Quantum dynamics study of energy requirement on reactivity for the HBr + OH reaction with a negative-energy barrier

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A time-dependent, quantum reaction dynamics approach in full dimensional, six degrees of freedom was carried out to study the energy requirement on reactivity for the HBr + OH reaction with an early, negative energy barrier. The calculation shows both the HBr and OH vibrational excitations enhance the reactivity. However, even this reaction has a negative energy barrier, the calculation shows not all forms of energy are equally effective in promoting the reactivity. On the basis of equal amount of total energy, the vibrational energies of both the HBr and OH are more effective in enhancing the reactivity than the translational energy, whereas the rotational excitations of both the HBr and OH hinder the reactivity. The rate constants were also calculated for the temperature range between 5 to 500 K. The quantal rate constants have a better slope agreement with the experimental data than quasi-classical trajectory results.

The title reaction HBr + OH → Br + H2O has attracted great interest with many experimental and theoretical studies during the past several decades. From the practical aspect, this reaction plays an important role in atmospheric chemistry because it produces bromine atoms, and the bromine atoms can very effectively destroy the ozone by a catalytic cycles in the stratosphere: Br + O3 → BrO + O2, BrO + BrO → 2Br + O2. In addition, the reaction also plays a key role in combustion chemistry as some brominated compounds act as fire retardants. From experimental studies, there is a number of measurements of rate constants mainly at room temperature (298 K)1–6. Moreover, the rate constants were also measured at the temperature ranges 249–4167, 23–2958, 76–2429, 230–36010, 120–22411, 20–35012, 53–135 K13, and at a high temperature of 1925 K14. Among them, four studies4,10,11,13 have also measured the rate constants for the isotopomers system and found the primary kinetic isotope effect (KIE) is independent of temperature between 53 and 135 K. The results of these investigations reveal that the HBr + OH reaction's rate constants are extremely negative temperature-dependent below 150 K and nearly independent temperature between ~400 K and room temperature. Furthermore, Butkovskaya and Setser15 studied the vibrational distributions for H2O, HOD and D2O produced in reactions of OH and OD with HBr and DBr. Che et al.16 observed the negative collision energy dependence of reaction cross section for the HBr + OH/OD reaction in a crossed molecular beam experiment. Tsai et al.17,18 reported the orientation dependence of the Br formation and found that O-end attack is more favored for this reaction.

From theoretical studies, Clary et al.19 provided the upper limit of rate constant for HBr + OH at low temperatures and predicted a maximum rate constant with the value of 3.5 × 10^-10 cm^3 molecule^-1 s^-1 at 20 K, using the statistical adiabatic capture theory with a long-range barrierless electrostatic interaction potential. After that, Clary et al.20 reported a three-dimensional quantum scattering calculation with the rotating bond approximation on a simple potential energy surface (PES) based on a LEPS function and an accurate H2O potential. The reaction cross sections are found to be dependent on (2j + 1)^-1, where j is the initial rotational quantum number of OH. And the calculated rate constant has a T^-1/2 dependence at low temperatures. Furthermore, Nizamov et al.21 redadjusted the LEPS PES20 to fit the experimentally measured H2O vibrational energy and the thermal rate constant, they performed a quasi-classical trajectory (QCT) study on the mechanism for excitation of the bending mode and isotopic effects on the energy disposal. In 2001, Liu et al.22 investigated the dynamic properties of the...
hydrogen abstraction reaction HBr + OH over a wide range of temperatures 23–2000 K, by employing the improved canonical variational transition-state theory (VTST)\(^22\) with a small-curvature tunneling correction.

Recently, Bowman’s group\(^23\) has developed a high-quality, full dimensional PES for the HBr + OH system based on 26,000 high-level \textit{ab initio} energies. There is a van der Waals (vdW) well in the entrance channel, as well as in the product channel respectively, and a negative energy saddle-point barrier on the PES. They carried out a QCT calculation to obtain the reaction’s rate constants over the temperature range from 5 to 500 K, and found an inverse temperature dependence of rates below 160 K and a nearly constant temperature dependence above 160 K. In addition, they also studied the reaction cross section, energy disposal and rate constant for the isotopomers reaction DBr + OH\(^25\). In 2015, Ree et al.\(^26\) reported the temperature dependence of the title reaction using analytic forms of two-, three-, four-body and long-range interaction potentials in a QCT calculation over the temperature range of 20–2000 K. In 2016, Coutinho et al.\(^27\) investigated the stereodirectional dynamics of the title reaction as the prominent reason for the peculiar kinetics on a multidimensional PES mechanically generated on-the-fly\(^28\).

Till now, there have been no full-dimensional, quantum dynamics studies on the HBr + OH reaction. Thus, in this paper, we carry out the first, full-dimensional, quantum dynamics time-dependent, wave-packet study on the PES developed by Bowman’s group. Our purpose of the present work is to (1) calculate the thermal rate constants over the temperature range of 5–500 K and compare our six degrees of freedom (6DOF) results with experiments\(^13\)–\(^15\) and the QCT results\(^23\), see the relationship of the rate constants with the temperature; (2) investigate the energy efficiency of the translational, vibrational, and rotational energy on a negative-energy barrier.

In recent years, studies on energy efficacy rules for more than three atoms systems show there does not exist a unified rule on the energy efficacy to reactivity regarding the location of the transition states. For example, the O + CH\(_4\)/CD\(_4\)/CHD\(_3\) reaction with a slightly late barrier, studies\(^29\)–\(^31\) on the reactions indicate that the translational energy is more effective in surmounting the slightly late barrier. Similar to the O + CH\(_4\) reaction, the H + CH\(_4\) also has a slightly late barrier. However, the quantum dynamics calculations\(^32\),\(^33\) show that the vibrational energy is more efficient in promoting the reaction than the translational energy.

The title reaction HBr + OH has a large exoergicity with an early barrier, however, the early barrier is −0.52 kcal mol\(^{-1}\) lower than the reactant on the PES. There has been no quantum reaction dynamics studies before on the energy efficacy for the negative early barrier. Since the ground-state energy of the reactant is already higher than the barrier height, there is no barrier for the reactant to surmount, one wonders whether any form of the reactant energy (translational, vibrational or rotational energy) is equal to enhance the reactivity; if not, it would be interesting to find the energy efficacy in surmounting this negative early barrier and to see what energy efficacy rule governs this reaction system.

### Results and Discussion

#### Vibrational excitation of HBr

Figure 1(a) and (b) give the integral cross sections (ICSs) comparison for the first four vibrational excitation states of HBr (\(v_1, j_1 = 0\)) with OH (\(v_1 = 0, j_2 = 0\)) at ground state as a function of translational energy and total energy, respectively. To converge the ICSs for the initial states: \((v_1 = 0, j_1 = 0)\), \((v_1 = 1, j_1 = 0)\), \((v_1 = 2, j_1 = 0)\) and \((v_1 = 3, j_1 = 0)\), 200, 230, 260 and 260 partial waves are calculated, respectively. For the partial waves of \(J \leq 100\), the reaction probability for every partial wave was calculated explicitly, and the reaction probabilities for different partial waves of \(J > 100\) were computed using the \(J\)-shifting method\(^24\) with a \(J\) interval of 5. The standard centrifugal sudden (CS) approximation\(^5\),\(^26\) was employed in calculation for \(J > 0\). This figure shows the cross sections decrease as the translational energy increases. Especially for the excited state ICSs, \(v_1 = 1, 2, 3\), the cross sections decrease significantly about 75% as the translational energy goes from 0.05 to 0.3 eV. On the other hand, the ground state cross section drops slower only about 45% for the same energy. Among these four cross sections, the ground state ICS is the smallest. For the translational energy lower than 0.1 eV, the amplitudes of the HBr \(v_1 = 1, 2, 3\) ICSs are about 3–4 times bigger than that of ground state; even for the collision energy is larger than 0.1 eV, the three excited-state ICSs are also about 2 times bigger than the ground state’s. As this reaction has a negative barrier height, even the ground state energy is higher than the barrier, it is surprising to see that the higher of the excited state the more reactive of this reaction.

In order to see the energy efficacy of the vibrational energy of HBr on the reactivity. We plot the ICS ratios of the HBr, \(\sigma(v_1 = 1)/\sigma(v_1 = 0)\), \(\sigma(v_1 = 2)/\sigma(v_1 = 1)\) and \(\sigma(v_1 = 3)/\sigma(v_1 = 2)\), on the basis of an equivalent amount of total energy in Fig. 2. The ICS ratio of \(\sigma(v_1 = 1)/\sigma(v_1 = 0)\) has a maximum ~10.8 at the initial translational energy 0.037 eV, then rapidly drops to 3.5 at the 0.145 eV until it reaches to 2.2 at 0.283 eV. In the whole energy range, the ratio is always bigger than 1.0, which means that vibrational energy is more effective to promote the reaction than translational energy. Furthermore, the ratio of \(\sigma(v_1 = 1)/\sigma(v_1 = 0)\) is much bigger than those of \(\sigma(v_1 = 2)/\sigma(v_1 = 1)\) and \(\sigma(v_1 = 3)/\sigma(v_1 = 2)\), and the ratio of \(\sigma(v_1 = 3)/\sigma(v_1 = 2)\) is just slightly larger than \(\sigma(v_1 = 2)/\sigma(v_1 = 1)\). Bases on the above results, we can conclude that the vibrational excitation from the ground state to the first excited state is the most effective one to promote the reactivity; however, there is no much reactivity change as the vibrational quantum numbers increase from \(v_1 = 1\) to \(v_1 = 2\) and from \(v_1 = 2\) to \(v_1 = 3\). Nevertheless, the comparison of the ICS ratios on the equal amount total energy indicates that the vibrational energy of HBr is more effective than translational energy on promoting the reactivity for this negative-barrier reaction.

#### Vibrational excitations of OH

In Fig. 3(a) and (b), we also compares the ICSs for those vibrational excitation states of OH with HBr (\(v_1 = 0, j_1 = 0\)) at ground state as a function of translation energy and total energy, respectively. There are 215, 210 and 210 partial waves needed to converge the vibrational excitation state of OH (\(v_2 = 1, j_2 = 0\), \(v_2 = 2, j_2 = 0\), \(v_2 = 3, j_2 = 0\), respectively. The results show that ICSs almost stick together in regard to the translation energy, and the ICSs decrease as the translational energy increases. The reaction path of the PES\(^24\) we used here has a vDW minimum at the entrance channel with a structure of the O–end of OH linked to HBr, HOD–HBr. This means the favorite route of this reaction is the two reactants enter the entrance vDW
minimum to form HO·HBr, then scale the transition state to make the reaction occur. This has been confirmed by the crossed beam scattering experiment\textsuperscript{17,18} by Tsai et al. They found the orientation dependence for the title reaction that the reaction is favored by OH re-orientating its O-end to face the HBr. Since the barrier height

Figure 1. The integral cross sections of the reaction HBr ($v_1, j_1 = 0$) for $v_1 = 0, 1, 2, 3$ with OH ($v_2 = 0, j_2 = 0$) at ground state as a function of translational energy (a) and total energy (b), respectively.

Figure 2. The ICS ratios of $\sigma(v_1 = 1)/\sigma(v_1 = 0)$, $\sigma(v_1 = 2)/\sigma(v_1 = 1)$ and $\sigma(v_1 = 3)/\sigma(v_1 = 2)$ of HBr with OH at ground state as a function of translational energy on the equal amount of total energy.
respect to the vdW minimum is 0.12 eV on the PES, thus as seen in Fig. 3(a), the reaction ICSs are almost the same at the translational energy larger than ~0.15 eV because the reactants would overpass the vdW minimum without reorientation in this large energy range; however, for the translational energy less than ~0.15 eV, the reactants will enter the vdW minimum to re-orientate themselves then surmount the barrier, therefore the ICSs are bigger for translational energy smaller than 0.15 eV. In order to investigate the vibrational energy efficacy of the OH, we need to check the ICS ratio of the excited state over the ground state in terms of equal amount of the total energy.

Figure 4 gives the ICS ratio, $\sigma(v_2 = 1)/\sigma(v_2 = 0)$, in terms of translational energy at the equal amount of total energy. This figure shows that the ratio is larger than 1 in the whole energy range and the ICS ratio curve of $\sigma(v_2 = 1)/\sigma(v_2 = 0)$ is very similar to the HBr's. The ratio is also inversely proportional to the energy and has a rapid decline at lower energy. This shows that the vibrational energy of OH is also much more efficient than the translational energy in promoting the reaction. Compared with the $F + CH_4$ and $F + H_2O$ reaction, the title reaction $HBr + OH$ has some similarities with them in regard to the PES. The three reactions all have an early saddle point located in the reactant channel, a vdW well in the entrance channel, and a relatively deep vdW minimum in the product valley. And the difference is that the $HBr + OH$ reaction has a negative energy barrier ($-0.52$ kcal/mol) on the PES, while $F + CH_4$ barrier height is 0.5 kcal/mol and $F + H_2O$'s is 3.8 kcal/mol. For the early barrier reaction $HBr + OH$, our calculation shows both the vibrational energies of HBr and OH are more effective than translational energy in enhancing the reactivity. While for the two early barrier of $F + H_2O$ and $F + CHD$, reactions, the study of $F + H_2O$ shows that the vibrational energy of $H_2O$ has higher efficacy in enhancing the reactivity than the translational energy; however, the vibrational excitation of C-H stretching motion of CHD, hinders the overall reaction rate. Thus, these investigations further prove that, Polanyi rules in which the translational energy is more effective to raise the reactivity for the early-barrier tri-atomic reaction systems cannot be extended to the ploy-atomic reaction systems. Nonetheless, this study shows that, for this negative, early barrier reaction, the vibrational energy is more efficient than the translational energy in promoting the reactivity.

Figure 3. The integral cross sections of the reaction HBr ($v_1 = 0, j_1 = 0$) at ground state with OH ($v_2, j_2 = 0$) for $v_1 = 0, 1, 2, 3$, as a function of translational energy (a) and total energy (b), respectively.
Rotational excitations ICSs. In addition, we also studied the rotational excitations on the reactivity for this reaction. For all the excited rotational ICSs' calculations, 200 partial waves were needed to converge the excited rotational ICSs, and the CS approximation\(^3\) was also used to calculate the partial wave reaction probabilities for \(J > 0\). Similar to the vibrational ICSs' calculations, for the reaction probabilities of partial waves for \(J \leq 100\), every \(J\) partial wave was calculated; and for \(J > 100\), the reaction probabilities were computed using the \(J\)-shifting method\(^4\) with a \(J\) interval of 5. Figure 5 presents the first five rotational excited ICSs of the HBr \((v_1 = 0, j_1)\) with OH \((v_2 = 0, j_2 = 0)\) at the ground state as a function of translational energy. As the figure shows that the ground state has the largest ICS among the 5 ICSs, and as the rotational quantum numbers \(j_1\) increases, these excited rotational states' ICSs significantly decrease. So the rotational-excited modes of the HBr greatly hinder the reactivity.

In Fig. 6, the first four ICSs of rotational excitations and the ground state of the OH were compared. It is shown that overall the OH rotational excitations greatly inhibit the reactivity, and the faster of the rotation, the smaller of the ICS. This can be explained due to the fact that the OH plays a receiver role; the faster rotation of OH will further add difficulty for H atom in HBr to attack the O atom in OH. And our 6DOF quantum dynamics results here are in agreement with the three-dimensional quantum scattering calculations by Clary et al.\(^5\) who found the reaction cross sections are proportional to \(2j + 1\) \(^{-1}\), where \(j\) is the initial rotational quantum number of OH.

Overall, the rotational excitations, both HBr and OH, hinders the reactivity. This indirectly proves Tsai et al. molecular beam study\(^6\) that Br formation of this reaction has orientation dependence which favors the O-end attack. On one hand, the faster rotation of HBr will make H in HBr cannot attack O-end easily; on the other hand, the faster rotation of OH will make O-end having difficulty to receive the H in HBr. Therefore, both the rotational excitations of HBr and OH hinder the reactivity.
Thermal rate constants. By summing over all the ro-vibrational states of HBr and OH, we can obtain the 6DOF cumulative reaction probability (CRP). In order to converge the rate constants up to 500 K to compare with the experimental results\(^2\)–\(^{13}\), the reaction probabilities of the ground vibrational state including 6 HBr rotational excitation states \((j_{1\text{max}} = 5)\) and 3 OH rotational excitation states \((j_{2\text{max}} = 3)\) were calculated.

The thermal rate constants are obtained using the \(J-K\) shifting rate expression from equation (8) in the Method Section. Note, in the QCT calculation by Bowman's group\(^{24}\), they compared their calculated rates, with the OH spin-orbit coupling (RR/SO rates), without the spin-orbit coupling (RR/nSO rates), and with a fully coupled partition function (Coupled rates), with the experimental measured ones. They found that, with the spin-orbit coupling neglected, the RR/nSO rates have the best overall agreement with the experimental results. Therefore, in the current study, we neglected the spin-orbit coupling to calculate our 6DOF quantal rate constants. In Fig. 7, our 6DOF results are compared with experimental\(^{2\text{–}13}\) and QCT results (RR/SO, RR/nSO, Coupled)\(^{24}\). And the QCT calculations of RR/SO, RR/nSO and Coupled are obtained from three different approaches to treat the reactant OH rotational and associated electronic partition function. As the comparison shows, our 6DOF rate constants have a good agreement with the experimental data and demonstrate a negative temperature dependence, which is in agreement with the experimental ones and the QCT results. However, at the very low temperature range up to about 50 K, our 6DOF results are bigger than the experimental and QCT results. Nonetheless, in general, our 6DOF results have a better agreement with the slope of the experimental data than the QCT results. In addition, our 6DOF results give a maximum at about 15 K just as RR/SO and RR/nSO results do. This agrees with Clary et al.'s\(^{19}\) prediction that a maximum rate constant should appear at 20 K. On the whole, Fig. 7 shows that our 6DOF rate constant has a better slope agreement with the experiments than the QCT results except at the extreme low temperature.
Discussion

In this work, we carried out a 6DOF quantum reaction dynamics, time-dependent wave packet propagation approach to study the HBr + OH → Br + H₂O reaction system on the PES developed by Bowman's group. This is the first, full-dimensional, quantum dynamical study on the title reaction. For the HBr + OH reaction system with a negative-early barrier, this study shows that not all forms of energy are equal in enhancing the reactivity. Even the ground state energy of the reactant is higher than the barrier, the calculation still shows that vibrational excitations of both the HBr and OH vibrational enhance the reactivity. Furthermore, the HBr and OH vibrational excitations are more effective in enhancing the reactivity than the translational energy. We also studied the rotational excitations of HBr and OH. The results show that both the rotational excitations hinder the reactivity. This is due to the fact that the faster rotation of HBr makes H having difficult to attack O in OH; and the faster rotation of OH makes O having difficult to receive H from HBr. Comparing with other two early barrier reaction systems, F + H₂O and F + CHD₃, we can see there are no general rules so far on the energy efficacy for the ploy-atomic reaction systems as the Polanyi rules do to the tri-atomic systems. We think this is due to the complexity of the PESs of the poly-atomic systems with vdW wells usually in both the reactant and product channels. These wells, especially the entrance channel well before the transition state, might also play a role that cannot be neglected on energy efficacy on surmounting the energy barrier.

Furthermore, the comparison of the thermal rate constants between our 6DOF quantum results and the experimental display that our data agree well with experimental measurement except at extreme low temperatures.

Theoretical Methods

6DOF approach. We performed a full dimensional, 6DOF time-dependent quantum dynamics study for the HBr + OH → Br + H₂O reaction. The 6DOF Hamiltonian in the reactant Jacobi coordinates, as shown in Fig. 8, can be written as,

\[ H = -\frac{\hbar^2}{2\mu_R} \frac{\partial^2}{\partial R^2} + \frac{(1 - J^2)}{2\mu_R R^2} + h_1(r_1) + h_2(r_2) + \frac{j_1^2}{2\mu_1 r_1^2} + \frac{j_2^2}{2\mu_2 r_2^2} + V_6D(R, r_1, r_2, \theta_1, \theta_2, \phi), \]

where, \( \mu_R \) is the reduced mass of the whole reaction system; \( R \) is the center-of-mass distance between HBr and OH, \( r_1 \) is the distance of H-Br and \( r_2 \) is the distance of O-H; \( \theta_1 \) and \( \theta_2 \) are the two Jacobi angles between \( r_1 \) and \( R \) and \( r_2 \) and \( R \), \( \Phi \) is the torsion angle; \( J \) is the total angular momentum operator of the reaction system, \( j_1 \) and \( j_2 \) are the rotational angular momentum operators for HBr and OH, respectively, \( j_{12} \) is the coupled angular momentum operator of \( j_1 \) and \( j_2 \); and \( V_{6D} \) is the interaction potential. The vibrational reference Hamiltonians \( h_1(r_1) \) and \( h_2(r_2) \) are defined as

\[ h_1(r_1) = -\frac{\hbar^2}{2\mu_1} \frac{\partial^2}{\partial r_1^2} + V(r_1), \]  

\[ h_2(r_2) = -\frac{\hbar^2}{2\mu_2} \frac{\partial^2}{\partial r_2^2} + V(r_2), \]

Here \( V(r_1) \) and \( V(r_2) \) are the one-dimensional reference potentials for HBr and OH, and \( \mu_1 \) and \( \mu_2 \) are the corresponding masses. These potentials correspond to the reactant at the asymptotic region with other coordinates fixed at the equilibrium geometry.

The split-operator method⁴³ is employed here to propagate the wave packet on the full-dimensional \textit{ab initio} PES for the quantum scattering calculation. And we expand the time-dependent wave-function in terms of the body-fixed (BF) rovibrational eigenfunctions defined in terms of the above reactant Jacobi coordinates. After the time-dependent wave function is propagated into the product region, we perform the standard reactive flux⁴⁴–⁴⁷ method to extract the initial-state-selected reaction probability. To obtain the initial-state-selected ICS, we first

![Figure 8. Reactant Jacobi coordinates for the reaction HBr + OH.](image)
calculated the partial wave reaction probabilities for the different initial total angular $J$. Then the ICS $\sigma_{v_0 j_0} (E)$ is obtained by summing over all the initial-state-selected reaction probability $P_{v_0 j_0 K_0}^J$ for all partial waves

$$\sigma_{v_0 j_0} (E) = \frac{1}{2J_0 + 1} \sum_{J}^\pi (2J + 1) P_{v_0 j_0 K_0}^J (E),$$

(3)

where $k = (2\mu E)^{1/2}$ is the wave number and $E$ is the translational energy, $v_0$ and $j_0$ denotes, respectively, the initial vibrational and rotational quantum numbers, $K_0$ is the projection of $J$ onto BF z axis of the reaction system.

**Cumulative reaction probability and thermal rate constant.** The CRP $N^{j=0} (E)$ is defined as the sum of all the initial state selected ro-vibrational reaction probabilities $P_{v_0 j_0 K_0}^J (E)$

$$N^{j=0} (E) = \sum_{v_0 j_0} \sum_{K_0} P_{v_0 j_0 K_0}^J (E),$$

(4)

Next, the $J$-shifting method$^{44}$ is employed here to calculate the CRP for the nonzero total angular momentum $J$.

The total CRP, $N(E)$ is defined as the sum of CRPs for all the open $J$ and $K$ channels

$$N (E) = \sum_{J=0}^j (2J + 1) \sum_{K=-J}^J N^{j=0} (E - E_{JK}^J),$$

(5)

where $E_{JK}^J$ is the rigid rotor rotational energy of the reaction system at the transition state. This energy is approximated by the expression for a symmetric top molecule,

$$E_{JK}^J = B^J (J + 1) + (A^J - B^J) K^2,$$

(6)

$A^J$ and $B^J$ are the rotational constants of OHHBr at the transition state.

Thus the thermal rate constant can be computed as

$$k(T) = \frac{1}{h Q_{\text{rot}}} \int_0^\infty N (E) e^{-\frac{E}{k T}} dE,$$

(7)

where $Q_r (T)$ is the reactant partition function, which is written as a product of vibrational, rotational, and translational partition functions. The equation (7) simplifies under the $J$-$K$ shifting approximation in terms of Equation (5),

$$k(T) = \frac{Q_r^J}{h Q_{\text{rot}}} \int_0^\infty N^{j=0} (E) e^{-\frac{E}{k T}} dE,$$

(8)

where $Q_{\text{rot}}^J$ is the rotational partition function of the reaction system at the transition state.

**Numerical aspects.** To converge the above 6DOF, wave-packet, quantum dynamics calculation, we used the following numerical parameters to expand the wavefunction: for the translational coordinate $R$ from 2.5 to 12.5 bohr, 240 sine basis functions are used, and among these, 150 are used for the interaction region; 30 potential-optimized vibrational discrete variable representation (DVR) points$^{48}$ are chosen for the $r_1$ coordinates in the range from 1.6 to 5.5 bohr; 40 spherical harmonic rotational functions are used for $\theta_1$ and 15 for $\theta_2$, which gives 4896 coupled parity adapted total angular momentum basis. The time-dependent wave packet is propagated for a total time of about 16,000 atomic unit time with a time step of 15 a.u.

For the thermal rate constant calculation in equation (6), the rotational constants $A^J$ and $B^J$ of OHHBr at the transition state are 15.46 cm$^{-1}$ and 0.14 cm$^{-1}$. For the reactant HBr and OH’s vibrational partition function, the used harmonic vibrational frequencies of HBr and OH are 2525 cm$^{-1}$ and 3611 cm$^{-1}$. For reactant rotational partition function, the HBr’s rotational constant is 8.46 cm$^{-1}$ and the OH’s 18.86 cm$^{-1}$.

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**Author Contributions**
D.Y.W. supervised the projects, Y.W. performed the time-dependent quantum dynamics calculations. Y.L. helped the analysis of calculation data. D.Y.W. and Y.W. wrote the paper.

**Additional Information**

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