed in significant quantities. Computation of relevant stationary points on the potential energy surfaces for the reactions of Rₛ and Rₛ with O₂ indicates this cyclic ether is formed via a resonance-stabilized hydroperoxyalkyl radical (QOOH) intermediate, formed from isomerization of the RₛO₂ radical.

I. Introduction

As concerns over global climate change, pollution and the stability of traditional petroleum supplies continue to intensify, finding and developing alternative sources of transportation fuels have become high environmental and national security priorities. As the most abundant biopolymer worldwide, cellulose has the potential to provide a major source of liquid fuels. Furthermore, because plants extract CO₂ from the atmosphere during photosynthesis, in principle their combustion can be part of a carbon-neutral cycle. However, the recalcitrance of plant material to either biochemical or thermochemical breakdown has proved to be a major impediment to efficient, large-scale production of cellulosic biofuels.3-4

Fungal conversion of lignocellulosic biomass is a promising new alternative to standard thermochemical and microbial decomposition methods.5-7 The relative simplicity of fungal genomes makes these organisms useful candidates for engineering specific metabolic pathways if particular products prove to be desirable. Among the products of fungal decomposition of cellulose are a variety of ketones and other oxygenates5,7,8 whose combustion chemistry and ignition behavior are not well understood. Novel fuels may be useful in advanced engine strategies that have the potential to combine greater efficiency and lower emissions when compared to spark ignition or diesel engines by employing autoignition of lean fuel-air mixtures.9 Homogeneous charge compression ignition (HCCI) is a limiting case of such strategies; ignition in an HCCI engine is controlled by gas-phase chemistry, and is sensitive to the initial oxidation steps of the fuel molecule outlined in Section II.

Detecting products from the initial steps of fuel oxidation is an important foundation in an investigation of the ignition
characteristics of an individual fuel. However, such studies also provide in-depth knowledge of the fundamental reaction mechanisms involved during ignition that will enable predictive capability across a family of fuel molecules. Specifically, ascertaining the role of functional groups in the complex reactivity of particular radicals can help identify molecular structures and chemical bonding motifs that will yield desired ignition characteristics. In this study we take diethyl ketone (DEK; 3-pentanone) as a representative molecule for the family of open-chain ketones that could be harnessed from conversion of lignocellulosic feedstocks. DEK has also been investigated recently\(^{10}\) as a prototype for oxidation of ketones in the troposphere, which follows many of the same channels that have been mapped out in the higher-temperature environment of autoignition.\(^ {11,12}\) In the present case, partially deuterated DEK isotopologs are used to investigate the various ‘chain-propagating’ cyclic ether + OH formation pathways from reactions of carbonyl-substituted alkyl radicals with O\(_2\). These products are formed via isomerization of the initial peroxy (RO\(_2\)) radicals to hydroperoxyalkyl (QOOH) radicals that subsequently lose OH. We show that vinylic resonance stabilization in the QOOH is critical in directing the outcome of the reaction, and discuss possible implications for more general description of ketone autoignition.

II. Background

In the last decades investigations of fundamental oxidation chemistry have yielded a detailed mechanism for reaction pathways that govern alkane autoignition.\(^ {11,13}\) A simplified general mechanism is given in Scheme 1 for an initial alkyl radical (R) reacting with O\(_2\). Formation of an alkyl peroxy radical (RO\(_2\)) typically proceeds without an energy barrier.\(^ {14,15}\) Of particular note in Scheme 1 is the formation of hydroperoxyalkyl radicals (QOOH) via internal hydrogen atom transfer from RO\(_2\). The unimolecular decomposition of QOOH can lead to chain-propagating formation of OH, often associated with formation of a stable cyclic ether coproduct.\(^ {11,16}\) Alternatively, these QOOH species can promote chain-branching via addition of a second O\(_2\) molecule and subsequent decomposition to two OH radicals and an ox radical.\(^ {17}\) QOOH radicals have, however, never been directly detected, so observation of the stable cyclic ether products is a key indication of QOOH chemistry for any particular fuel.

In competition with QOOH formation, RO\(_2\) can eliminate HO\(_2\), a much less reactive species than OH. In alkane oxidation, the HO\(_2\)-elimination channel generates an alkene by removal of an H atom initially bound to a carbon neighboring the peroxy group.\(^ {14,15,16,19}\) Due to the relatively unreactive nature of the HO\(_2\) radical, these channels are deemed ‘chain-terminating’.\(^ {11}\) Understanding the overall importance of chain-propagating and chain-branching pathways relative to the chain-terminating channels facilitates prediction of the low-temperature autoignition properties of a particular compound as a fuel in advanced engines. Though larger fuel molecules have been studied,\(^ {20}\) the majority of compounds in which these pathways have been investigated in detail are simple hydrocarbons with more tractable mechanistic pathways.

A useful method for near-instantaneous generation of a radical pool for time-resolved study of oxidation chemistry is the pulsed-photolytic generation of Cl\(^ +\) via photodissociation of Cl\(_2\). The Cl\(^ +\) then abstracts an H atom from the molecule of interest yielding R + HCl. However, secondary reactions involving both Cl\(_2\) and Cl\(^ +\) must also be characterized, especially if they yield products identical to those expected from oxidation chemistry. These undesired side and secondary reactions can be quenched by including sufficiently large concentrations of O\(_2\) and fuel in the reacting flow, although experimental considerations limit the maximum practical fuel concentration. In the case of DEK, the resonance-stabilized radicals that are not as readily consumed by O\(_2\) suffer a greater degree of secondary reactions, including chlorine chemistry. Therefore, care must be taken to identify and quantify these reactions and products so the oxidation channels can be accurately isolated and characterized.

III. Experiment

The chlorination and oxidation chemistry of DEK [(CH\(_3\)CH\(_2\))\(_2\)C\(\equiv\)O], \(d_4\)-DEK [(CH\(_3\)CD\(_2\))\(_2\)C\(\equiv\)O] and \(d_6\)-DEK [(CD\(_3\)CH\(_2\))\(_2\)C\(\equiv\)O] was studied at the Advanced Light Source (ALS) at Lawrence Berkeley National Laboratory. The experiment has been described in detail previously\(^ {21,22}\) and will only be summarized briefly here. Calibrated mass flow controllers introduce reactants and buffer gas (He) into a heated 1.05 cm i.d. quartz flow tube. The pressure in the tube is maintained at 8 Torr by feedback control of a butterfly valve in the exhaust line. The reactive mixture is continuously sampled through a \(\sim 650 \mu\)m aperture in the sidewall of the reactor. A collimated molecular beam is directed into a differentially pumped ionization chamber by passing the effluent through a 1.5 mm diameter skimmer. Quasi-continuous, tunable ionizing vacuum ultraviolet radiation provided by the ALS synchrotron intersects the molecular beam orthogonally and

![Scheme 1](image_url)  
Scheme 1: Simplified mechanism for low-temperature oxidation of a hydrocarbon radical R.
ions are accelerated and orthogonally extracted into a linear
time-of-flight mass spectrometer. Products are detected as a
function of mass, time and photoionization energy yielding a
3-dimensional dataset that can be sliced and integrated to obtain
photoionization spectra and temporal profiles for individual
species. Photolysis of Cl$_2$ by a pulsed 351 nm excimer laser
operating at 4 Hz yields the initial Cl$^+$ atom pool. Unless noted
otherwise, concentrations of DEK, Cl$_2$, Cl$^+$ and O$_2$ are given in
Table 1. A conversion of 1.5% of the Cl$_2$ to Cl$^+$ is estimated based
on the laser fluence and the photodissociation cross section of
Cl$_2$ at 351 nm.$^{23}$ Average pre-photolysis background signal is
subtracted, resulting in difference mass spectra that show
positive signal for products generated post-photolysis. Signals
are normalized to the synchrotron photon flux measured by a
calibrated photodiode.

Along with DEK, partially deuterated forms of DEK were isolated to isolate particular reaction pathways. The non-deuterated diethyl ketone was obtained commercially at a stated purity of >99%. The $d_4$- and $d_6$-diethyl ketone isotopologs were obtained commercially and assayed at 99.8% and 99.6% chemical purity, respectively. Vinyl ethyl ketone, 2-methyltetrahydrofuran-3-one and tetrahydro-4H-pyran-4-one were obtained commercially at stated purities of >95%, 98% and 99% respectively. All samples were freeze–pump–thawed to remove dissolved gases before use.

### IV. Results and discussion

Vinoxylic resonance stabilization plays a key role in the oxidation chemistry of ketones, both in the formation of the initial radicals and in the isomerization pathways of the R + O$_2$ reactions. Chlorine-atom initiation of diethyl ketone oxidation forms both non-resonance-stabilized radicals at the primary carbons (R$_p$) as well as resonance-stabilized secondary radicals (R$_s$) (Scheme 2) for which the equilibrium R + O$_2$ ↔ RO$_2$ lies further to the left. This shift increases the likelihood that the secondary radicals will undergo other reactions besides oxidation, for example radical–radical reactions or chlorine chemistry.

Because reactions of R$_p$ with Cl$^+$ can form vinyl ethyl ketone (VEK), which is a product of HO$_2$ elimination from R$_p$O$_2$ and R$_2$O$_2$, it is important to quantify to what extent these secondary chlorine reactions influence the product spectrum under our experimental conditions. The next section describes the influence of secondary chlorine reactions, and the subsequent section presents the results from oxidative chemistry.

#### A. Chlorine reactions

In our experiments, the photolytically generated Cl$^+$ atoms should ideally react rapidly and quantitatively with the fuel molecules. This requirement can be achieved by operating at high fuel concentration, which ensures that Cl$^+$ is dominantly consumed by reaction with the fuel and effectively suppresses secondary Cl$^+$ reactions with radical intermediates (R$_p$, R$_s$, R$_p$O$_2$, R$_s$O$_2$) and products. However in the present case, because of the efficient ionization of the ketone at the highest photon energies employed here, high DEK concentrations would unacceptably increase the total ion count rates and [DEK] was kept ≤7.2 × 10$^{13}$ cm$^{-3}$ (see Table 1). Therefore the contributions of side and secondary chlorine reactions remain non-negligible and must be quantified.

The reactions of DEK and its radicals with atomic$^{24-27}$ and molecular$^{25}$ chlorine have been studied previously under a variety of temperature and pressure conditions. Fig. 1 shows current results for the time profile of DEK depletion when the concentration of Cl$^+$ immediately following photolysis is 13% of [DEK]$_0$ (Table 1). A rapid drop in DEK concentration occurs in the first ~3 ms, followed by a more gradual depletion for the extent of reaction. Hydrogen abstraction from diethyl ketone can give rise to the two distinct initial radicals defined in Scheme 2 along with calculated C–H bond dissociation enthalpies (BDE; 0 K). Unless noted otherwise, calculations presented in this work are at the CBS-QB3$^{28,29}$ level using Gaussian 09.$^{30}$ Because of the greater degree of alkyl substitution and the vinoxy-type resonance stabilization, the secondary radical (R$_s$) that results from abstraction of a β hydrogen is 13 kcal mol$^{-1}$ more stable than the primary radical (R$_p$) originating from abstraction of a γ hydrogen. Using the bond dissociation enthalpy (BDE) of H–Cl at 0 K (102.3 ± 0.1 kcal mol$^{-1}$)$^{31}$ abstraction of a

![Scheme 2](image-url)
primary hydrogen to give \( R_p + \text{HCl} \) is only slightly exothermic (\( \Delta H_{\text{rxn}} \approx -2 \text{ kcal mol}^{-1} \)), but formation of \( R_s + \text{HCl} \) is much more exothermic (\( \Delta H_{\text{rxn}} \approx -15 \text{ kcal mol}^{-1} \)). Kaiser et al.\(^{25}\) investigated the reactions of atomic chlorine with DEK at temperatures from 297 K to 515 K, reporting rate constants of \( 4.0 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \) for primary H atom abstraction and \( 6.3 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \) for secondary H atom abstraction. These expressions are extrapolated to predict branching ratios in the present experiments: \( R_p/R_s = 0.40 \) at 550 K and \( R_p/R_s = 0.43 \) at 650 K.

The resonance stabilization of \( R_s \) is lost upon addition of \( \text{O}_2 \) and as a result the \( R_s\text{O}_2 \) well is shallow (24 kcal mol\(^{-1}\); Fig. 2) relative to that of \( R_p\text{O}_2 \) (36 kcal mol\(^{-1}\); Fig. 3). Because of the stability of \( R_s \), formation of bimolecular products from \( R_s\text{O}_2 \) must proceed \textit{via} saddle points that are above the energy of the \( R_s + \text{O}_2 \) entrance channel, which limits their importance such that back dissociation to reactants is favored. Kaiser et al.\(^{25}\) observed the \( R_s + \text{O}_2 \leftrightarrow R_s\text{O}_2 \) equilibrium shifting towards reactants at even modestly elevated temperatures, but detected no back dissociation of \( R_p\text{O}_2 \) to \( R_p + \text{O}_2 \) up to 510 K under their experimental conditions. The temperatures of the current study are still higher (550–650 K). From the CBS-QB3 calculated equilibrium constant and the \( \text{O}_2 \) concentration in our experiments (Table 1), the equilibrium ratio \([R_s\text{O}_2]/[R_s]\) at 550 K is 0.067, while at 650 K it is only 0.002, showing that the equilibrium heavily favors the \( R_s + \text{O}_2 \) reactants at these conditions.

During the timescale of the initial reaction of \( \text{Cl}^* \) with DEK, the following reactions (1–6) involving atomic and molecular chlorine can also occur (\( d_e\)-DEK is employed to highlight different isomers):

\[
\begin{align*}
(1) \quad \text{Cl}_2 + \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{CH}_2 \rightarrow \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{Cl} + d_e\text{R}_p \\
(2) \quad \text{Cl}_2 + \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{CH}_2 \rightarrow \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{Cl} + d_e\text{R}_p\text{Cl} \\
(3) \quad \text{Cl}_2 + \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{CH}_2 \rightarrow \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{Cl} + d_e\text{R}_p\text{Cl}
\end{align*}
\]

\[
\begin{align*}
(4) \quad \text{Cl}_2 + \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{CH}_2 \rightarrow \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{Cl} + d_e\text{R}_p\text{Cl} \\
(5) \quad \text{Cl}_2 + \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{CH}_2 \rightarrow \text{H}_3\text{C}_2\text{CD}_2\text{CD}_2\text{Cl} + d_e\text{R}_p\text{Cl}
\end{align*}
\]
Of particular interest are reactions 3 and 6, H-abstraction or addition–elimination reactions of Cl*/C15 with Rp and Rs to form vinyl ethyl ketone (VEK). Because VEK is also the product expected from the chain-terminating oxidation channel outlined in Scheme 1, accurate branching fraction determination in R + O2 requires characterization of the Cl*/C15 + R side reactions.

The top panel of Fig. 4 shows the product mass spectrum resulting from the reaction of d4-DEK (m/z = 90) + Cl2 + Cl at 550 K in the absence of O2, measured at an ionizing photon energy of 11.0 eV. The bottom panel shows the results in the presence of O2. Both curves are normalized to depletion of the precursor d4-DEK (Fig. 1) both in the presence and absence of O2. The identical rise times of d4-RpCl and d3-RsCl support the interpretation that these chlorinated products largely originate from R + Cl*/C15 and not from chain chlorination in light of the vastly different chain chlorination rates of the two radicals. Kaiser et al. report that chain chlorination reaction 1 involving Rp is nearly 150 times faster than chain chlorination reaction 4 involving Rs (k1 = 2.7 × 10−12 cm3 molecule−1 s−1 and k4 = 1.85 × 10−14 cm3 molecule−1 s−1).

An estimate of the proportion of initial radicals that react with Cl*/C15 can be derived from comparing the amplitudes of chlorinated and oxygenated product signals as a function of O2 concentration. Fig. 5 shows the effect of O2 on the chlorinated product signals at 550 K taken at 11.0 eV resulting from the reaction of d4-DEK + Cl*/C15. These spectra result from integrating the ion signal over only the first 3 ms immediately following photolysis, and are dominated by the products of Cl*/C15 reactions. The red dotted curve of Fig. 5 shows chlorinated product formation in the presence of [O2] = 2.8 × 1016 cm−3 and the black solid curve shows products in the absence of O2. Both curves are normalized to depletion of the precursor d4-DEK (note differing y-axis scales) and displayed so that the peak heights of the m/z = 123 feature (RsCl) coincide. The concentration of RsCl is found to diminish by a factor of 2.6 due to the presence of O2, whereas RpCl diminishes by a factor of 23.3. The greater reduction of RpCl relative to RsCl is consistent with rapid back dissociation of RsO2 and lack of low-lying RsO2 exit channels (Fig. 2). The introduction of [O2] = 2.8 × 1016 cm−3 intercepts ~ 60% of Rp but > 95% of Rp.

Fig. 6 shows the ratio of cyclic ether (m/z = 100) signal to that of VEK (m/z = 84) as a function of [O2] for Cl-initiated oxidation.
of non-deuterated DEK. As expected, with no O₂ added to the reacting flow, no cyclic ether is produced and the ratio \( m/z = 100 : m/z = 84 \) is 0. As O₂ is added, the relative concentration of cyclic ether increases until reaching a plateau near a ratio of \( [O₂] : [Cl^*]_0 = 3300 \). At this point, the ratio \( m/z = 100 : m/z = 84 \) changes very little as a function of increasing \([O₂] : [Cl^*]_0\). To minimize the influence of secondary chlorine chemistry while maintaining sufficient signal for oxidation products, we used \([O₂] : [Cl^*]_0\) ratios of 3300 and 6600, indicated by the red and blue boxed data points highlighted in Fig. 6. However, because of the difference in reactivity of the two initial fuel radicals, the plateau in Fig. 6 does not imply that VEK yield from chlorine reactions has been eliminated. Even in the presence of \([O₂] : [Cl^*]_0 > 3300\), substantial signal is observed for R₅Cl (\(m/z = 123\) and 125; Fig. 4 and 6), indicating that reaction 5 (R₅ + Cl* → R₅Cl) and therefore also reaction 6 (R₅ + Cl* → VEK + HCl) still contribute, although reaction 2 (R₅ + Cl* → R₅Cl) and therefore reaction 3 (R₅ + Cl* → VEK + HCl) have been effectively quenched, giving the plateau. If it were feasible to increase the fuel concentration in these experiments, higher \([DEK]\) would reduce the role of Cl + R reactions. However, because of its low reactivity with O₂, even at low Cl⁺ concentrations the R₅ radical is still more likely than R₅ to react via other competing pathways, such as self reaction, rather than with O₂.
This assignment is corroborated by the observation of the Rₚ dimer, arising from recombination of Rₚ (see below).

To determine the fraction of VEK produced by chlorine reactions vs. HO₂ reactions, one can compare the ratio of RₚCl and RₛCl signals to VEK in the absence and presence of O₂ using d₄-DEK, where RₛCl and RₚCl molecules have different masses due to deuterium substitutions. Using the known chain chlorination rates for reactions 1 and 4 and estimates for the rates of reactions 2, 3, 5 and 6, these ratios yield the VEK branching estimate. The model is detailed in the ESI† and we estimate that at the high O₂ concentrations used in this work ~1/4 of the total VEK signal is due to reaction 6, with the remaining VEK signal arising from Rₛ + O₂ and Rₚ + O₂ reactions (see below). In the presence of O₂, reaction 3 gives a negligible contribution to VEK. Finally, reactions of Cl* + RO₂ to form ClO + oxy radical could also affect the interpretation of these experiments. However, peaks at 51 (³⁵ClO) and 53 (³⁷ClO) are barely discernible in our spectra, suggesting that such reactions are unimportant in these experiments.

B. Reactions of RO₂

Fig. 7 shows the product mass spectra for Cl-initiated oxidation of non-deuterated diethyl ketone (DEK) at 550 K (bottom) and 650 K (top). Major peaks are observed at m/z = 141, m/z = 122, m/z = 120, m/z = 100, m/z = 84, m/z = 72, m/z = 57, m/z = 44, m/z = 42, m/z = 30 and m/z = 28. The figure inset shows an integrated slice at higher energy (10.8–11.0 eV) demonstrating formation of ethene (C₂H₄; m/z = 28) and formaldehyde (H₂CO; m/z = 30). Peaks at m/z = 122, m/z = 120 and m/z = 57 are assigned to RCl. The signal at m/z = 141 is due to a fragment ion of the Rₛ dimer. ESI† Fig. S2 is an extended mass spectrum which shows that in the absence of O₂ RₛRₛ and RₛRₚ dimers are observed and give rise to fragment ions corresponding to loss of neutral ethyl radical. In the presence of O₂ only peaks associated with the RₛRₛ dimer and its fragment remain, consistent with rapid consumption of Rₛ by O₂ and a significant concentration of Rₛ even in the presence of a large excess of O₂. Fig. 8 shows the time profiles for several products observed in the 550 K Cl-initiated oxidation of d₄-DEK, demonstrating that important oxidation products are generated on a significantly slower time scale than the rapid accumulation of chlorinated products.

i. Cyclic ether formation. The peak at m/z = 100 (Fig. 7) is assigned to cyclic ether products and serves as a marker for decomposition of QOOH radicals (Scheme 1). A number of possible pathways and cyclic ether products are available in DEK oxidation. Reactions 7 and 8 show these channels for Rₛ and Rₚ and Table 2 gives the potential products along with relative energies (0 K) and adiabatic ionization energies (AIEs) calculated at the CBS-QB3 level. The third column indicates the possible initial radical(s) that could yield the corresponding cyclic ether product. For example, the three-membered-ring cyclic ether shown in the first row, 1-(oxiran-2-yl)propan-1-one, could be formed via O₂-addition to Rₚ, followed by internal hydrogen transfer from the neighboring β carbon to yield a secondary hydroperoxalkyl (QOOH) radical, which can then decompose to OH + cyclic ether (channel 1, reaction 7). Similarly, O₂-addition to Rₛ, followed by internal hydrogen transfer from the neighboring γ carbon would yield a primary QOOH and subsequently the same cyclic ether (channel 4, reaction 8).

Fig. 9a shows the photoionization spectrum for the cyclic ether products at m/z = 100 in DEK oxidation. The ionization onset near 9.1 eV matches well the calculated values for the 4- and 5-membered ring cyclic ethers shown in Table 2,
Table 2: Relative energies at 0 K (ΔE₀,rel) of neutral cyclic ethers associated with diethyl ketone oxidation and the corresponding adiabatic ionization energies (AIEs). All energies were calculated at the CBS-QB3 level of theory.

<table>
<thead>
<tr>
<th>Exp. IE = 9.1 eV</th>
<th>ΔE₀,rel (kcal mol⁻¹)</th>
<th>AIE (eV)</th>
<th>H-abstraction (s-secondary; p-primary)</th>
<th>m/z (d₀, d₁, d₆-DEK)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H₂C==C==CH₂</td>
<td>20.8</td>
<td>9.6</td>
<td>Rₚ, Rₛ</td>
<td>100, 103, 105</td>
</tr>
<tr>
<td>1-(Oxiran-2-yl)propan-1-one</td>
<td>14.1 (14.3)</td>
<td>9.1</td>
<td>Rₛ</td>
<td>100, 102, 106</td>
</tr>
<tr>
<td>2,4-Dimethyl-oxetan-3-one</td>
<td>0.0</td>
<td>9.1</td>
<td>Rₚ, Rₛ</td>
<td>100, 103, 105</td>
</tr>
<tr>
<td>2-Me-THF-3-one</td>
<td>1.2</td>
<td>9.6</td>
<td>R₀</td>
<td>100, 104, 104</td>
</tr>
<tr>
<td>THP-4H-4-one</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*trans* isomer.

2,4-dimethyloxetan-3-one and 2-methyltetrahydrofuran-3-one (2-MeTHF-3-one), respectively. Also shown in Fig. 9a are the standard photoionization spectra recorded for 2-MeTHF-3-one and tetrahydro-4H-pyran-4-one (THP-4H-4-one), the 5- and 6-membered ring cyclic ether products shown in Table 2 and reactions 7 and 8. The 3- and 4-membered ring cyclic ethers were not commercially available nor did we attempt to synthesize them. The photoionization spectrum of 2-MeTHF-3-one matches almost exactly that of the m/z = 100 product in DEK oxidation. No evidence of an onset in the range 9.4–9.6 eV is observed, as would be expected from a contribution from THP-4H-4-one or the oxirane (Table 2). Based on the results of undeuterated DEK alone, and without knowledge of the shape of the photoionization spectrum of 2,4-dimethyloxetan-3-one (4-membered ring, Table 2), formation of this compound via channel 5 (reaction 8) cannot be definitively eliminated from consideration.

However, studying d₄- and d₆-DEK allows separating the cyclic ether isomers by mass. The final column of Table 2 indicates the expected values of m/z for the various cyclic ether isotopologs under consideration. Based on the above discussion, we would expect to see a signal at m/z = 103 (d₁-2-MeTHF-3-one) from d₁-DEK oxidation and analogously at m/z = 105 (d₆-2-MeTHF-3-one) in d₆-DEK oxidation. In contrast, formation of the oxetane (4-membered ring) would yield products at m/z = 102 in d₄-DEK oxidation and at m/z = 106 in d₆-DEK oxidation. Fig. 9b shows the difference mass spectra for the series of diethyl ketones studied here. Both deuterated samples show a major peak at the m/z value expected from 2-MeTHF-3-one and only minor contributions at masses associated with 2,4-dimethyloxetan-3-one and THP-4H-4-one. Furthermore, as Fig. 9a shows, the photoionization spectra of m/z = 103 in d₁-DEK and m/z = 105 in d₆-DEK also agree well with the 2-MeTHF-3-one standard. The slight difference in shape may be due to the presence of deuterium atoms in the d₄-DEK and d₆-DEK oxidation products. Not shown in Fig. 9a are the dissociative photoionization products of 2-MeTHF-3-one and THP-4H-4-one. Each displays a strong fragment ion at m/z = 72. This m/z = 72 peak of 2-MeTHF-3-one has a shallow onset near 9.4 eV, and agrees well with the m/z = 72 feature observed in DEK oxidation. This feature is assigned to loss of CO from the 2-MeTHF-3-one cation. Similar features are observed at the expected masses (m/z = 75 and m/z = 77) in the oxidation spectra of d₁- and d₆-DEK. Taken as a whole, it is clear from the experimental data that one or both of the 5-membered ring cyclic ether channels of reactions 7 (channel 3) and 8 (channel 6) is dominant and thus expected to be a prominent chain-propagation mechanism in the ignition chemistry of diethyl ketone. We return to channels 3 and 6 in a discussion of computed potential energy surfaces below.

ii. HO₂-elimination. Continuing discussion in reference to Fig. 7, the m/z = 84 signal is assigned to a combination of the
HO₂-elimination channels and reaction 6 (Cl⁺ + Rₚ). The m/z = 84 product could originate from peroxy radicals associated with either Rₚ or Rₛ (reactions 9 and 10). These channels have been investigated in a recent article discussing 1,2-acyl group migration reactions in open-chain ketones and will not be discussed at length here. Briefly, the photoionization spectrum of the m/z = 84 product agrees very well with that of a vinyl ethyl ketone (VEK) standard and is given in Fig. 10, confirming the VEK assignment. Good agreement with the VEK standard is also observed for the product photoionization spectra for m/z = 87 (d₃-VEK) and m/z = 89 (d₅-VEK) in d₄- and d₆-DEK oxidation. The figure inset shows a small signal with an early onset (~8.4 eV) for m/z = 84 in DEK oxidation. This signal may be due to 3-hydroxy-penta-1,3-diene (AIE = 8.24 eV), however, its origin is unclear.

iii. Potential energy surfaces. Fig. 2 and 3 show potential energy surfaces calculated for the HO₂-elimination and cyclic ether formation channels associated with the Rₛ + O₂ and Rₚ + O₂ systems (reactions 7–10). Pathways for formation of the 3-, 5- and 6-membered ring cyclic ethers (reaction 7) as well as HO₂-elimination to form VEK (reaction 9) are all energetically plausible from RₚO₂. Note that no 4-membered ring cyclic ether is expected from unimolecular reaction of RₚO₂. Formation of 2-MeTHF-3-one from RₛO₂, is the lowest energy channel and involves a secondary, resonance-stabilized QOOH intermediate. This resonance stabilization may be expected to lower the barrier associated with the RO₂ → QOOH isomerization. However, the calculated CBS-QB3 barrier (20.4 kcal mol⁻¹) is similar to that for the analogous channel in n-pentylperoxy calculated at the same level of theory, potentially due to increased ring strain in the RₛO₂ reaction arising from the presence of the sp²-hybridized carbonyl carbon atom in the cyclic saddle-point structure when compared to the n-pentylperoxy system. The QOOH associated with formation of the oxirane in Fig. 3 is also resonance-stabilized, but the degree of strain in the 5-membered ring RO₂ → QOOH transition state leads to a significantly larger barrier (34.1 kcal mol⁻¹), making this channel unfavored. In contrast to Rₚ + O₂, every anticipated unimolecular RₛO₂ exit channel has at least one saddle-point energy above the entrance channel (Fig. 2). Considering also that the Rₛ + O₂ ↔ RₛO₂ equilibrium favors reactants, it is unclear how important the pathways of Scheme 1 will be in the oxidation of Rₛ. All stationary point energies and geometries for the potential energy surfaces of Fig. 2 and 3 are given in ESI† along with values of the T1 diagnostic of Lee et al. 34 The T1 diagnostic for all structures is within the range for which single reference methods are appropriate (<0.03) with the exception of the saddle points associated with QOOH → cyclic ether + OH, which have T1 diagnostic values of 0.033–0.045. For the latter points the electronic structures are expected to have higher multireference character leading to somewhat larger uncertainties in the total energies.

C. Small molecule products

Fig. 11a-d shows photoionization spectra confirming that peaks at m/z = 28, m/z = 30, m/z = 42 and m/z = 44 in the Cl-initiated oxidation of DEK at 550 K are due to closed-shell products ethene (H₂C=CH₂), formaldehyde (CH₂O), ketene (CH₂CO) and acetaldehyde (CH₃CHO). These products are also seen in d₄-DEK oxidation (Fig. 4) in which ethene is observed at m/z = 30.05 (H₂CD₂). Formaldehyde is observed in roughly the same abundance at m/z = 30.01 (H₂CO) and m/z = 32.01 (D₂CO).
Ketene is observed in roughly the same abundance at $m/z = 42$ (H$_2$CCO) and $m/z = 44$ (D$_2$CCO) and acetaldehyde is observed at exclusively at $m/z = 45$ (CH$_3$CDO). Similarly, for $d_6$-DEK, the products are observed as follows: $d_4$-ethene ($m/z = 30.05$), $d_0$- and $d_2$-formaldehyde ($m/z = 30.02$ and $m/z = 32$), $d_0$- and $d_2$-ketene ($m/z = 42$ and $m/z = 44$) and $d_3$-acetaldehyde ($m/z = 47$; CD$_3$CHO). These results are summarized in Table 3 and all product photoionization spectra are included in Fig. 11. ESI,† Fig. S3 shows mass spectra of $d_4$-DEK oxidation in the presence and absence of O$_2$ in the region of ethene and formaldehyde and demonstrates the mass resolution capability of the current MPIMS experiment. No H/D atom scrambling is observed; for instance, no $d_1$-formaldehyde or $d_1$-ketene is seen in either the $d_4$- or $d_6$-DEK reactions.

The top panel of Fig. 4 shows that when O$_2$ is removed, ethene is still formed. This is consistent with formation via the $R_p$ $\beta$-scission reaction 11, yielding ethene and propionyl radical. Propionyl is expected to decarbonylate to form ethyl radical, which will most likely add O$_2$ and undergo HO$_2$-elimination, forming an additional ethene molecule (reaction 12). The first step of reaction 11 proceeds with a calculated barrier of 21 kcal mol$^{-1}$. Similarly, $R_s$ can undergo $\beta$-scission reaction 13 to form methyl ketene and ethyl radical. Methyl ketene is obscured by dissociative photoionization of DEK so that we cannot quantify its formation. However, calculations indicate this reaction proceeds with a large barrier (43 kcal mol$^{-1}$) and is expected to be far slower than reaction 11 at 550 K.

\begin{equation}
H_3C\overset{\cdot}{C}\overset{\cdot}{D}_2 + O_2 \rightarrow HO_2 + H_3C\overset{\cdot}{C}\overset{\cdot}{D}_2
\end{equation}
Formaldehyde, ketene and acetaldehyde are formed only in the presence of O\textsubscript{2}, and likely arise from oxy radicals formed in secondary reactions of RO\textsubscript{2}. Self-reaction of RO\textsubscript{2} to yield two oxy radicals + O\textsubscript{2} is known to be important in many oxidative systems.\textsuperscript{11} In the case of DEK, the resonance-stabilized R\textsubscript{s} reacts relatively slowly with O\textsubscript{2} and undergoes significant side reactions, e.g., dimerization both in the presence and absence of O\textsubscript{2}. The R\textsubscript{s} may also react with R\textsubscript{p}O\textsubscript{2} to yield oxy radicals that would be expected to undergo further decomposition to generate acetaldehyde, formaldehyde and ketene, as shown in reactions 14–16 for d\textsubscript{4}-DEK.

Self-reaction of d\textsubscript{4}R\textsubscript{p}O\textsubscript{2} would yield two oxy radicals identical to the reactant of reaction 16. These reactions account for CH\textsubscript{3}CDO (m/z = 45), H\textsubscript{2}CO (m/z = 30) and D\textsubscript{2}CCO (m/z = 44) in d\textsubscript{4}-DEK oxidation. However, D\textsubscript{2}CO (m/z = 32) and H\textsubscript{2}CCO (m/z = 42) (Table 3) are still unexplained.

Table 3 Assignments for the small molecule products in the Cl-initiated oxidation of diethyl ketone (DEK) d\textsubscript{4}-diethyl ketone (d\textsubscript{4}-DEK) and d\textsubscript{6}-diethyl ketone (d\textsubscript{6}-DEK) at 550 K

<table>
<thead>
<tr>
<th>m/z</th>
<th>DEK</th>
<th>d\textsubscript{4}-DEK</th>
<th>d\textsubscript{6}-DEK</th>
</tr>
</thead>
<tbody>
<tr>
<td>28</td>
<td>H\textsubscript{2}CCH\textsubscript{2}</td>
<td></td>
<td></td>
</tr>
<tr>
<td>29</td>
<td>CH\textsubscript{3}CH\textsubscript{2} + CH\textsubscript{3}CH\textsubscript{2}O\textsubscript{2}</td>
<td>H\textsubscript{2}CO</td>
<td>H\textsubscript{2}CO</td>
</tr>
<tr>
<td>30.02</td>
<td>H\textsubscript{2}CO</td>
<td>D\textsubscript{2}CH\textsubscript{2}</td>
<td></td>
</tr>
<tr>
<td>30.05</td>
<td>H\textsubscript{2}CO</td>
<td>CH\textsubscript{3}CD\textsubscript{2} + CH\textsubscript{3}CD\textsubscript{2}O\textsubscript{2}</td>
<td>D\textsubscript{2}CO</td>
</tr>
<tr>
<td>31</td>
<td>D\textsubscript{2}CO</td>
<td></td>
<td></td>
</tr>
<tr>
<td>32.02</td>
<td>D\textsubscript{2}CO</td>
<td>CD\textsubscript{3}CH\textsubscript{2} + CD\textsubscript{3}CH\textsubscript{2}O\textsubscript{2}</td>
<td>H\textsubscript{2}CCO</td>
</tr>
<tr>
<td>32.06</td>
<td>D\textsubscript{2}CO</td>
<td>D\textsubscript{2}CCO</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 11 Comparison of the photoionization spectra of the indicated peaks observed in Cl-initiated oxidation of undeuterated DEK, d\textsubscript{4}-DEK and d\textsubscript{6}-DEK at 550 K with calibration spectra (a) ethene (b) formaldehyde (c) ketene (d) acetaldehyde. The early onset observed in the acetaldehyde plot may be due to a small contribution from vinyl alcohol.
In recent work from our group, the gas phase rearrangement of primary ketone radicals via 1,2-acyl group migration was shown to be fast relative to O₂-addition at the conditions of this study. This rearrangement is shown in reaction 17 for d₄-R₄O₂. This rapid equilibration is expected to yield both d₀- and d₂-formaldehyde as well as d₀- and d₂-ketene via reactions 15 and 16.

Other possible routes for the formation of acetaldehyde appear less likely, for example, the β-scission reaction of QOOH radicals derived from R₄O₂, shown for d₄-DEK in reactions 18 and 19:

The second step of reactions 18 and 19 proceeds with calculated barriers of 33 and 22 kcal mol⁻¹, respectively. Isomerization of QOOH back to RO₂ and ring closure to form cyclic ether + OH will both be more favorable.

Recently a reaction has been characterized that involves H₂O-elimination from QOOH radicals to yield alkoxy radical co-products, which in turn typically rapidly undergo unimolecular decompositions via β-scission. We have considered (see also ESI†) whether any similar water elimination could contribute to aldehyde and ketene formation in the DEK oxidation. However, the barrier heights are larger than 40 kcal mol⁻¹ (in contrast to the ~12–17 kcal mol⁻¹ barriers found for alcohols), and we conclude that water elimination does not play a significant role in the oxidation of ketones.

D. Product branching fractions

Relative product branching fractions can be determined by integrating product signals over a range of kinetic times, accounting for mass discrimination in the TOF mass spectrometer and using previously known or currently measured photoionization cross sections of the pure compounds. The results are given for oxidation and β-scission products in Table 4. Ethene becomes the major product at 650 K, reflecting increasing dominance of β-scission reactions from the initial radicals. At 550 K, both chain-propagating OH-elimination and chain-terminating HO₂-loss are observed to be major oxidation channels, each responsible for roughly 15–20% of the quantified products as correlated by the concentrations of 2-MeTHF-3-one and VEK co-products. As shown in Fig. 3, although the barrier height for RpO₂ → QOOH is similar to analogous saddle points in saturated hydrocarbon oxidations (20 kcal mol⁻¹), the reverse reaction of QOOH → RpO₂ has a higher barrier (16 kcal mol⁻¹), reflecting the resonance stabilization of the QOOH.
stability of the QOOH intermediate shifts the RO₂ + O₂ equilibrium to the right, and may favor reactions that arise from the QOOH isomer, such as unimolecular reaction to form cyclic ethers or bimolecular reaction with a \( \text{O}_2 \) molecule. Therefore, whereas resonance stabilization of the fuel radical \( \text{Rs} \) tends to inhibit reactivity with \( \text{O}_2 \) in the oxidation of the non-resonance-stabilized radical isomer \( \text{Rp} \), the associated resonance stabilization of the QOOH radical may tend to increase the chain propagation and chain branching pathways. This behavior will be general in the initial oxidation steps of fuels that can form both resonance-stabilized and non-resonance-stabilized radical isomers following hydrogen abstraction.

### V. Conclusions

The initial reactions in the Cl-initiated oxidation of diethyl ketone and a series of deuterated diethyl ketones have been studied with multiplexed photoionization mass spectrometry using tunable synchrotron ionizing radiation. The secondary DEK radical generated by H-abstraction reactions, \( \text{Rs} \), is resonance-stabilized and thus has a shallow well for \( \text{O}_2 \) addition, which makes the radical much less reactive with \( \text{O}_2 \) at the temperatures considered here (550–650 K).

The oxidation of DEK at 550 K yields chain-terminating HO₂ + VEK and chain-propagating OH + cyclic ether products in roughly equal abundance. A number of possible cyclic ether species are available in principle. However, nearly exclusive production of 2-methyltetrahydrofuran-3-one (2-Me-THF-3-one) is observed. This product could result from \( \text{O}_2 \)-addition to either \( \text{Rs} \) or \( \text{Rp} \). Reactions with selectively deuterated DEK allow us to rule out any significant formation of the other possible cyclic ethers. Calculated potential energy surfaces reveal that barriers to formation of oxidation products of \( \text{R}_n \text{O}_2 \) lie above the \( \text{R}_n + \text{O}_2 \) entrance channel. Conversely, the lowest energy chain-propagating pathway for decomposition of \( \text{R}_n \text{O}_2 \) is energetically favorable and proceeds via a resonance-stabilized QOOH to the observed 2-Me-THF-3-one product.

The small-molecule products observed provide evidence for the generation of \( \text{O} \) radicals from the reaction of \( \text{Rs} + \text{RsO}_2 \) and self-reaction of \( \text{R}_n \text{O}_2 \). Observation of \( \text{d}_0 \)- and \( \text{d}_2 \)-formaldehyde, \( \text{d}_0 \)- and \( \text{d}_2 \)-ketene in both \( \text{d}_2 \text{-DEK} \) and \( \text{d}_0 \text{-DEK} \) provides additional corroboration of rapid 1,2-acyl group migration reactions in ketone radicals.\(^\text{33}\)

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