Reaction Pathway Control via Reactant Vibrational Excitation and Impact on Product Vibrational Distributions: The O + HO₂ → OH + O₂ Atmospheric Reaction

Qixin Chen, Shuwen Zhang, Xixi Hu,¹ Daiqian Xie, and Hua Guo

ABSTRACT: Chemical reactions often have multiple pathways, the control of which is of fundamental and practical importance. In this Letter, we examine the dynamics of the O + HO₂ → OH + O₂ reaction, which plays an important role in atmospheric chemistry, using quasi-classical trajectories on a recently developed full-dimensional potential energy surface (PES). This reaction has two pathways leading to the same products: the H abstraction pathway (O_1 + HO_2 → OH + O_3) and the O abstraction pathway (O_2 + HO_2 → O_1H + O_1O). Under thermal conditions, the reaction is dominated by the latter channel, which is barrierless, leading to vibrational excitation of the O₂ product. However, we demonstrate that excitation of the HO₂ reactant in its O–H (νᵢ) vibrational mode results in dramatic switching of the reaction pathway to the activated H abstraction channel, which leads to a highly excited OH product vibrational state distribution. The implications of such dynamical effects in the atmospheric chemistry are discussed.

Atomic oxygen (Oa), leaving the ObH group as a spectator. The

H(νᵢ) acts as a spectator. Although both pathways lead to nominally the same products, it is readily seen from Figure 1 that they are not expected to have equal contributions under thermal conditions, due to their different barrier heights. At conditions relevant to atmospheric chemistry, the OA channel dominates. Furthermore, it is clear from the transition state configurations that the products are likely to have different extents of vibrational excitation. Indeed, the O₃–O₂ bond length at SP1 is much larger than the one for oxygen molecule, which may produce highly vibrationally excited O₂, same for the OH radical in the HA pathway. As described below, the control of the product branching of a chemical reaction. The relative yields of various product channels are often influenced by the differing barrier heights in the corresponding pathways and dynamics could also play an important role.¹ The rapid progress of experimental and theoretical understanding of the dynamics of multichannel reactions in recent years has greatly advanced our understanding of the dynamical control of reactivity and product branching.²⁻⁵ In a recent study, for example, the branching ratio between the CH₃O + HF and CH₂OH + HF channels in the F + CH₃OH reaction was investigated theoretically, and it was shown that stereodynamics has a more significant impact than energetics.⁶

In this Letter, we examine a slightly different scenario in which two reaction pathways lead to the same products, as manifested by the following reaction:

HO₂(X'A') + O(³P) → OH(X'Π) + O₂(X'Σ⁺)  (R1)

This radical–radical reaction is of great importance in atmospheric chemistry as it provides a key atomic oxygen destruction pathway in the stratosphere (215–270 K) and mesosphere (180–270 K) and plays a major role in controlling the partitioning among the HO₂ species.¹ The vibrational state distribution of the OH radical produced by reaction R1 could potentially influence an array of atmospheric phenomena. For instance, several authors have proposed that the nascent OH product may have significant vibrational excitation, thus responsible for hydroxyl night airglow.⁸⁻¹¹ However, this proposal has been disputed by others who believe that the OH vibrational excitation is small.¹²⁻¹⁵ Definitive evidence to resolve issues like this requires the determination of the OH vibrational distribution for the title reaction.

As shown in Figure 1, this exothermic reaction has two pathways.¹⁶,¹⁷ The O abstraction (OA) pathway (O₃ + HO₂ → O₁H + O₃O₂), featuring a submerged barrier (SP1), proceeds by abstracting the terminal oxygen of HO₂ (O₃) by the atomic oxygen (O₃), leaving the O₁H group as a spectator. The alternative H abstraction (HA) pathway (O₂ + HO₂ → O₂ + O₃O₂), which has a small barrier (SP2), proceeds by abstracting the H atom from HO₂ by the atomic oxygen (O₃), with the O₂ moiety as a spectator. Although both pathways lead to nominally the same products, it is readily seen from Figure 1 that they are not expected to have equal contributions under thermal conditions, due to their different barrier heights. At conditions relevant to atmospheric chemistry, the HA channel dominates. Furthermore, it is clear from the transition state configurations that the products are likely to have different extents of vibrational excitation. Indeed, the O₃–O₂ bond length at SP1 is much larger than the one for oxygen molecule, which may produce highly vibrationally excited O₂, same for the OH radical in the HA pathway. As described

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below, these observations are indeed born out from our dynamics calculations on a recently developed global potential energy surface (PES). We, further, demonstrate that it is possible to control the relative contributions of the two pathways by exciting a vibrational mode of the HO2 reactant, which also leads to drastically different product state distributions.

In 1998, Varandas and co-workers\textsuperscript{18} published a pioneering study on the dynamics of O + HO\textsubscript{2} reaction, reporting the ro-vibrational distributions OH and O\textsubscript{2} at the collision energy of $E_c = 4.184$ kJ/mol. Using a quasi-classical trajectory (QCT) method, the dynamical calculations were performed on the first full-dimensional PES for this system, which was fitted using the double many-body expansion (DMBE) method to \textit{ab initio} points calculated using unrestricted configuration interaction with single and double excitations (UCISD) and the 6-311G+ + (d, p) basis set.\textsuperscript{18} On this DMBE I PES, the majority of the OH product were found in the vibrational ground state, via the OA pathway. In 2004, these authors further studied the role of HO\textsubscript{2} vibrational excitation in the title reaction on the DMBE I PES\textsuperscript{17} and an improved DMBE II PES\textsuperscript{19} and concluded that the impact of HO\textsubscript{2} vibrational excitation is relatively minor even for highly vibrationally excited HO\textsubscript{2}.\textsuperscript{20} The DMBE PESs have also been used in transition-state theory calculations of the rate coefficients, but the results are not in good agreement with known experimental values.\textsuperscript{21}

Very recently, we have developed a more accurate full-dimensional global PES for the HO\textsubscript{3} system constructed from more than 21,000 multireference configuration interaction (MRCI) points\textsuperscript{22,23} using the high-fidelity permutation invariant polynomial-neural network (PIP-NN) method.\textsuperscript{24} The active space used in the CASSCF calculations includes 13 electrons in 10 orbitals with 1s and 2s orbitals of O atoms closed, while the full-valence active space was used in subsequent MRCI calculations. The MRCI treatment of the PES was found to be essential due to the strongly correlated nature of this system.\textsuperscript{25,26} This new PES has been demonstrated to reproduce various spectroscopic, kinetic, and dynamics properties of the system.\textsuperscript{22,23,27,28} In several comparisons, the results on the new PES were found to differ significantly from those obtained from the DMBE PESs,\textsuperscript{17,19} presumably due to the neglect of multireference effects in the earlier \textit{ab initio} calculations and the relatively small number of \textit{ab initio} points used in the DMBE fitting. As a result, it is not clear if the previous dynamics results for the title reaction obtained from the DMBE PESs are reliable. In this Letter, we report a detailed QCT dynamical study of R\textsubscript{1} on our new PES, focusing on the impact of vibrational excitation of the HO\textsubscript{2} reactant on the reaction mechanism as well as the product internal distributions, which might impact atmospheric modeling.

Standard QCT calculations were performed using VENUS\textsuperscript{29} on the PIP-NN PES with the reactants prepared in various vibrational states. For each vibrational state of HO\textsubscript{2}, the normal mode coordinates and momenta were transformed to the atomic counterparts.\textsuperscript{30} The initial relative translational energy and initial rotational energies of reactants were sampled from the Boltzmann distribution at 200 K, which is relevant for the atmospheric conditions. 120,000 trajectories were propagated for each vibrational state with a time step of 0.1 fs and the maximum impact parameter $b_{\text{max}}$ of 5.5 Å. The vibrational state distributions of the diatomic products were obtained using the Einstein–Brillouin–Keller method,\textsuperscript{31} by binning the vibrational actions to the nearest integers. Mechanistic insights on the particular pathway taken by a trajectory can be readily identified by examining the connectivity of the products (HO\textsubscript{3} + O\textsubscript{2}O\textsubscript{2} and HO\textsubscript{2} + O\textsubscript{2}O\textsubscript{2} for the HA and OA channels, respectively.) In reality, such channels can be distinguished by isotope labeling, as done by Sridharan et al.\textsuperscript{32}

The energetics of the title reaction on the PIP-NN PES is shown in Figure 1, including all calculated geometries and energies of stationary points along both the HA and OA pathways. The detailed structures and harmonic frequencies of stationary points are listed in Tables S1 and S2, respectively. The agreement with available experimental data\textsuperscript{33,34} is quite good, which is a testament of the quality of the PES. The title reaction is highly exergonic, with a reaction energy of $\sim 52.91$ kcal/mol with zero-point energy (ZPE) corrections, which allows the highest accessible vibrational level of $v = 6$ or $v = 13$ for the OH or O\textsubscript{2} product. For the reactant HO\textsubscript{2}, the OH stretching mode ($v_1$) has the highest calculated harmonic frequency of 3658.7 cm$^{-1}$, while the bending mode ($v_2$) and...
OO stretching mode ($v_3$) are 1431.9 and 1116.7 cm$^{-1}$, respectively, in reasonably good agreement with experiment.

At 200 K, which is relevant to atmospheric chemistry, the HA pathway is essentially closed for the HO$_2$ radical in the vibrational ground state, consistent with the isotopic experiment of Sridharan et al.,$^{13}$ who found no participation of the HA channel at room temperature. At room temperature, the calculated rate coefficient is $2.17 \times 10^{-11}$ cm$^3$ s$^{-1}$, which compares reasonably well with the experimental values of $5\sim6 \times 10^{-11}$ cm$^3$ s$^{-1}$,$^{13}$ further validating the accuracy of the PES.

The calculated rate coefficients of the two pathways at 200 K for HO$_2$ in both the ground and excited vibrational states are shown in Table 1. The reaction is completely dominated by the OA channel when HO$_2$ is in its vibrational ground state, as HA channel constitutes only a tiny percentage. It is also clear from the table that the excitation of either the bending ($v_2$) or OO stretching ($v_3$) mode of HO$_2$ has little effect on the rate coefficients as the HA channel remains closed. However, the HA rate coefficient increases drastically with the excitation of the OH stretching mode ($v_1$) of the HO$_2$, reactant, while the OA reaction rate decreases at the same time. When the OH mode is excited beyond its first overtone, the HA channel becomes the dominant one, accompanied by an increased total rate. Our observations are in sharp contrast to the earlier work of Varandas and co-workers, who reported almost no vibrational enhancement of the reactivity.$^{20}$

The effectiveness of the $v_1$ mode of the HO$_2$ reactant in promoting the HA channel can be readily understood in terms of the Sudden Vector Projection (SVP) model,$^{35}$ which stipulates that mode specificity of a chemical reaction is controlled by the projection of the corresponding reactant normal mode vector onto the reaction coordinate at the relevant transition state. The SVP model has been successfully applied to many direct reactions in which the sudden conditions are satisfied.$^{36}$ In Table 2, the SVP values for various reactant modes are listed for the HA channel and it can be readily seen that the reaction coordinate at SP2 is well aligned with the $v_1$ mode of HO$_2$, reflected by a large SVP value of 0.745. The impact of exciting the $v_2$ or $v_3$ mode is moderate, consistent with the slightly increased rate coefficient for this channel. On the other hand, none of the HO$_2$ vibrational modes is very effective in promoting the OA channel.

The calculated vibrational distributions for the OH and O$_2$ products from different initial states at 200 K are shown in the Figures 2 and 3, respectively. For comparison, the theoretical results of Varandas and co-workers on the DMBE I PES$^{18}$ and the DMBE II PES$^{20}$ are also included in figures, which were calculated for the ground HO$_2$ vibrational state (panel a) and for $E_{	ext{orb}} = 48.0$ kcal/mol (panel d).

For the reaction from the vibrational ground state of HO$_2$, it is clear that the OH product is almost exclusively in its vibrational ground state, while the vibration of the O$_2$ product is highly excited. Our results are similar to the 1998 values of Varandas and co-workers$^{18}$ but quite different from their 2004 results.$^{20}$ The inconsistent OH distributions in the previous theoretical calculations are probably attributable to large differences in the two DMBE PESs used in the calculations. Our product vibrational distributions can also be explained by the SVP model, which takes advantage of the microscopic reversibility in predicting the product energy disposal.$^{36}$ As shown in Table 2, the O$_2$ vibrational mode has a good overlap with the reaction coordinate at SP1 for the OA channel, which dominates the reaction under thermal conditions. For this saddle point, on the other hand, the OH vibration has a small SVP value as it is essentially a spectator in the OA channel.

For reaction with the $v_1$ mode of HO$_2$ excited, the OH product distribution becomes bimodal, as shown in Figure 3. The peak for the low-lying states is clearly correlated with the OA channel, where the OH moiety is a spectator and its excitation is largely preserved, as discussed above. On the other hand, populations at high vibrational states of the OH product are associated with the HA pathway, which can also be rationalized by the SVP model. As shown in Table 2, the O$_2$ vibrational mode has a good overlap with the reaction coordinate at SP1 for the OA channel, which dominates the reaction under thermal conditions. For this saddle point, on the other hand, the OH vibration has a small SVP value as it is essentially a spectator in the OA channel.

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Table 2. SVP Projections for the Reactant and Product Modes of the O + HO$_2$ → OH + O$_2$ Reaction for the HA and OA Channels

<table>
<thead>
<tr>
<th>Species</th>
<th>Mode</th>
<th>H abstraction</th>
<th>O abstraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reactant</td>
<td>relative translation</td>
<td>0.453</td>
<td>0.229</td>
</tr>
<tr>
<td>HO$_2$</td>
<td>OH stretch ($v_1$)</td>
<td>0.745</td>
<td>0.004</td>
</tr>
<tr>
<td>HO$_2$</td>
<td>bend ($v_2$)</td>
<td>0.464</td>
<td>0.012</td>
</tr>
<tr>
<td>HO$_2$</td>
<td>OO stretch ($v_3$)</td>
<td>0.321</td>
<td>0.001</td>
</tr>
<tr>
<td>Product</td>
<td>relative translation</td>
<td>0.258</td>
<td>0.301</td>
</tr>
<tr>
<td>OH</td>
<td>rotation</td>
<td>0.147</td>
<td>0.022</td>
</tr>
<tr>
<td>OH</td>
<td>vibration</td>
<td>0.950</td>
<td>0.006</td>
</tr>
<tr>
<td>O$_2$</td>
<td>rotation</td>
<td>0.026</td>
<td>0.236</td>
</tr>
<tr>
<td>O$_2$</td>
<td>vibration</td>
<td>0.130</td>
<td>0.540</td>
</tr>
</tbody>
</table>

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in the OA channel is 1.78 kcal/mol lower than the reactant asymptote, accessible from a vdW well, which is 3.08 kcal/mol below the reactant asymptote. For the DMBE I PES, however, the vdW well is only 1.39 kcal/mol below the reactants, which

Figure 2. Calculated vibrational distributions of the product OH from different initial vibrational excitations in the OH stretching mode ($v_1$) of the reactant HO$_2$. The previous theoretical distributions of Varandas and co-workers$^{18,20}$ are also shown for comparison.

Figure 3. Calculated vibrational distributions of the product O$_2$ from different initial vibrational excitations in the OH stretching mode ($v_1$) of the reactant HO$_2$. The previous theoretical distributions of Varandas and co-workers$^{18,20}$ are also shown for comparison.
Figure 4. Calculated rotational distributions of the product OH from different initial vibrational excitations in the OH stretching mode ($v_1$) of the reactant HO$_2$.

Figure 5. Calculated rotational distributions of the product O$_2$ from different initial vibrational excitations in the OH stretching mode ($v_1$) of the reactant HO$_2$. Only odd quantum numbers are included assuming the oxygen is the $^{16}$O isotope in the ground electronic state, for which the even quantum numbers are not allowed because of its bosonic nuclear spin.
impacts the dynamics at low collision energies. When it comes to the HA channel, the barrier height of SP2 was controversial for many years. Our new PIP-NN PES shows a very tight saddle point, which has a barrier of 6.96 kcal/mol, significantly lower than the 17.9 kcal/mol value reported by Varandas and co-workers. We have, further, examined the barrier height by recalculating the minimum energy path using a larger basis set. The barrier obtained at the level of MRCI-F12/VQZ-F12 is only 0.96 kcal/mol higher than the value of our PES, as discussed in the Supporting Information. These are only some representations of the differences between the PESs.

Figures 4 and 5 examine the rotational energy disposal in the products. It is clear that the rotational excitation in the OH and O2 products is also closely related to the two reaction pathways. For the OA channel, the rotational excitations in both OH and O2 are moderate. For the HA channel, on the other hand, OH radicals are significantly more excited in their rotational degrees of freedom. The relative contributions of the two channels to the rotational distributions are clearly observable in the figures. As shown in the Table 2, the SVP model also predicts how much the energy flows into the rotational modes. For the OA channel, the O2 rotation is more easily excited than the OH rotation, meanwhile, a reversal occurs for the HA channel. In SI, the energy partition in the products along the different reaction pathways is listed in Table S3.

The results reported here have important implications in the origin of the OH based nightglow. From our results, it is clear that the title reaction at 200 K will hardly produce any OH with v > 1. However, highly excited OH up to v = 9 can be produced if the HO2 reactant has excitation in its v1 mode. Such vibrationally hot HO2 may indeed exist in the atmosphere, via for example the OH + O3 → HO2 + O2 reaction, as pointed out by Varandas and co-workers. Although little direct evidence exists for a significant presence of v1-excited HO2 in the middle atmosphere, it is possible such scenarios are relevant in local thermodynamic disequilibrium situations. In such cases, the contribution of the HA channel should be included in the atmospheric model. For situations at thermal equilibrium, the nightglow is most likely due to the H + O3 → OH(v) + O2 reaction, which occurs on the same PES.

In summary, we investigated the detailed dynamics of the O + HO2 → OH + O2 reaction using a QCT method on a recently developed PIP-NN PES to obtain nascent ro-vibrational distributions of the two products and to understand their dynamical origins. To explore vibrational control of two reaction pathways to the same products, we show that the excitation of the v1 vibrational mode of HO2 opens the HA channel, which has a higher barrier, and significantly enhances the reaction rate. The v1 excitation of the reactant also leads to highly vibrationally exited OH product, as the reaction coordinate at the corresponding transition state is strongly aligned with the OH vibration, leading to facile energy disposal into this mode. This study thus presents an example how the vibrational excitation in the reactant controls product formation.

The elucidation of the dynamics of the O + HO2 reaction helps to shed valuable light on its role in Earth’s stratosphere and mesosphere. Under atmospheric conditions, our results indicate that the OH product is mostly in its vibration ground state with moderate rotational excitation, thanks to the spectator nature of the OH in the barrierless OA channel, which dominates the reaction at low temperatures. Hence, our results do not support the hypothesis that the title reaction produces vibrationally excited OH radicals that are responsible for OH nightglow, unless vibrationally hot HO2 are involved.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcl.2c00053.

Additional information for the potential energy surface and energy partition in the products (PDF)

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**Notes**

The authors declare no competing financial interest.

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