CRDS Study of Tropospheric Ozone Production Kinetics: Isoprene Oxidation by Hydroxyl Radical

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ABSTRACT

The tropospheric ozone production mechanism for the gas phase additive oxidation reaction of hydroxyl radical (OH) with isoprene (2-methyl-1,3-butadiene) has been studied using cavity ring-down spectroscopy (CRDS) at total pressure of 50 Torr and 298 K. The applicability of CRDS was confirmed by monitoring the shorter (~4%) ring-down time in the presence of hydroxyl radical than the ring-down time without the photolysis of hydrogen peroxide. The reaction rate constant, $(9.8 \pm 0.1) \times 10^{-11}$ molecule$^{-1}$ cm$^{3}$ s$^{-1}$, for the addition of OH to isoprene is in good agreement with previous studies. In the presence of O$_2$ and NO, hydroxyl radical cycling has been monitored and the simulation using the recommended elementary reaction rate constants as the basis to OH cycling curve gives reasonable fit to the data.

Keywords: cavity ring-down spectroscopy, ozone, isoprene, oxidation, rate constant

I. Introduction

Among the issues in regional air quality, the tropospheric ozone production through the photochemical oxidation of unsaturated hydrocarbons has been drawing attention of many atmospheric scientists. Isoprene (2-methyl-1,3-butadiene, C$_5$H$_8$) is the most abundant non-methane biogenic organic compound emitted by vegetation into the atmosphere (~450 Tg/yr) and its oxidation serves as a major source of tropospheric ozone.\(^1,2\) During the daytime, isoprene reacts predominantly with the hydroxyl radical (OH) which has tropospheric lifetime of a few seconds or less and the concentration of OH in the sunlit troposphere is near $10^6$ molecules/cm$^3$.\(^3\)

In the metropolitan city, the atmosphere is polluted by automobile exhaust gases including nitrogen oxides (NO$_x$)\(^4\) and isoprene is oxidized by OH in the presence of moderate concentration of NO as well as O$_2$. The OH regeneration during isoprene oxidation in the presence of NO and O$_2$ is abbreviated shown in Fig. 1. The first intermediate radical product, hydroxy alkyl radical (HOCC$_5$H$_8$), reacts with O$_2$ to produce the hydroxy peroxy radical (HOCC$_5$H$_8$O). The reaction between hydroxy peroxy radicals and NO yields alkoxy radical (HOCC$_5$H$_8$O) and nitrogen dioxide. The decomposition of alkoxy radicals, followed by the HO$_2$ + NO reaction regenerates OH radical. It is worth noting that each cycle of the isoprene oxidation produces two molecules of ozone from the reaction between O$_2$ and O followed by photolysis of nitrogen dioxide.

Fig. 1. Schematic diagram showing the OH initiated isoprene oxidation cycle which results in ozone production.

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Furthermore, the following oxidation processes of isoprene oxidation products such as methyl vinyl ketone, methacrolein, acrolein, and 3-methylfuran produce more ozone in the troposphere. It is also crucial to determine the yield distribution of these first oxidation products since the impacts on the elevation of the tropospheric ozone concentration by each of them are different. In this regard, careful tracing of isoprene oxidation process through theoretical or experimental kinetic studies is important for understanding detail isoprene oxidation mechanism which reveals the end product distribution. From the kinetic study, the rates of elementary reactions are determined and the information on the dominant reaction path can be provided by comparing the reaction rates when there are several possible reactions competing.

Initial reaction of isoprene and OH can result in four possible adducts and the initial branching ratios among these hydroxy alkyl radicals strongly influence the final product distribution. There have been several theoretical and end product analysis studies suggesting a preference for OH addition to the terminal carbons although a recent mass spectrometry study indicates comparable yields of all four radicals. Despite the discrepancy in the initial branching ratios, there is general consensus regarding the rate constant of \( (1.0 \pm 0.1) \times 10^{-13} \text{ molecule}^{-1} \text{ cm}^3 \text{ s}^{-1} \) at 298 K.\(^1\)\(^{2-5}\)

In the presence of \( \text{O}_2 \), the hydroxyl alkyl radicals react with \( \text{O}_2 \) forming peroxy radicals (RO\(_2\)). The reported overall reaction rate constants\(^6\)\(^{17}\) are in the range from \( 1 \times 10^{-13} \) to \( 8 \times 10^{-14} \text{ molecule}^{-1} \text{ cm}^3 \text{ s}^{-1} \) and there is little of direct measurement studies on the formation rate of RO\(_2\) due to the difficulty in monitoring RO\(_2\). A direct measurement study by Zhang et al.\(^18\) using fast flow reactor coupled to chemical ionization mass spectrometry detected the rate constant of \( (7 \pm 3) \times 10^{-13} \text{ molecule}^{-1} \text{ cm}^3 \text{ s}^{-1} \) and it is more than an order of magnitude larger than the previous report of \( (2.8 \pm 0.7) \times 10^{-15} \text{ molecule}^{-1} \text{ cm}^3 \text{ s}^{-1} \) obtained by indirect OH cycling experiments. In this regard, UV absorption spectroscopy shall be considered as an efficient method to determine RO\(_2\) formation rate using OH cycling experiments. There have been difficulties in monitoring the fluorescence of OH due to the high fluorescence quenching efficiency of \( \text{O}_2 \) when laser photolysis/laser-induced fluorescence (LP/LIF) spectroscopy is used with high concentration of \( \text{O}_2 \). High concentration of \( \text{O}_2 \) provides more dramatic OH signal rise (cycling) at earlier time of the isoprene + OH + \( \text{O}_2 \) reaction and this gives more detail information of \( \text{O}_2 \) addition to hydroxy alkyl radical. On the contrary, the absorption method is free from the quenching problem and the temporal profile of OH can be efficiently detected. Recently, various studies\(^20\)\(^{23}\) using cavity ring-down spectroscopic (CRDS) detection method which basically uses absorption property of the object species have been adopted due to its simplicity: absorption is derived from the increased decay rate of laser radiation within a stable optical cavity induced by absorbing species. Especially, CRDS is an excellent detection method for measuring kinetics of radical species and the sensitivity of CRDS is very high. Lin et al.\(^24\)\(^{25}\) have used CRDS for kinetics measurement for phenyl radical and Atkinson and Hudgens\(^26\)\(^{28}\) have adopted CRDS for ethyl radicals, propargyl radicals and chloroally radicals.

In this study, a CRDS apparatus has been tested for measuring the rate constant of OH + isoprene addition reaction and it has been found the rate constant is in good agreement with previously reported values. OH cycling experiment also has been performed in the presence of \( \text{O}_2 \) and NO at the total pressure of 50 Torr and 295 K and early OH cycling has been monitored with \( \text{O}_2 \) partial pressure of 6 Torr.

### II. Experimental Section

The experimental determination of the rate constants was performed using a slow flow reactor with a length of 60 cm and two highly reflective mirrors (reflectivity > 99.5%) which have a 0.56 m radius of curvature were installed at both ends of the reaction cavity. The cavity was evacuated by a mechanical pump and argon was used as a
buffer gas. Throughout the experiment, the total pressure and temperature were maintained at 50 Torr and 295 K, respectively. 90–95% hydrogen peroxide (H$_2$O$_2$) was prepared by vacuum distillation by checking the density of the solution and H$_2$O$_2$ was transferred into the cavity by flowing argon through a bubbler containing concentrated H$_2$O$_2$. Isoprene buffered with argon gas was prepared in a 5L bulb via freeze-pump-thaw method and introduced into the cavity using a mass flow controller to yield typical concentrations of 1×10$^{14}$ molecules cm$^{-3}$. The NO was buffered with argon in a 5L bulb to known concentrations and was introduced into the cavity through a flow controller. The concentrations of NO were varied from 0 to 1.6×10$^{15}$ molecules cm$^{-3}$ and O$_2$ concentration was fixed at 6 Torr (1.96×10$^{17}$ molecules cm$^{-3}$).

Fig. 2 is a schematic diagram of cavity ring-down apparatus. The frequency doubled probe beam was generated by the Nd:YAG pumped dye laser with a pulse duration of 6–8 ns and OH from H$_2$O$_2$ photolysis was produced by the 248 nm beam from the excimer laser. The delay between the pump (excimer laser) and probe (dye laser) beam was adjusted by a delay generator. The probe beam leaking out of the exit mirror has been pointed to a photomultiplier tube which transferred analog signal proportional to the photon density to a digital oscilloscope. A computer was interfaced to the oscilloscope to analyze the ring down time. The temporal photon decay can be described by the equation

$$I(t) = I_0 e^{-\frac{(1-R)\alpha}{cL}t}$$  \hspace{1cm} (1)

where $I$ is the number of photons detected by the photomultiplier tube, $I_0$ is the number of photons injected into the cavity, $L'$ is the length of the absorbing medium which is the same as the length of cavity ($L$) in this study, $c$ is the speed of light, and $R$ and $\alpha$ are mirror reflectivity and the absorption coefficient, respectively. When the averaged waveform is transformed to a log[intensity] vs. time profile, it was fit to a line using a least squares algorithm. The resulting slope of the line is the inverse of the time constant for the single exponential decay of light intensity from the cavity and it can be written as

$$\frac{1}{t} = \frac{L' \alpha c + (1-R)c}{L}$$  \hspace{1cm} (2)

where intrinsic ring-down time (without absorbing species) is given as

$$\frac{1}{t_0} = \frac{(1-R)c}{L}$$  \hspace{1cm} (3)

Fig. 3 shows the temporal the ring-down profiles obtained in the absence (probe beam only) and the presence (pump and probe beam) of OH and the intrinsic ring-down time ($t_0$) of 546 ns which corresponds mirror reflectivity of 99.6% with a cavity length of 60 cm was measured whereas the shorter (~4%) ring-down time of 527 ns was
monitored due to the absorption by OH.

### III. Results and Discussion

The absorption coefficient \( (\alpha) \) is the product of the extinction coefficient \( (\sigma) \) and the concentration of the hydroxyl radical, \([OH]\), at a time, \( t \), after the firing of the pump or photo-dissociation beam from the excimer laser and an initial concentration of hydroxyl radical, \([OH]_0\), decays exponentially in the pseudo-first-order reaction condition.

\[
\alpha = \sigma \cdot [OH] = \sigma \cdot [OH]_0 \cdot e^{-k't}
\]

where \( k' \) is the pseudo-first-order reaction rate constant.

From the above equations, a relation can be expressed as

\[
\frac{1}{t_c} - \frac{1}{t_0} = \frac{\sigma [OH]_0 cL}{L} \cdot e^{-k't}
\]

or

\[
\ln \left( \frac{1}{t_c} - \frac{1}{t_0} \right) = \ln \left( \frac{\sigma [OH]_0 cL}{L} \right) - k't
\]

Fig. 4 shows the plot of Eq. (6) as a function of the delay time between pump and probe beam in the presence of excess isoprene compared with OH which satisfies the pseudo-first-order reaction condition. As the isoprene concentration increases, the slope gets steeper and the trend is similar to the previous results obtained by the laser photolysis / laser-induced fluorescence (LP/LIF) method.

The absolute value of bimolecular reaction rate constant, \( k \), was obtained from the slopes of the linear plots of \( k' \) versus isoprene concentration and it is depicted in Fig. 5. The pressure dependence of the isoprene + OH reaction rate has been reported by a number of groups and it is generally accepted that the reaction is in high pressure limit regime even at 10 Torr. Hence, it is assumed that the reaction condition with the pressure of 50 Torr is in high pressure limit regime.

The determined rate constant of \((9.8 \pm 0.1) \times 10^{-11}\) molecule\(^{-1}\) cm\(^3\) s\(^{-1}\) is in good agreement with the previous recommendation of \(1.0 \times 10^{-10}\) molecule\(^{-1}\) cm\(^3\) s\(^{-1}\) by Atkinson et al.

As shown in Fig. 1, OH regeneration process happens in the presence of NO and O\(_2\), and Fig. 6 shows a OH cycling data using concentration of isoprene and O\(_2\) of \(5.6 \times 10^{15}\) molecules cm\(^{-3}\) and \(2.0 \times 10^{17}\) molecules cm\(^{-3}\), respectively. A numerical program, KINTECUS, was used to simulate the data from the experiments and the previously reported 14 reactions\(^{27}\) and corresponding reaction rate constants given in Table 1 were used in the simulation. The fits of the simulation to the data are shown as solid lines in Fig. 6. The upper curve was obtained with NO concentration of \(2.0 \times 10^{15}\) molecules cm\(^{-3}\) and exhibits the typical exponential decay at early times until the onset of recovery signal at around 50 µs.\(^{27,30}\) Although there is
Concentration effect of OOH improved to obtain a set of data showing the apparent signal regeneration due to the cycling of OH, relatively large signal to noise needs to be improved to obtain a set of data showing the concentration effect of O2 and NO on the recovery time. In the absence of NO (lower line), the observed decay shows pseudo-first order kinetics with a rate of 1.0×10−10 molecule−1 cm3 s−1, consistent with the recommendation value.20

IV. Conclusion

This work focuses on the verification of CRDS measurements for reaction kinetics of isoprene oxidation which results in the elevation of ozone concentration in the troposphere. The measured initial rate constant for the isoprene+OH reaction is in excellent agreement with previously reported value and OH cycling was successfully monitored. Although a direct monitoring of hydroxy peroxy radicals was not available in this study due to the lack of adequate highly reflective mirrors ranging 210–260 nm where the UV absorptions of typical peroxy radicals appear,31 CRDS could be utilized to study the reaction kinetics by monitoring the ring-down time of radical species whose absorption is in the visible range in the future.

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References


