On the theory of the CO+OH reaction, including H and C kinetic isotope effects

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The effect of pressure, temperature, H/D isotopes, and C isotopes on the kinetics of the OH + CO reaction are investigated using Rice-Ramsperger-Kassel-Marcus theory. Pressure effects are treated with a step-ladder plus steady-state model and tunneling effects are included. New features include a treatment of the C isotope effect and a proposed nonstatistical effect in the reaction. The latter was prompted by existing kinetic results and molecular-beam data of Simons and co-workers [J. Phys. Chem. A 102, 9559 (1998); J. Chem. Phys. 112, 4557 (2000); 113, 3173 (2000)] on incomplete intramolecular energy transfer to the highest vibrational frequency mode in HOCO. In treating the many kinetic properties two small customary vertical adjustments of the barriers of the two transition states were made. The resulting calculations show reasonable agreement with the experimental data on (1) the pressure and temperature dependence of the H/D effect, (2) the pressure-dependent 12C/13C isotope effect, (3) the strong non-Arrhenius behavior observed at low temperatures, (4) the high-temperature data, and (5) the pressure dependence of rate constants in various bath gases. The kinetic carbon isotopic effect is usually less than 10 per mil. A striking consequence of the nonstatistical assumption is the removal of a major discrepancy in a plot of the $k_{\text{OH+CO}}/k_{\text{OD+CO}}$ ratio versus pressure. A prediction is made for the temperature dependence of the OD+CO reaction in the low-pressure limit at low temperatures. © 2005 American Institute of Physics. [DOI: 10.1063/1.2031208]

I. INTRODUCTION

The reaction of CO with the OH radical plays a major role in both atmospheric and combustion chemistry.1,2 The reaction is known to be the principal oxidation which converts CO to CO$_2$ in the atmosphere,1 and is also the key reaction controlling the OH radical concentration in the lower atmosphere.1 Reflecting its importance and the unusual temperature and pressure dependence of its rate constant, the CO+OH→CO$_2$+H reaction has been examined extensively in many experimental3–17 and theoretical studies,17–28 including experimental studies over a very wide range of temperatures and pressures.4–13,17 Nevertheless, carbon and oxygen isotope effects have been observed29–33 and the anomalous effects for these heavy-atom isotopes have not yet been treated in the literature. Again, Simons and co-workers34–36 observed in their molecular-beam study of the reverse reaction, H+CO$_2$→OH+CO, that the vibrational excitation of the CO product was far below that expected from statistical theory for the HOCO intermediate. While the energies in their experiments are higher than those normally occurring in the OH+CO reaction, it is interesting to look for other anomalies in that reaction which may be better explained by a nonstatistical modification of the Rice-Ramsperger-Kassel-Marcus (RRKM) theory. Furthermore, there is a discrepancy in the H/D isotope effect in the literature.27 The present work treats these various observations.

The radical intermediate HOCO has been observed both in the gas phase and in a matrix.16,34–36 The ionization potential, energy of formation, and vibrational and rotational spectroscopy of the radical have also been characterized.3,35 Studies of the kinetics of HOCO formation have firmly established the connection between the reactants (OH+CO) and the products (H+CO$_2$).16 The currently accepted reaction mechanism includes an OH and CO bimolecular association step producing a vibrationally excited trans-HOCO, followed by cis-trans isomerization. The final reaction steps in competition with each other are dissociation to H and CO$_2$, back reaction to OH and CO, and collisional stabilization.5,13,16,17,21,22 As the pressure increases, the collisional stabilization of the cis- and trans-HOCO intermediates competes favorably with the dissociation channel and the back reaction. When OH and CO react in oxygen, both the dissociation channel and the collision stabilization lead to HO$_2$ and CO$_2$, because of the follow-up reactions of H and HOCO with O$_2$.37–38 The present paper treats the dependence of the rate constants on temperature, pressure, foreign gases, and H/D and 12C/13C isotope effects. Interpretation of the oxygen isotope effects31–33 is complex and is discussed in a second paper.

The experimental study of the OH+CO reaction covers a temperature range from 80 to 3150 K.4–13,17 A marked change of activation energy occurs near 500 K. The pressure and bath gas dependence of the reaction rate have also been measured,11–15,17,40 as has the pressure and temperature dependence of the OD+CO reaction.11,17,41 The $k_{\text{OH+CO}}/k_{\text{OD+CO}}$ ratio shows a pressure-dependent H/D isotope effect. Further
experimental kinetic isotopic studies with $^{18}$OH, $^{13}$CO, C$^{17}$O, and C$^{18}$O have also been performed. These results provide a body of data for tests of ab initio or other potential-energy surfaces and of reaction-rate theory. In particular, there is a serious discrepancy between the observed and calculated plot of OH/OD rates versus pressure.

Stevens et al. appear to be the first group to study experimentally the kinetic isotope effects for carbon and oxygen in the reaction. They measured the isotopic ratios of the unreacted CO and of the CO$_2$ formed in the reaction mass spectrometrically and obtained pressure-dependent fractionation of $^{13}$C/$^{12}$C and $^{18}$O/$^{16}$O at pressures from 150 to 800 torr. In a similar but independent study Smit et al. obtained results for the fractionation between carbon isotopes, which were consistent with those of Stevens et al. Their investigations were prompted by the need for understanding the global budget of atmospheric $^{12}$CO and $^{13}$CO. The enrichment of oxygen isotopes in CO was found to be nonconventional in its mass dependence, i.e., it did not obey the usual “mass-dependent” theoretically based and usually applicable rule of Bigeleisen and Mayer.

Results similar to those of Stevens et al. were obtained by Röckmann et al. Laboratory studies of isotope effects were also made by Feilberg et al. whose results do not agree with those of Röckmann et al. and Stevens et al. The experimental precursors generating the OH radicals are different. The experimental conditions of Röckmann et al. are simpler in that ozone was not added and so there were fewer chemical reactions in the overall scheme. While HO$_2$ may be a dominant product in the studies of Stevens et al. and Röckmann et al., due to the reaction of the source material H$_2$O$_2$ with OH, the rate constant for the reaction of CO with HO$_2$ is extremely small, less than $10^{-27}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ at room temperature, and so is about 14 orders of magnitude smaller than that of CO with OH. The HO$_2$ is therefore not expected to interfere in the analysis.

Dynamical and statistical treatments of the CO+OH reactions have been made with various theoretical methods, including quasiclassical trajectories, quantum dynamics, transition state theory, and RRKM theory. In the present article we focus on the statistical and nonstatistical aspects of the reaction over a wide temperature region for the different experimental conditions, as well as the hydrogen and carbon isotope effects.

The carbon isotope effect is typically less than 10 per mil at these pressures and temperatures. In the present study, RRKM theory and a nonstatistical modification are used, together with nuclear tunneling corrections and a step-ladder model for collision stabilization. Like the previous study by Senosiain et al., the resulting theory reproduces a large body of experimental data, apart from the potentially complex O-isotope effect discussed elsewhere. In addition, we treat the C-isotope effect and the large OH/OD rate constant discrepancy described in Ref. 27.

The paper is organized as follows: The potential-energy surface used is summarized in Sec. II A. The kinetic scheme for RRKM calculations, approximate tunneling probabilities, step-ladder model for pressure effects, and a possible nonstatistical distribution model for energy transfer at the exit channel are treated in Secs. II B–II E, respectively. Results are given in Sec. III and discussed in Sec. IV.

### II. CALCULATION METHODS

#### A. Potential-energy surface

An ab initio potential-energy surface (PES) was calculated by Zhu et al., who used the modified GAUSSIAN-2 method (G2M). To obtain the more accurate vibrational frequencies and rotational constants of all stable and TS structures for various isotopes we use a coupled-cluster method, abbreviated as CC. The calculated geometries and vibrational frequencies of normal isotopes are listed in Table I. To test the robustness of certain calculations, particularly of the H/D isotope effects, the results are compared in Appendix A with another modified ab initio potential by Lakin-Troya-Schatz-Harding (LTSH) by Lakin et al. and with two other ab initio methods, the MP2 method and a density-functional method (B3LYP), for the vibrational and rotational constants. The GAUSSIAN 98 program was employed for all ab initio calculations to obtain the principal stable and TS equilibrium structures and their rotational and vibrational constants.

#### Table I. Calculated geometries and vibrational frequencies of normal isotopes by CC. The units are in Å for bond length, in degree for angles and dihedral angles, and in cm$^{-1}$ for vibrational frequencies.

<table>
<thead>
<tr>
<th></th>
<th>OH+CO$'$</th>
<th>TS$_0$</th>
<th>cis-HOCO$'$</th>
<th>trans-HOCO$'$</th>
<th>TS$_0$</th>
<th>CO$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R_{OH}$</td>
<td>0.976</td>
<td>0.975</td>
<td>0.976</td>
<td>0.967</td>
<td>1.345</td>
<td></td>
</tr>
<tr>
<td>$R_{CO}$</td>
<td>1.148</td>
<td>2.033</td>
<td>1.342</td>
<td>1.358</td>
<td>1.227</td>
<td>1.177</td>
</tr>
<tr>
<td>$R_{CO'}$</td>
<td>1.150</td>
<td>1.199</td>
<td>1.194</td>
<td>1.184</td>
<td>1.177</td>
<td></td>
</tr>
<tr>
<td>$\theta_{HOC}$</td>
<td>94.86</td>
<td>107.74</td>
<td>107.37</td>
<td>116.27</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\theta_{OCO'}$</td>
<td>123.52</td>
<td>129.61</td>
<td>126.38</td>
<td>155.28</td>
<td>180.00</td>
<td></td>
</tr>
<tr>
<td>$d_{HOCO'}$</td>
<td>180.00</td>
<td>0.00</td>
<td>180.00</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$v_1$</td>
<td>2154$^a$</td>
<td>312$^i$</td>
<td>596</td>
<td>533</td>
<td>2125$^i$</td>
<td></td>
</tr>
<tr>
<td>$v_2$</td>
<td>3763$^b$</td>
<td>197</td>
<td>594</td>
<td>606</td>
<td>518</td>
<td></td>
</tr>
<tr>
<td>$v_3$</td>
<td>244</td>
<td>1091</td>
<td>1090</td>
<td>640</td>
<td>1345</td>
<td></td>
</tr>
<tr>
<td>$v_4$</td>
<td>752</td>
<td>1321</td>
<td>1262</td>
<td>945</td>
<td>2410</td>
<td></td>
</tr>
<tr>
<td>$v_5$</td>
<td>2122</td>
<td>1868</td>
<td>1903</td>
<td>1298</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$v_6$</td>
<td>3784</td>
<td>3707</td>
<td>3869</td>
<td>2157</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

$^a$POH
$^b$PCO
A major candidate for the origin of the strong non-Arrhenius effect in the rate constant is the tunneling at the transition state TS₂ in Fig. 1 for forming H+CO₂ from cis-HOCO. This interpretation is consistent with there being a large imaginary frequency at TS₂, 2126i cm⁻¹ calculated by CC. The RRKM rate constant in the low-pressure limit at room temperature calculated with this PES was larger than the experimental value by a factor of about 2. Accordingly, to match the experimental and calculated rate constants, the potential energy of TS₂ was shifted vertically slightly by two independent constants, E₁ at TSₐ and E₂ at TS₋. (Vertical shifts are commonly used in conjunction with ab initio surfaces.) E₁ and E₂ were determined from the rate constants for the CO+OH reaction at 100 and 300 K in the low-pressure limit (Sec. III A). At lower temperatures the barrier at TSₐ dominates the rate constant due to the large tunneling at TS₂. At room temperature the barriers in both exit and entrance channels of HOCO⁺ influence the rate constant. With these two constants, E₁ and E₂, and an energy-transfer parameter α per collision (Sec. II D), all the remaining properties of the reaction were calculated and compared with the experimental data. When different methods of varying accuracy were used and the shifts E₁ and E₂ were made to fit two pieces of data above, the resulting calculated rate constants are similar in all methods, as shown in Appendix A.

**B. Kinetic scheme and rate constants**

Microcanonical RRKM theory is used with tunneling corrections⁴⁸ for each quantum state of the two transition state structures. The steady-state equation for pressure effects is then solved. The coordinate assumed for the transition state structures is the center-to-center distance, and hence, of the two largest moments of inertia. (A more elaborate variational calculation of selected points will be given elsewhere.) There are van der Waals complexes of CO and OH and the loose transition structures to form the TSₐ, but calculations show that they have only a minor effect on the reaction rates.⁴⁹ Thus, TSₐ is the only transition state structure for the entrance channel included in the kinetic scheme. The potential-energy barrier between trans- and cis-HOCO is much lower than the potential energies of the entrance and exit channels, and so the two isomers interconvert in the energetic HOCO⁺ intermediate. The density of states of HOCO⁺ is therefore evaluated by summing ρ_{cis-HOCO}(EJ) and ρ_{trans-HOCO}(EJ), which includes the hindered rotation as one of the coordinates in the calculation. In the kinetic scheme, we can simply write HOCO⁺ for the sum of cis-HOCO⁺ and trans-HOCO⁺. At zero pressure, the following scheme for the reaction of CO+OH is used for reaction at any energy E and total angular momentum J:

\[
\begin{align*}
\text{CO} + \text{OH} & \rightarrow \text{HOCO}^+(EJ), \\
\text{HOCO}^+(EJ) & \rightarrow \text{H} + \text{CO}_2.
\end{align*}
\]

At finite pressures an additional series of kinetic equations are added to the scheme:

\[
\omega(EJ → E'J') \text{HOCO}^+(EJ) + M \rightarrow \text{HOCO}(E'J') + M,
\]

where M is a third body and E and J are conserved in reaction steps (1) and (2). The rate constants k₁(EJ) are all E and J dependent. The ω(EJ → E'J') is the rate constant per unit E for forming HOCO at (E'J') from HOCO(EJ) by collision with a third body M.

The rate equation for the distribution function, g(EJ), for HOCO⁺ at any specified (EJ) per unit energy is

\[
\frac{dg(EJ)}{dt} = k₁(EJ) - [k₁(EJ) + k₂(EJ) + \omega]g(EJ)
\]

\[
+ \sum_{E'} \int E' \omega(E'J' → EJ) g(E'J') dE',
\]

where g(EJ) denotes [HOCO(EJ)]/[CO][OH]. The latter has units of (energy concentration)⁻¹. The total collision frequency ω is

\[
\omega = \sum_{J'} \int E' \omega(EJ → E'J') dE'.
\]

This ω(EJ → E'J') is described by a step-ladder model employed in Sec. II D.

In the dissociation of the intermediate via TSₜₚ, the RRKM rate constant k₂(EJ) appearing in Eqs. (2) and (4) is

\[
k₂(EJ) = \frac{N_a(EJ)}{\hbar \rho(EJ)},
\]

where ρ(EJ) is the density of states of the corresponding dissociating energetic molecules and N_a(EJ) is the sum of states of TSₜₚ, with the tunneling correction⁵⁰,⁵¹ for a microcanonical ensemble of
\[ N_d(EJ) = \int_{-\infty}^{E} \kappa_d(E') \rho_d(E-E')dE'. \]  

Here, \( \kappa_d(E') \) is the tunneling transmission probability of \( TS_d \) when the energy in the H-tunneling coordinate is \( E' \), and \( \rho_d(E-E')d \) denotes the density of states of \( TS_d \) at an energy \( E-E' \) and at total angular momentum \( J \).

Harmonic vibrational frequencies and rotational constants for counting \( N(EJ) \) or \( \rho(E) \) were obtained by electronic structure calculations described in Sec. II A. To account approximately for the anharmonicity of the energetic intermediates, the density of states of \textit{trans}- and \textit{cis}-HOCO were increased by a constant factor, 1.5.52 For TS, a “loose” TS was assumed. Anharmonicity effects for the vibrations of OH and CO are minor. In the calculation of quantum states of the internal rotation about the HO\textendash{}CO bond in TS,53 \textit{cis}- and \textit{trans}-HOCO were treated as one-dimensional hindered rotors.

C. Tunneling calculations with statistical and nonstatistical models

All \textit{ab initio} calculations show a sharp potential-energy barrier in the product channel.18,21 However, the magnitude of the imaginary frequency of \( TS_d \) depends on the electronic structure method used. Although these values are substantially different from each other (they are tabulated later), all of them indicate that the tunneling must be included in calculating the low-temperature reaction rates.54 In these systems, in mass-weighted coordinate space the tunneling through the potential barrier occurs entirely in the products’ channel and so there is no “corner cutting.”55 As a first approximation to reduce computational time, the tunneling probability \( \kappa(E) \) was estimated for passage through an Eckart potential.56 The potential was fitted to the paths of steepest ascent and descent through the saddle point. The Eckart-fitted approximation was checked with a sampling of more accurate calculations for selected points. In general, the Eckart barrier provides a more accurate but still simple representation than an inverted parabolic potential. However, the tunneling probability in the significant tunneling regions calculated by the Wigner approximation with an inverted parabolic potential is very similar to that calculated using the Eckart approximation. The difference of tunneling probabilities is less than 10%. However, compensation by barrier-height adjustment almost completely removes this discrepancy. Therefore, the bottom shape of the barrier in the exit channel only has a minor effect in the tunneling effect. With a slightly different vertical shift \( E \) of \( TS_d \) such difference have only a minor effect over the final results.

D. Pressure effects with statistical and nonstatistical models

At high temperatures the effect of pressure on all properties in the present region of focus, below 1000 torr, is minor both experimentally and theoretically. However, at low temperatures in this pressure region both the pressure and the nature of the foreign gas affect the reaction rate. The collision frequency for the transition from \( EJ \) to \( E'J' \), \( \omega(EJ \rightarrow E'J') \), is approximated as the product of the total collision frequency \( \omega \), the energy transfer probability \( P_{E \rightarrow E'} \), and the rotational angular momentum transfer probability \( P_{J \rightarrow J'} \): \( \omega(EJ \rightarrow E'J') = \omega P_{E \rightarrow E'} P_{J \rightarrow J'} \). For angular momentum transfer a strong collision is assumed. Thereby, the transfer probability \( P_{J \rightarrow J'} \) from \( J \) to state \( J' \) equals the thermal distribution of \( J' \) states of the intermediate, HOCO, at the given temperature.

In the step-ladder model used for the energy-transfer probability \( P_{E \rightarrow E'} \),57 a certain amount of energy \( \alpha \) is transferred in each collision, in the present case between HOCO7 and a bath molecule. The energy-transfer parameter \( \alpha \) (\( \alpha \) is positive) is dependent on the bath gas, but is independent of pressure. The \( \alpha \) value of each bath gas, determined by best agreement with experimental results11,13,14,17,40 of the CO +OH reaction at pressures below 800 torr, are given later in Table III.

By microscopic reversibility the ratio of activating collisions divided by the deactivating collisions in a step is equal to \( [\rho(E)/\rho(E-\Delta E)] e^{-\alpha kBT} \), where \( \rho(E) \) is the density of states of the intermediate at a total energy \( E \). When \( \rho(E) \) is approximated by a constant \( \Delta \) for a range of energies near the dissociation threshold,58 the probabilities can then be written as

\[ P_{E \rightarrow E'} = \begin{cases} \frac{e^{-\alpha kBT}}{1 + e^{-\alpha kBT}} \cdot E' = E + \alpha \\ \frac{e^{-\alpha kBT}}{1 + e^{-\alpha kBT}} \cdot E' = E - \alpha \\ 0, \text { otherwise.} \end{cases} \]  

The Lennard-Jones potential was used for interaction between a bath molecule and HOCO. The \( \epsilon \) and \( \sigma \) parameters for the various bath gases in the Lennard-Jones potential are the same as those calculated by Zhu \textit{et al.}21 The lower limit for the energy range in the present calculations is 3000 cm\(^{-1}\) below the dissociation limit of CO and OH formation at the given temperature and pressure; even energies below \(-2500 \text{ cm}^{-1}\) are sufficiently lower than the dissociation threshold that the probability of the intermediate reacquiring enough energy to dissociate is negligible. To ensure convergence in the high-temperature and high-pressure region, the upper limit of the energy used in the calculations was 12 000 cm\(^{-1}\) above the dissociation limit, and the total angular momentum \( J \) covered the range from 0 to 220 in the \( E, J \)-resolved calculation. The energy grain size used for the energy integration in computing the pressure effect on the reaction rate constant was \( 1 \text{ cm}^{-1} \), 59 (For calculations restricted to 500 K or less, an upper limit of 5000 cm\(^{-1}\) in \( E \) and of 100 in \( J \) would have sufficed.)

E. OH energy partitioning and nonstatistical modification

In the RRKM theory it is assumed that the rate of energy redistribution in the intramolecular states of the intermediate is much faster than that of the dissociation. The molecular-beam results of Simons and co-workers for the reaction H + CO\(_2\) → HOCO\(^{-}\) → OH + CO show, albeit at higher energies than the present, that the CO formed has less energy than would be expected from statistical microcanonical behavior,
i.e., that there was a limited energy exchange between the newly formed OH bond and the other coordinates in H⋯OCO'. The reaction of CO+OH→H+CO2 requires a concentration of energy in the OH bond stretch in the cis intermediate (Fig. 1). Since the exit channel for the CO2 +H→CO+OH reaction is flat (no sharp repulsive drop), there is no major dynamical effect in this exit channel and so the energy distribution of the CO and OH products should reflect that expected from statistical theory,60–62 applied now to HOCO, with relatively minor corrections for dynamics.

Although the formation of HOCO' from OH+CO is a highly exothermic reaction, the OH group would still be present largely in its vibrational ground state in a microcanonical distribution ρ(EJ) at room temperature, because of the high OH-stretching vibrational frequency. However, since the potential-energy barriers in the entrance and exit channels are similar, a large amount of internal energy in an energetic HOCO molecule needs to be transferred into the OH bond in order for the HOCO to dissociate into CO2+H. After this barrier at exit channel is overcome, the H atom then dissociates from the OCO. The harmonic energy steps of OH stretching in HOCO are about 3600 cm−1, which is larger by 1700 cm−1 than that for the other vibrational modes. Because of this large frequency difference between OH stretching and the other modes, the internal energy transfer to OH-stretching motion via internal resonances with the other modes may be less rapid than needed for internal equilibration. (A higher-order resonance is possible.) Prompted by the experimental results of Brouard et al.63–65 and other results, we have explored, with details in Appendix C, the effect of a reduced intramolecular energy transfer between the OH-stretching-mode motion and the other coordinates.

In this first trial calculation, an intramolecular energy-transfer parameter ξ in HOCO is defined in Eq. (C2) for the transfer to and from an OH vibration and is assumed to be 4000 cm−1. It is somewhat larger than the vibrational frequency of the OH stretching in the HOCO', so allowing the intramolecular transfer of roughly one OH vibrational quantum. The nonstatistical modification significantly improves the agreement with the data for the rate constants of CO+OH at high temperatures, the rate constants of CO+OD at low pressures at room temperature, and the k_{OH,CO}/k_{OD,CO} ratio at room temperature. The ε^{13}C values at around 1 atm are also improved. The results are given and discussed in the following sections.

III. RESULTS

A. Temperature effect

The experimental values of the rate constants at 100 and 300 K in the low-pressure limit are about 1.1×10^{-13} and 1.5×10^{-13} cm³ molecule⁻¹ s⁻¹, respectively. The two parameters E₁ and E₂ used to adjust potential energies of TS_a and TS_d to obtain agreement with these two low-pressure rate constants are 0 and 340 cm⁻¹, respectively, using the usual RRKM theory. When the nonstatistical modification is employed, they are instead 10 and 30 cm⁻¹, respectively. (Since the low-pressure rates are independent of the parameter α, we used them for this purpose.) The classical barrier with zero-point energy added for TS_a is about 280 cm⁻¹ in both RRKM theory and the nonstatistical modification. For TS_d it is 965 and 655 cm⁻¹, respectively. Although there are some differences between the barriers of the unmodified and modified RRKM theory, their temperature-dependent rate constants at low pressures shown in Fig. 2(a) are similar. The calculations in Figs. 2(a) and 3 show that the slopes of the calculated Arrhenius plot are comparable to those of the experiment, and the change in the slopes occurs in both cases between 300 and 500 K. However, as seen in Fig. 3, the nonstatistical modification significantly improves...
the agreement between experimental and calculated results of the rate constants for CO+OH in the high-temperature region.

B. Pressure effect

Low and high-pressure rate constants \( k^0 \) and \( k^\infty \) were studied experimentally by Fulle et al. at various temperatures. Their fitted values are summarized in Table II, together with the present calculations and the step-down energy-transfer model. The effect of bath gases on the reaction of CO+OH has been investigated for He, N\(_2\), air, CF\(_4\), and SF\(_6\). The calculated rate constants are compared with the experimental data in Fig. 4(a). The \( \alpha \) values obtained by unmodified and modified RRKM theory are similar and are listed in Table III.

C. OD and OH

Pressure-dependent rate constants were measured for the OD+CO reaction with different bath gases by Paraskevopoulos and Irwin, Golden et al., and Frost et al. The three sets of measurements and the calculations for He, Ar, N\(_2\), air, CF\(_4\), and SF\(_6\) are compared in Fig. 4(b). For simplicity, the \( \alpha \)'s for OD+CO are assumed to be the same as those for OH+CO. The fitted curves for OD+CO using the nonstatistical modification, shown in Fig. 5, agree well with experiment, except for N\(_2\), which is similar to the results of Golden et al. and Senosiain et al. and are discussed later. Unmodified RRKM theory underestimates the rate constant of OD+CO at low pressures at room temperature, and so the fitted RRKM curves in Fig. 4(b) do not agree well in the low-pressure region. However, the nonstatistical modification significantly improves the agreement, as seen in Fig. 5.

The ratio \( k_{\text{OH}}/k_{\text{OD}} \) calculated using RRKM theory is plotted in Fig. 6 as a function of the pressure of He. The theoretical results agree with the experimental data relatively well at pressures higher than 300–500 torr, but the nonstatistical modification significantly improves the rate constant ratios in the low-pressure region and agrees with the experimental data in the entire pressure region, as seen by the dashed line in Fig. 6. The pressure-dependent rate constants for the OD+CO reaction with different bath gases calculated with the nonstatistical model in Fig. 5 also improve the agreement with experimental data in the low-pressure region [cf. RRKM theory used in Fig. 4(b)].

The OH+CO and OD+CO systems in Fig. 2(a) show a marked difference in their dependence on temperature at low pressures and are calculated both for the RRKM theory and the nonstatistical modification. The OD+CO rate is predicted to have a minimum value around 1000/T=4. Only Frost et al. appear to have studied the OD+CO reaction at low pressures. The present calculation is in general agreement with their experimental results.

Jonah et al. measured the temperature dependence of the OH+CO and OD+CO rate constants at pressures \( \sim 760 \) torr at moderate and low temperatures. Fulle et al. did a similar experiment of the OH+CO reaction at the lower-temperature...
The temperature dependence of the calculated rate constants for both OH+CO and OD+CO reactions at 785 torr of Ar in Fig. 2 are similar in both RRKM theory and its nonstatistical modification models and are in good agreement with experiment.

D. Kinetic isotope effect for carbon

The isotopic fractionation of $^{13}$C, $\varepsilon^{13}$C, is defined as

$$\varepsilon^{13}C = \left( \frac{12_k}{13_k} - 1 \right) \times 1000 \text{ per mil},$$

where $12_k$ and $13_k$ are the rate constants for the reaction of $^{12}$CO and $^{13}$CO with OH, respectively. Pressure-dependent $\varepsilon^{13}$C’s from 75 to 840 torr at room temperature were measured by Stevens et al., Röckmann et al., and Smit et al. The data of Stevens et al. and Röckmann et al. show that $\varepsilon^{13}$C depends slightly on bath gases. Our calculations for the pressure dependence of the $\varepsilon^{13}$C values by both RRKM and the nonstatistical models, shown in Fig. 7, agree reasonably well with the experimental data. As the pressure is increased, the calculated $\varepsilon^{13}$C values also increase and depend slightly on bath gases. We discuss these results in Sec. IV.

IV. DISCUSSION

A. The barriers at TS$_a$ and TS$_d$

1. RRKM theory

As noted in Sec. III A two parameters $E_1$ and $E_2$ are needed to adjust the potential energies of TS$_a$ and TS$_d$ to fit the experimental values of the rate constants at 100 and 300 K. The results of the barrier adjustments are consistent with recent previous estimates, the literature values varying from 100 to 380 cm$^{-1}$ for TS$_a$ and from 0 to 1000 cm$^{-1}$ for TS$_d$. The fitted barriers for TS$_d$ in the literature are more scattered than those for TS$_a$ because of the difference in tunneling arising from large differences in imaginary frequencies at TS$_d$. In most theoretical calculations including the present, the exit channel barrier, $\Delta E_d$, including zero-point energy, is higher than that for the entrance channel, $\Delta E_a$, when the same calculational method is applied to both TS$_s$.

### Table III. The $\alpha$ values of several bath gases. The units are in cm$^{-1}$.

<table>
<thead>
<tr>
<th>Bath Gas</th>
<th>G2M</th>
<th>Nonstatistical$^a$</th>
<th>Zhu et al. (Ref. 21)$^b$</th>
<th>Golden et al. (Ref. 22)$^b$</th>
<th>Senosiain et al. (Ref. 27)$^b$</th>
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</thead>
<tbody>
<tr>
<td>He</td>
<td>140</td>
<td>140</td>
<td>150</td>
<td>150$^f$</td>
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<tr>
<td>Ar$^d$</td>
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<td></td>
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<td>190</td>
<td></td>
</tr>
<tr>
<td>N$_2$</td>
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<td>210</td>
<td>250</td>
<td>300$^f$</td>
<td>100$^f$</td>
</tr>
<tr>
<td>CF$_4$</td>
<td>300</td>
<td>310</td>
<td>450</td>
<td>680</td>
<td>160$^f$</td>
</tr>
<tr>
<td>SF$_6$</td>
<td>400</td>
<td>410</td>
<td>1000</td>
<td>800</td>
<td>190$^b$</td>
</tr>
</tbody>
</table>

$^a$The nonstatistical model is used.

$^b$The pressure effect is calculated using the exponential down model.

$^c$180 cm$^{-1}$ for OD+CO in He.

$^d$Fitted to the data from OD+CO reaction.

$^e$480 cm$^{-1}$ for OD+CO in N$_2$.

$^f$140 cm$^{-1}$ for OD+CO in N$_2$.

$^g$150 cm$^{-1}$ for OD+CO in CF$_4$.

$^h$200 cm$^{-1}$ for OD+CO in SF$_6$.
Because of the large width and low barrier of TS.a, tunneling through TS.a at the low-pressure limit is important only at temperatures below 400 K. The calculated tunneling effects at TS.a increase the rate constant by about 10% in the low-pressure limit for a temperature of 400 K and less at higher temperatures. When any tunneling at TS.a is neglected, the reaction rate at room temperature in the low-pressure limit is less by about 20%. This effect is much smaller than that at TS.d, where the neglect of tunneling reduces the rate constant by a factor of 20.

The rate constants calculated with variation of rotational constants along the minimum reaction path near both TSs reduced the rate constants at low pressure by less than 1% at room temperature, as compared with fixing the position of TS.a at the local potential-energy maximum along this reaction coordinate. In the low-pressure limit the rate constant was reduced by 5% and 10% at 1000 and 2500 K, respectively. At high pressures this effect of varying the rotational constants along the reaction coordinate reduced the rate constant by 10% at 1000 K and 20% at 2500 K. The variational effects by Senosiain et al.27 reduced the rate constants by 23% at 1000 K and 42% at 2500 K at low pressures, and by 11% at 1000 K and 21% at 2500 K at high pressures. The decrease is smaller in our calculations, presumably due to their taking into account the changing hindering rotation of OH and CO in TS.a along the reaction path.

2. Nonstatistical modification

The barrier of TS.a in the modification is similar to that used for the usual theory, reflecting the small sensitivity of the rate to TS.a. The barrier of TS.d now needed is about 300 cm⁻¹ less than before. This lowering compensates for the constraint placed on the intramolecular energy transfer in the HOCO⁺ when it passes through the TS.d in this nonstatistical modification.

B. Non-Arrhenius behavior

1. RRKM theory

The strongly non-Arrhenius behavior of the CO+OH reaction in Fig. 2 was observed in many earlier studies, both experimentally5,6,11–13,15 and theoretically17,20–22,25. The experimental rate constants are slightly temperature dependent at low pressures in the temperature range 100–500 K, but increase steeply at higher temperatures.5,6,11 Because the barriers to form products (H+CO₂) or revert to reactants (OH + CO) from HOCO⁺ are similar, the competition between \( k_{-1} \) in Eq. (1) and \( k_2 \) in Eq. (2) for HOCO⁺ disappearance serves to explain the change of slope of ln \( k \) versus \( 1/T \) of the reaction in Fig. 2(a); in the present calculations at high temperatures the rate constants are slightly temperature dependent at low pressures in the temperature range 100–500 K, but increase steeply at higher temperatures.
temperatures the $k_{-1}$ in Eq. (1) is much larger than the $k_2$ in Eq. (2). So the barrier in the exit channel $TSD$ then dominates the rate. However, at temperatures below ~500 K $k_{-1}$ and $k_2$ are comparable. Due to a large tunneling through $TSD$ at lower energies, $k_2/k_{-1}$ ultimately increases as the temperature is decreased and so the contribution to the barrier from the rate at the entrance channel $TSD$ increases. In summary, the change in slope in Fig. 2(a) is due to the similar barrier heights of both entrance and exit channels together with a large tunneling effect in the exit channel $TSD$.

2. Nonstatistical modification

The results calculated using the modification are similar to those obtained with the unmodified theory. However, as seen in Fig. 3, the modification describes the experiments at the high temperatures better than before.

C. Pressure effect

1. RRKM theory

The calculated low- and high-pressure rate constants, $k^0$ and $k^a$, at the various temperatures listed in Table II, were compared with the extrapolated experimental values of Fulle et al. As mentioned earlier, the calculated rate constants in the low-pressure limit are very similar to the extrapolated experimental values. The rate constants at the high-pressure limit are largely controlled by the properties of $TSD$, and the extent of agreement with the extrapolated experimental values by Fulle et al. in the high-pressure limit is seen in Table II. Any difference may reflect a shortcoming of the calculation of $TSD$.

The effect of bath gases on the reaction of CO+OH has been studied for He, N$_2$, air, CF$_4$, and SF$_6$. It has been repeatedly documented that polyatomic bath gases are better energy-transfer agents than monatomic gases. Possible reasons include higher densities of intramolecular energy states for the collisional energy transfer, larger collision radii, and more attractive interactions leading to a closer and so more steeply repulsive interaction. The fitted energy-transfer parameter $\alpha$ for He, Ar, N$_2$, CF$_4$, and SF$_6$ is 140, 190, 200, 300, and 400 cm$^{-1}$, respectively. The calculated rate constants and the experimental data are compared in Fig. 4(a). The $\alpha$ values listed in Table III are close to the fitted $\alpha$ values of Zhu et al. and Senosiain et al., especially for monatomic and diatomic gases, but are larger than the fitted values of Senosiain et al. Strictly speaking the $\alpha$ values are not exactly equivalent since the exponential down model is used by all other groups. In the results of other groups and ours the somewhat larger deviations of the $\alpha$ values for the polyatomic gases may arise from the different pressure models employed.

The pressure-dependent measurements of OD reacting with CO in various bath gases, He, Ar, N$_2$, air, CF$_4$, and SF$_6$ are compared in Fig. 4(b) with calculations using the unmodified RRKM theory. Assuming the $\alpha$’s for OD +CO to be similar to those used for OH+CO, the fitted curves for OD+CO are in general agreement with experiment. The results indicate that $\alpha$ depends not only on the bath gas but also on whether the isotope is D or H. Senosiain et al. optimized their calculations to agree with the experimental data by adjusting the $\alpha$ values for both OD+CO and OH+CO. Their $\alpha$’s for OD+CO are somewhat higher than those for OH+CO, but the difference is small, except for N$_2$, for which the results are similar to ours. For simplicity, the same $\alpha$ value for the C-isotope-substituted OH+CO reactions in each bath gas is assumed in the study of the kinetic C-isotopic effect discussed in Sec. IV F.

2. Nonstatistical modification

The $k^0$ values at high temperatures obtained with the RRKM modified theory are smaller than those obtained with conventional theory and are closer to the extrapolated experimental results. The $k^a$ values are similar in both treatment, because they are mainly determined by the properties of $TSD$, which is treated in the same way in both models. The energy-transfer parameter $\alpha$ for various bath gases by both models are similar and are listed in Table III. The nonstatistical theory improves the agreement of the pressure dependence of OD reacting with CO in various bath gases significantly, especially for the extrapolated values at zero pressure, as shown in Fig. 5. The most important improvement arising from the nonstatistical model is in the plot of $k_{OD+CO}$ versus pressure, especially at lower pressures, as in Fig. 6. In that figure, the large discrepancy between experiment and the conventional RRKM theory is clear. This discrepancy with conventional RRKM theory was also seen in Fig. 9 of Ref. 27 and is now removed by the nonstatistical modification (Fig. 6). The robustness of this result is shown in Appendix A.4 (Fig. 10).

D. Negative temperature effect in OD+CO

1. RRKM theory

The calculated rate constants for OD+CO in Fig. 2(a) display a minimum around 250 K. Only Frost et al. appear to have studied the OD+CO reaction as a function of temperature at low pressures. In their study at 178, 216, and 295 K, Fig. 2(a), they observed nearly temperature-independent reaction rates. Their RRKM calculations also showed a slight decrease in rate in going from 80 to 300 K. There is general agreement between the experimental and calculated results.

The relatively large negative temperature effects observed in the OD+CO reaction, shown in Fig. 2(a), can be explained by the lowered barrier at $TSD$ and the increased height of the $TSD$ barrier in OD+CO relative to that in OH +CO. The energy with the zero-point corrections of $TSD$ in OD+CO is around 150 cm$^{-1}$ and that of $TSD$ is around 1320 cm$^{-1}$. Although the value of the imaginary frequency of OD at $TSD$ is smaller than that of OH, the barrier change increases the tunneling window (energy difference between $TSD$ and $TSD$) by about 500 cm$^{-1}$ in OD+CO. The decreased barrier in the entrance channel allows the formation of more intermediates at low temperatures. At low pressures, the intermediates can only tunnel through $TSD$ to products or revert to reactants. The number of states is approximately independent of temperature, but the partition function of the collision pair decreases as temperatures increase. From our calculations in Fig. 2(a), the negative temperature effect is signifi-
cant at temperatures lower than 250 K. The experiments on OD+CO at low pressures and temperatures would permit a test of the prediction of a negative temperature effect in the present calculation.

When different \textit{ab initio} theories are used there is a difference of imaginary frequency at TS\textsubscript{d} in various force constants. As a result, the temperature where the minimum in the calculated rate constant occurs is spread from 200 to 400 K, as shown in Appendix A 2, the higher temperature occurs with the larger value of the imaginary frequencies. The studies on OD+CO at low pressures and low temperatures would help determine better the value of the imaginary frequency at TS\textsubscript{d} and so determine more precisely the magnitude of tunneling effects.

2. Nonstatistical modification

Comparing the Arrhenius plot of the OD+CO reaction at low pressures for the unmodified and modified theories, the former has a weaker temperature dependence in the low-temperature region. This result is consistent with the results in the experiment of Frost \textit{et al.} There are only two measurements of the rate constants below 250 K. More measurements at low temperatures would permit a test of the trend predicted in Fig. 2(a).

E. OH and OD

1. RRKM theory

In the measurements of Paraskevopoulos and Irwin, Frost \textit{et al.},\textsuperscript{11} and Golden \textit{et al.},\textsuperscript{17} the pressure-dependent kinetic H/D isotope effect is given at about 300 K. At low pressures their experimental values for \( k_{\text{OH}+\text{CO}} / k_{\text{OD}+\text{CO}} \) are 3.6, 2.7, and 2.3, respectively. This kinetic effect can be understood in terms of tunneling, together with the zero-point energy difference of TS\textsubscript{a} and TS\textsubscript{d}. Compared with the ratios in the data of Paraskevopoulos and Irwin, the ratio \( k_{\text{OH}+\text{CO}} / k_{\text{OD}+\text{CO}} \) calculated using the unmodified theory, the solid line in Fig. 6, significantly overestimates it at pressures below 500 torr, but is comparable with experiments at higher pressures. The robustness of this result is seen later in Appendix A.

Jonah \textit{et al.} measured the rates of OH+CO and OD+CO from 340 to 1250 K at 1 atm of Ar and 15 torr of water vapor.\textsuperscript{2} Both rates have minima around 600 K [Fig. 2(b)]. This negative dependence on temperature was also observed by Fulle \textit{et al.}\textsuperscript{3} in the rates of OH+CO from 91 to 288 K at 750 torr of He. The rate constants measured by the latter group showed a maximum around 170 K. Our calculations in Fig. 2(b) for both OH+CO and OD+CO reactions at 785 torr of Ar are close to their results. The minimum rate of OH and OD occurs at about 400 and 500 K, respectively, which are slightly lower than the measured values. The calculated negative temperature dependence of the CO+OH reaction rate between 200 and 400 K has a maximum around 160 K, which is close to the maximum in the measurements of Fulle \textit{et al.}

2. Nonstatistical modification

When the intramolecular energy-transfer restriction is introduced using the modified theory, the agreement of the rate constant ratios at the low-pressure region is significantly improved, as seen by the dashed line in Fig. 6. The modified theory also improves agreement with experiment for the pressure dependence of the low-pressure rate constants of OD+CO as in Fig. 5. The temperature dependence of the rate constants at pressures \( \sim 1 \text{ atm} \) are similar to those in the unmodified theory and in the experiment. At temperatures higher than 1000 K, the rate constants in Fig. 3 obtained using the modified theory agree better with the experiment than before and are lower than the conventional RRKM result by about 60%.

The calculations for the unmodified theory tend to underestimate the rate constants of OD+CO in the low-pressure region, perhaps because any difference in the energy-transfer ability between HOCO\textsuperscript{−} and DOCO\textsuperscript{−} is not considered. The vibrational frequency of the OD stretching in DOCO, about 2700 cm\textsuperscript{−1}, is much less than the OH-stretching frequency. Accordingly, the energy transfer between the OD stretching and other modes in energetic DOCO\textsuperscript{−} may well be easier than in HOCO\textsuperscript{−}. In the nonstatistical modification, the relation between the energy-transfer parameter \( \xi \) in HOCO and that in DOCO is simply assumed to be

\[
\xi_D = \frac{\nu}{\nu_D} \xi, \tag{10}
\]

where \( \nu \) is the OH (OD) stretching frequency in the intermediate, the subscript \( D \) and no subscript indicating the DOCO in HOCO systems, respectively. Equation (10) implies that the energy redistribution in the energetic DOCO is faster than that in HOCO. In counting the number of states for TS\textsubscript{D} with a tunneling correction using Eq. (C2), the range of integration used for DOCO is about 30% larger than that for HOCO due to the \( \nu_D \) in Eq. (10) being smaller than \( \nu \).

F. Kinetic isotope effect for carbon

1. RRKM theory

The reaction at atmospheric pressure was found to favor a positive \( \delta^{13}\text{C} \) by 6 per mil, while negative values were observed at pressures below \~300 torr. These experimental data also show that \( \delta^{13}\text{C} \) depends slightly on the bath gas.\textsuperscript{29,31} The values in monatomic gases, such as He or Ar, are lower than those in diatomic gases, such as air, O\textsubscript{2}, and N\textsubscript{2}. Our calculations in Fig. 7 for \( \delta^{13}\text{C} \) agree reasonably well with these data. The calculated \( \delta^{13}\text{C} \) at low pressures is negative, around \~4 per mil. As the pressure increases, the calculated \( \delta^{13}\text{C} \) increases to positive values, similar to the pressure dependence observed in the experiment. Although the values of collision stabilization energy, the \( \alpha\text{'s} \) for He and N\textsubscript{2} are different, the trend in pressure dependence of \( \delta^{13}\text{C} \) in Fig. 7 is similar. As pressure increases, the \( \delta^{13}\text{C} \) in N\textsubscript{2} increases faster than in He, but the difference is small.

The calculated negative \( \delta^{13}\text{C} \) at low pressures can be understood in terms of the higher density of the vibrational and rotational states for heavier isotopic molecules. The
pressure dependence of $\varepsilon^{13}$C can be explained in terms of the difference of vibrational energies at the TSs and of the collision frequencies, $\omega_i$, for the $^{12}$C and $^{13}$C intermediates; due to the lower barrier and higher number of states in both TS$_{a}$ and TS$_{d}$ when $^{13}$C is substituted for $^{12}$C, $^{13}k$ is larger than $^{12}k$ in the low-pressure region, causing $\varepsilon^{13}$C to be negative in the present theory. The collision frequencies are proportional to $\mu^{-1/2}$, where $\mu$ is the reduced mass of the bath gas and HOCO$^*$. As the pressure is increased the rate of collision stabilization of intermediates is larger due to the larger collision frequency. Therefore, the presently calculated $\varepsilon^{13}$C values in Fig. 7 vary from slightly negative at low pressures to about +5 per mil at 1000 torr, which is similar to the experimental data. The collision factor can introduce a maximum of +7 per mil when the bath gas is N$_2$ and the rate is proportional to the collision frequency.

2. Nonstatistical modification

Our calculations in Fig. 7 for $\varepsilon^{13}$C in both the unmodified and the modified theory agree reasonably well with the experimental data. The $\varepsilon^{13}$C calculated using the modified theory are a little larger than the values obtained before. It increases from −4 per mil to about 0 per mil at 100 torr, which is higher than the independent experimental results of Stevens et al.$^{29}$ and Smit et al.$^{30}$ given in Fig. 7. Even though the values obtained using the model in the low-pressure region tend to overestimate the carbon fractionation, as compared with experiment, they are still in good overall agreement with the data, especially for pressures higher than 500 torr. Similar to the results obtained for the pressure dependence of the $\varepsilon^{13}$C values of two bath gases with the unmodified theory, the values in N$_2$ using the modified theory increase faster than those in He, again in agreement with experiments.

V. CONCLUSIONS

The use of a nonstatistical modification of the RRKM theory significantly improves the agreement with experiment for (1) the pressure-dependent $k_{OD+CO}/k_{OD+CO}$ ratio, (2) the pressure dependence of the OD+CO reaction rate, and (3) the non-Arrhenius behavior at high temperatures. In (1) a major discrepancy between experiment and theory has been removed, assuming the validity of the nonstatistical effect. A negative temperature effect at low temperatures in the OD+CO reaction is predicted in both unmodified and modified RRKM theories. The measured $\varepsilon^{13}$C effect is very small and pressure dependent. The calculations with unmodified and modified RRKM theories are in reasonable agreement with $^{13}$C rate, considering the extremely small magnitude of the effect (from −4 per mil at low pressures to 6 per mil at −1 atm).

ACKNOWLEDGMENTS

It is a pleasure to acknowledge the support of this research by the National Science Foundation. We also thank Dr. Muckerman for providing the F matrix for the structures in their paper,$^{18}$ which were used to check the accuracy of rate constant calculation with our previous vibrational frequencies.

APPENDIX A: RESULTS FOR DIFFERENT PES’S

Calculations with different and less accurate force constant methods were used to test the sensitivity of certain aspects of the calculations and are discussed in this Appendix. The results for the $k_{OD+CO}/k_{OD+CO}$ ratios between 200 and 800 torr and for the non-Arrhenius effect and the pressure effect below 1000 torr all show a robustness toward the different calculations. Apart from the transition state structure of the entrance channel (TS$_{a}$),$^{67}$ the vibrational frequencies and rotational constants of all structures for the various isotopes were calculated by three methods in this comparison,
TABLE V. \(k^0\)'s and \(k^\infty\)'s at temperatures obtained from values extrapolated to \(p=0\) by Fulle et al. (Ref. 5) and calculated with various PESs. The rate constants have units of \(10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}\).

<table>
<thead>
<tr>
<th>PES (ab) initio</th>
<th>G2M</th>
<th>LTSH</th>
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<tbody>
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<td></td>
<td>DFT-CC</td>
<td>DFT-MP</td>
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<tr>
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<td>1.1</td>
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<table>
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<tr>
<td></td>
<td>63.7</td>
<td>37.7</td>
</tr>
</tbody>
</table>

\(^a\)Extrapolated results to \(p=0\) and \(p=\infty\) by Fulle et al. (Ref. 5).
\(^b\)Measured at 98 K.
\(^c\)measured at 190 K.
\(^d\)Measured at 717 K.
\(^e\)Measured at 819 K.

CCSD(T) with 6-31G(\(d,p\)), B3LYP with cc-pVTZ, and MP2 with 6-31+ +G(\(d,p\)). The properties of TS\(_a\) are obtained only from the results of MP2 or CC. The acronym DFT will denote the calculation by B3LYP. DFT-MP will denote the calculations combining the structure data of TS\(_a\) from MP2 and the other statures from B3LYP, while DFT-CC is similar but with the data for TS\(_a\) obtained from CCSD(T). The LTSH potential, given by Lakin et al.,\(^{24}\) involves an analytical function based on recent high-level \(ab\) initio methods and was also used to explore the robustness.

1. Temperature and pressure effects

Some robustness is expected to occur partly because there are two adjustable parameters \(E_1\) and \(E_2\) of the potential energy of TS\(_a\) and TS\(_d\) determined from the two rate constants. These \(E_1\) and \(E_2\) adjustments are listed for each specific PES in Table IV. Although the parameters differ between G2M and LTSH, the resulting classical barriers (i.e., no tunneling) with zero-point corrections of TS\(_a\) and TS\(_d\) are similar at the same \(ab\) initio level. All the barriers of TS\(_a\) with zero-point corrections in our fitted results agree well with the recent estimates derived from kinetic measurements and from barrier theoretical calculations.\(^{17-19,21,24}\) Due to the difference of the imaginary frequencies of TS\(_d\), the \(E_2\) parameter differs most among \(ab\) initio methods. All of them are, as expected, within or near the range of earlier theoretical studies.\(^{17-19,21,24}\) The calculated temperature dependence of rate constants has a non-Arrhenius effect at low temperatures and is quite robust in the present calculations.

The extrapolated low- and high-pressure rate constants \(k^0\) and \(k^\infty\) from the data of Fulle et al.\(^5\) and the calculations

TABLE VI. \(a\) values of several bath gases with various PESs. The units are cm\(^{-1}\).

<table>
<thead>
<tr>
<th>PES (ab) initio</th>
<th>G2M</th>
<th>LTSH</th>
</tr>
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<td>150</td>
</tr>
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<td>SF(_6)</td>
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\(^a\)180 cm\(^{-1}\) for OD+CO in He.
\(^b\)Fitted to the data from the OD+CO reaction.
\(^c\)480 cm\(^{-1}\) for OD+CO in N\(_2\).
The last observation is similar to the results of Senosiain et al. The high-pressure limit of the rate constants is largely controlled by the properties of TS\(_a\). Since there are two kinds of \textit{ab initio} calculation levels, CC and MP2, for TS\(_a\), the high-pressure rate constants \(k^\alpha\)'s in the calculations can be divided into two groups. Both sets of \(k^\alpha\) are close to the \(k^\alpha\) values independently fitted by Zhu et al.\(^{21}\) and Senosiain et al.\(^{22}\) (A caveat in comparing the step ladder \(k^\alpha\) with the exponential down \(\alpha\) was noted earlier.) The pressure-dependent rates in various bath gases calculated by various \textit{ab initio} methods are similar to those in Fig. 4(a). Using the same force constants, the fitted \(k^\alpha\) values from the G2M potential are slightly larger than those from LTSH, since the intermediate in the latter is stabler by about 1.5 kcal mol\(^{-1}\). The last observation is similar to the results of Senosiain et al.\(^{22}\) Their fitted \(k^\alpha\) value for He was reduced from 150 to 100 cm\(^{-1}\) when they lowered the energy of the HOCO\(^*\) intermediates by 5.5 kcal mol\(^{-1}\). Thus, the pressure dependence of the rate constants is controlled by the properties of both TS\(_a\) and the stability of the HOCO\(^*\) intermediate.

FIG. 8. Arrhenius plots for the reaction of CO+OH (black lines) and CO +OD (gray lines) in Ar at 785 torr. All curves in the plot are obtained with G2M, and the LTSH potentials show similar curves. The black and gray open circles are the experimental data of OH and OD reacting with CO, respectively (Ref. 4). The bath gas in the experiment consists of 760 torr of Ar and 15 torr of H\(_2\)O. The open triangles are Fulle's experiments at 760 torr of He. The solid, dot-dashed, dashed, and dotted lines are calculated with DFT-CC, CC, DFT-MP, and MP2, respectively.

FIG. 9. The \(13C\) values for the system of OH+CO as a function of total pressure at room temperature. Both G2M and LTSH potentials give similar results, only the calculations with G2M are shown in the plot. The gray and black lines are for He and N\(_2\) gases, respectively. The solid, dot-dashed, dashed, and dotted lines are calculated with DFT-CC, CC, DFT-MP, and MP2, respectively. The open circles are obtained for N\(_2\) by Smit et al. (Ref. 30). The gray solid and black open squares are obtained by Röckmann et al. (Ref. 31) for He, and N\(_2\) (or N\(_2\)+O\(_2\)), respectively. Others are obtained by Stevens et al. (Ref. 29), where the gray and black indicate that the bath gases are He or Ar, and N\(_2\), O\(_2\), or air, respectively.

3. Kinetic isotope effect for carbon

Since the fractionation of \(13C\) is very small, the different sets of force constants yield slightly different results. However, all the calculated results show similar trends of the pressure dependence of the \(13C\) values between different bath gases, in agreement with the experimental data. The calculated pressure dependence of fractionation of \(13C\) is shown in Fig. 9. All calculations except MP2 give similar agreement.

The negative temperature effects in the OD+CO reaction in the low-pressure limit are obtained for all PESs. The temperature for the minimum rate strongly depends on the imaginary frequency of TS\(_d\). The temperature with the minimum rate is around 400 K in MP2, 300 K in CC, and 200 K in DFT-CC and DFT-MP, and is higher the higher the imaginary frequency. All Arrhenius plots of the OH and OD reacting with CO in Ar at 785 torr, shown in Fig. 8, are similar.

The calculated \(k_{\text{OH+CO}}/k_{\text{OD+CO}}\) ratio decreases to about 2 at 200 torr and about 1.2 at 700 torr using the various force constants for the various PESs. DFT-CC, DFT-MP, and CC overestimate the ratio at pressures lower than 100 torr.

2. OH and OD

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The parameters methods together with the G2M potential. The value of the nonstatistical modification, namely, the CC and DFT-CC two most accurate methods were used to test the accuracy of +CO relative to OD+CO. In the nonstatistical effect only the substituted HOCO, it decreases the ratio of the rate for OH HOCO in both CC and DFT-CC is assumed to be 4000 cm−1.

The solid and dot-dashed lines are calculated with DFT-CC and CC, respectively. The modified and unmodified RRKM theories are used in the calculation. The solid and dot-dashed lines are calculated with DFT-CC and CC, respectively.

The reason for the inadequate pressure dependence calculated with MP2 is probably due to an overestimation of the imaginary frequency in TSd.

4. Nonstatistical effect

Since Eq. (C4) is used to approximate ξ in the isotope-substituted HOOCO, it decreases the ratio of the rate for OH +CO relative to OD+CO. In the nonstatistical effect only the two most accurate methods were used to test the accuracy of the nonstatistical modification, namely, the CC and DFT-CC methods together with the G2M potential. The value of ξ in HOCO in both CC and DFT-CC is assumed to be 4000 cm−1. The parameters E1 and E2 adjusted to fit the rates of OH +CO at 100 and 300 K at low pressures and the energy-transfer parameter α in various bath gases are listed in Table VII. The α values for the model are also similar to those for the conventional RRKM model, hence the pressure dependence of rate constants in both models are similar. However, for both CC and DFT-CC methods the nonstatistical model describes better the rate constant ratios between OH and OD reacting with CO, as seen in Fig. 10. The negative temperature effect of the OD+CO reaction is also observed in the model, shown in Fig. 11.

The kinetic isotope effects of carbon of DFT-CC are in better agreement with experiment at pressures under 400 torr when this nonstatistical correction is used, as seen in Fig. 12. Although the values obtained with CC agree better at pressures higher than 500 torr, both DFT-CC and CC give a similar trend for the pressure dependence of e 13C.

APPENDIX B: THE TUNNELING PROBABILITY FOR AN ECKART BARRIER

For the Eckart potential, frequently used to estimate the tunneling probability of the reaction, the transmission probability as a function of energy E is

![Equation](B1)

where

![Equation](B2)

V1 is the barrier height relative to the beginning of the barrier and V2 is the barrier height relative to the products. The
The CO+OH reaction

Profiles near TS$_g$

FIG. 13. Comparison of different barrier profiles with Eckart potential functions and with IRC calculations. The open diamonds and open squares denote the IRC calculations at TS$_g$ calculated with B3LYP and MP2, respectively. The solid and dotted lines were obtained using the Eckart potential for the exit channel using B3LYP and MP2, respectively.

potential is written for an exoergic reaction, i.e., $V_1 \equiv V_2$, and $\nu$ is the absolute value of the imaginary frequency.

The GAUSSIAN 98 program supports the intrinsic reaction coordinate (IRC) calculation for the MP2 and DFT methods, but not for the CC method. Since the value of the imaginary frequency of TS$_g$ by CC is intermediate between those obtained using the other methods, the difference between the barrier profiles estimated by the Eckart potential functions and calculated by the IRC method with DFT or MP2, shown in Fig. 13, would be expected to reflect the nature of any error in using the Eckart potential to approximate calculations by CC. The Eckart potential is sharper than that obtained by the IRC calculation, especially when the imaginary frequency is large. (The imaginary frequency is 1526 cm$^{-1}$ in DFT and 3372 cm$^{-1}$ in MP2.) However, most of the tunneling region occurs between the top of the barrier at TS$_g$ and about 1500 cm$^{-1}$ below it. In comparing the tunneling probabilities through Eckart and parabolic barriers, the shape of the bottom of the barrier has a negligible effect on the reaction rate constants. We infer that an Eckart potential is a reasonable barrier to approximate the tunneling effect in the current study.

APPENDIX C: EQUATION FOR THE NONSTATISTICAL MODEL

In a trial intramolecular energy-transfer model, we assume that only a limited amount of energy, $\xi$, can be transferred into or out of OH stretching just before the HOCO* passes through TS$_g$. The dissociation rate of HOCO* at a total energy $E$, rotational quantum state $J$, and quantum number $n$ of OH stretching in HOCO* is then

$$k_n(EJ) = \frac{N_n(EJ)}{h\rho(EJ)}, \quad (C1)$$

where

$$N_n(EJ) = \int_{\text{Min}[E,E_0+n\hbar\nu-\xi]}^{\text{Max}[E_0,E_0+n\hbar\nu+\xi]} \kappa_n(E')\rho_n(E')dE'. \quad (C2)$$

The definitions of $\kappa_n(E')$ and $\rho_n(E')$ are the same as in Eq. (7). $\nu$ is the OH-stretching frequency of cis-HOCO, and the lower and upper bounds of the integral correspond respectively to the maxima value between $E_0$ and $E_0+n\hbar\nu-\xi$ and the minima value between $E_0$ and $E_0+n\hbar\nu+\xi$. The dissociation rate of HOCO* at total energy $E$ and rotational state $J$ can be expressed from Eq. (C1) as

$$k(EJ) = \frac{1}{n_{\text{max}} \sum \rho_n(EJ)} \rho(EJ) k_n(EJ), \quad (C3)$$

where $n_{\text{max}}$ is the maximum allowed quantum number of the OH stretching in the energetic intermediate at $(E,J)$. $\rho_n(EJ)$ is the density of states of the intermediate at OH vibrational state $n$, and $\rho(EJ)$ is the total density of states at total energy $E$ and rotational state $J$.

The energy-transfer parameter $\xi$ depends on the difference in resonances of the OH-stretching vibration with other modes and on the anharmonic coupling of the PES. It can also influence the isotope effects. As a first approximation, and similarly as in Eq. (10) in the deuterium case, $\xi_{iso}$ is written as

$$\xi_{iso} = \frac{\nu}{\nu_{iso}}, \quad (C4)$$

where $\nu$ is the OH-stretching frequency in the intermediate and the subscript “iso” and no subscript indicates the isotope-substituted HOCO and normal isotopologue of HOCO, respectively. Although a very crude approximation, the equation includes the fact that the internal energy transfer would be more rapid when the stretching frequency is closer to that of the other modes. For example, the $\xi$ value in DOCO calculated from the approximation is about 40% larger than that in HOCO.

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1. With the same values of $\alpha$ for various bath gases, the difference of calculated rates by using the rotational constants and vibrational frequencies between CC and Yu et al. (Ref. 18) is less than 1% as pressure is under 1000 torr. The values of carbon-isotope fractionation by CC is higher than the other by around 1 per mil at low pressure and around 2.5 per mil at 1 atm. Thus, the structures and vibrational frequencies by CC is good enough for our study.

2. Recent high-level ab initio calculations are considered in the energies of stationary points in the LTSH potential. Since the imaginary frequency of TS$_2$ in the LTSH potential is too low, the vibrational frequencies and rotational constants of all stationary structures are calculated by ab initio methods. Only the energies without ZPE corrections of the stationary structures are obtained from the LTSH potential.


4. The tunneling corrections for microcanonical systems were described by Marcus (Ref. 50). Instead of the usual sum over states $N(E,J)\Sigma n\epsilon(\epsilon)$, one sums over the TS-dependent tunneling over probabilities $\Sigma \epsilon(n|\epsilon)\epsilon$ for $\epsilon < E$. A similar prescription is given by Miller (Ref. 51). The details about evaluating the tunneling probability are described in Sec. II C.

5. Including the loose transition structure forming OH$\cdots$CO, the rates are decreased by about 5% at room temperature. It has negligible effects on the rate constant ratios for OH and OD reacting with CO at pressures below 1000 torr. The difference in fractionation of the carbon isotope calculated with and without the additional loose transition structure in the calculation for the present pressures 0–1000 torr and room-temperature regions is about 0.5 per mil.


9. The rotational barrier was obtained from the high-level ab initio calculations by Yu et al. (Ref. 18).

10. The sharp barrier at TS$_1$ is not included in any analytical potential functions by Schatz et al. (Refs. 25 and 46).


15. As the exponential model is used to test the convergence, the results of grain size in 5 and 10 cm$^{-1}$ are the same.


22. The experimental rate constants at low pressure near 300 K are in the range of $(1.3-1.8)\times10^{-13}$ cm$^3$ molecule$^{-1}$ s$^{-1}$. The rate constant is chosen by a best fit to the pressure dependence at 300 K.

23. The HO$\cdots$CO distance of TS$_2$, optimized by B3LYP, is too long, which is around 2.99 Å in B3LYP. So doubtful frequencies would be expected. A similar result happened when Lin used B3LYP/6-311G(d,p) to calculate TS$_2$. (Ref. 21).