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Reaction of NO₂ with Selected Conjugated Alkenes

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Supporting Information

ABSTRACT: The gas phase reactions of selected alkenes (isoprene, myrcene, ocimene, and 1,3-cyclohexadiene) with NO₂ under dark condition have been investigated at $T \sim 298$ K and $P \sim 760$ Torr of purified air. The kinetic studies were performed under pseudo-first-order conditions using a large excess of NO₂ concentration to those of the alkenes. The rate coefficients (in 10^{-19} cm³ molecule⁻¹ s⁻¹) obtained are 1.1 ± 0.2 for isoprene, 2.5 \pm 0.3 for myrcene, 8.5 \pm 1.2 for ocimene, and 15 ± 1 for 1,3-cyclohexadiene. Several products were identified by using in situ Fourier transform infrared (FT-IR) spectrometry, and acetone was found to be the major product from the reactions of NO₂ with myrcene and ocimene, with a formation yield of $22 \pm 3\%$ and $26 \pm 7\%$, respectively. The oxidation products from the reactions of NO₂ with isoprene and 1,3-cyclohexadiene were found to be mainly nitro compounds identified by FT-IR spectroscopy. Reaction mechanisms were proposed to account for the products observed.



1. INTRODUCTION

In the atmosphere, alkenes are known to react with the main gas phase oxidants (OH and nitrate radicals, ozone, and chlorine atoms).^{1,2} It is commonly assumed that the reaction with NO₂ is a negligible atmospheric removal process for these species because the reaction rate coefficients of NO₂ with alkenes are slow under atmospheric conditions ($<10^{-20}$ cm³ molecule⁻¹ s⁻¹).³⁻⁶ However, this process could be important with dienes and polyconjugated alkenes during the smog chamber experiments where very often high concentrations of reactants are used. Indeed, some of the previous studies on the reactions of NO₂ with dienes and polyconjugated alkenes indicate that these reactions could contribute to their consumption under high NO_X conditions.^{4,5}

In this work, we report a kinetic and mechanistic study of the reaction of NO₂ with isoprene, 1,3-cyclohexadiene, myrcene, and ocimene. These VOCs have been chosen according to their emission rates into the atmosphere,⁷ and their reactivities toward NO₂ under dark conditions.^{8,9} The reactions rate coefficients were determined and the oxidation products tentatively assigned and quantified using FT-IR spectroscopy. So far, the reactions of NO₂ with ocimene and myrcene have been investigated only as interfering reactions but not as central systems.^{10,11} Acetone has been identified as a product from both reactions in the dark.^{10,11} However, the NO₂-initiated degradation mechanism of these two compounds is still not defined. Each of the NO₂-initiated oxidation of isoprene and 1,3-cyclohexadiene has been subject to only one study.^{3,8}

2. EXPERIMENTAL METHODS

The experiments have been carried out using the 7300 L ICARE Teflon chamber^{12,13} in $P \approx 1013$ mbar of purified air at $T = 298 \pm 2$ K with a relative humidity of ~5%. Reactants that are liquids under ambient conditions were introduced into the chamber by injecting aliquot amounts of liquid in a stream of purified air. Gaseous reactants were injected into the chamber using a calibrated gas cylinder equipped with pressure sensors. Two fans installed into the chamber ensured rapid mixing of reactants in few minutes. The concentrations of NO_X (NO and NO₂) were continuously monitored by a NO_X-NO-NO₂ analyzer using chemiluminescence method (Thermo Environment 42*i* or/and Horiba APNA-360). SF₆ was added to the gas mixture to assess the dilution rate due to added flow in the chamber compensating sampling flows by the monitors.

Organic reactants and products were monitored by an in situ Fourier transform infrared spectrometer (Nicolet 5700 Magna) coupled to a white-type mirror system, resulting in a 129–148 m optical path length. Infrared spectra were recorded every 2– 10 min by coadding 40–300 interferograms with a resolution of 1 cm⁻¹. A gas chromatograph coupled to photoionization detector (GC-PID, DXDZ GC-4400) was also used for the measurements of the reactants and gas phase products. The

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GC-PID is equipped with a VUV lamp with transparent light as low as 110 nm. The organic compounds were separated on a chromatographic column of 10% SE 30, PAW-HMPS (capillary column, 20 m length and 0.5 mm i.d.). An acquisition time of GC-PID was fixed at 30 min. Synthetic air (>99.999%) was used as the carrier gas with a flow rate of 10–20 mL min⁻¹. A six-port valve equipped with an injection loop of 1 mL connected to the GC, and the loop was flushed by the reaction gas mixture at 350 mL min⁻¹ for 1 min before injection. The temperature of the column was controlled from 40 to 60 °C according to the studied chemical systems.

2.1. Kinetic Measurements. Most of the kinetic experiments were carried out under pseudo-first-order conditions with $[NO_2]_0 \gg [unsaturated organics]_0$. Under these conditions, the unsaturated organics are subject to the following removal processes:

unsaturated org +
$$NO_2 \rightarrow products$$
 k

unsaturated org
$$\rightarrow$$
 losses (dilution and wall loss) $k'_{\rm L}$

where k is the bimolecular rate coefficient of the NO₂ reaction with the unsaturated organic and $k'_{\rm L}$ is the loss rate of the unsaturated organic due to both dilution and wall losses. Under these conditions, the unsaturated organics concentration—time profiles followed the pseudo-first-order rate law:

[unsaturated org]

= [unsaturated org]₀ × exp(
$$-(k'_{\rm L} + k')$$
) × t (I)

with $k' = k[NO_2]_0$. $(k' + k'_L)$ is derived from the slope of the plot of $ln([unsaturated org]_0/[unsaturated org]_t)$ versus reaction time. k'_L is obtained by following the decay rate of the unsaturated organics in the absence of NO₂. Plots of $(k' + k'_L)$ versus the concentration of the unsaturated organics for the reaction of NO₂ were linear and the absolute rate coefficients, k, were derived from the least-squares fit of the straight lines. The quoted error originates from two-standard deviation from the slope (2σ) .

The rate coefficients of both isomers of ocimene (cis and trans) were also measured using the relative rate method in which the relative disappearance rate of these species and that of myrcene used as reference compound were monitored in parallel in the presence of NO₂. The reaction rate coefficients can then be derived from the equation:

$$\ln\left(\frac{\left[\text{ocimene}\right]_{t_0}}{\left[\text{ocimene}\right]_t}\right) - k_{\text{L}}(\text{ocimene}) \times t$$
$$= \frac{k_{\text{ocimene}}}{k_{\text{myrcene}}} \left(\ln\left(\frac{\left[\text{myrcene}\right]_{t_0}}{\left[\text{myrcene}\right]_t}\right) - k_{\text{L}}(\text{myrcene}) \times t\right)$$

where $[\text{ocimene}]_{t_0}$ $[\text{ocimene}]_t$ $[\text{myrcene}]_{t_0}$ and $[\text{myrcene}]_{t_0}$ represent the concentrations of ocimene (cis - or trans - ocimene)and myrcene at reaction time t_0 , $[\text{ocimene}]_t$ and $[\text{myrcene}]_t$ are the corresponding concentrations at time t, k_{ocimene} and k_{myrcene} are the rate constants of the reaction of NO₂ with *cis*- or *trans*ocimene and myrcene, respectively. To account for dilution and wall processes, the pseudo-first-order decay rates of *cis*- and *trans*-ocimene and myrcene, $k_{\text{L}}(\text{ocimene})$ and $k_{\text{L}}(\text{myrcene})$, respectively, have been estimated in the absence of NO₂. Plots of $\ln[(\text{ocimene})_{t_0}/(\text{ocimene})_t] - k_{\text{L}}(\text{ocimene}) \times t$ versus $\ln[(\text{myrcene})_{t_0}/(\text{myrcene})_t] - k_{\text{L}}(\text{myrcene}) \times t$ should be linear and pass through the origin and have a slope $k_{\text{ocimene}}/k_{\text{myrcene}}$. The quoted uncertainties on the obtained rate coefficients originate from the uncertainties of the reference rate coefficient associated to the value of the slope (two standard deviation, 2σ).

2.2. Product Identification. Experiments were performed in conditions similar to those of the kinetic studies (pseudo-first-order conditions). Time–concentration profiles of identified products have been corrected from dilution losses using the dilution rate constant of SF₆. When SF₆ was not added to the mixture, a value of 1.1×10^{-5} s⁻¹ was used. Experimental time duration was extended to 4 h. Quantitative measurement of oxidation products using FT-IR spectroscopy was based on calibrated IR spectra when available. Time–concentration profiles of nitro compounds have not been corrected for the loss due to further reaction and loss to the wall. The molar product yields were derived from the average of each individual experiment and the resulted error corresponds to one-standard deviation (1 σ) on this average.

3. CHEMICALS

The purity and the origin of the chemicals used in this work are isoprene (99%, Sigma-Aldrich), 1,3-cyclohexadiene (98%, Alfa Aesar), myrcene (90%, Sigma-Aldrich), ocimene (\geq 90%, composed of \geq 25% of the *cis*-ocimene and \geq 58% for the *trans*-ocimene, International Flavors and Fragrances), di-*n*-butyl ether (\geq 99%, Sigma Aldrich), and NO₂ (1% in N₂, Air liquide). The sample of ocimene contains about 68% for the *trans*-ocimene and 32% for the *cis*-ocimene.

4. RESULTS AND DISCUSSION

4.1. Kinetic Measurements. The loss rates of the organic compounds in absence of NO₂ were on the order of $k_{\rm L} = (1.1 - 1.1)$ 1.6) \times 10⁻⁵ s⁻¹. To achieve the pseudo-first-order conditions, the initial concentrations (in molecules cm^{-3}) were [alkenes]₀ = $(1.8-16.3) \times 10^{13}$; $[NO_2]_0 = (0.31-3.0) \times 10^{15}$. Unsaturated organics were monitored using FT-IR spectroscopy over the following wavenumbers (in cm⁻¹): isoprene, 900; myrcene, 900; ocimene, 900 and 989; 1,3-cyclohexadiene, 3055 and 2837. Experimental durations were from 10 to 110 min. Typical plots of $\ln([alkenes]_0/[alkenes]_t)$ versus time under different concentrations of NO2 are displayed in Figure 1. The rate coefficients, k, are derived from the slopes of the linear fits originated from the plots of k' versus NO₂ concentrations (Figure 2). Experimental conditions for each run and the corresponding results can be found in the Supporting Information. The rate coefficient values derived (in cm³ molecule⁻¹ s⁻¹) at $T = 299 \pm 3$ K and P = 760 Torr are

isoprene + $NO_2 \rightarrow products$	$k = (1.1 \pm 0.2) \times 10^{-19}$
myrcene + $NO_2 \rightarrow products$	$k = (2.5 \pm 0.3) \times 10^{-19}$
ocimene + $NO_2 \rightarrow products$	$k = (8.5 \pm 1.2) \times 10^{-19}$

1,3-cyclohexadiene + NO₂ \rightarrow products $k = (1.5 \pm 0.1) \times 10^{-18}$

The quoted errors represent two-standard deviation (2σ) from the least-squares analysis.

In the gas phase, NO₂ coexists with its dimer N₂O₄ according to the equilibrium reaction $2NO_2 \leftrightarrow N_2O_4$. Using the equilibrium constant, $K = 2.5 \times 10^{-19}$ cm³ molecule⁻¹ s⁻¹, for this equilibrium, it is estimated that N₂O₄ in our system under the highest NO₂ concentrations used in this work (up to



Figure 1. Pseudo-first-order experiments from the NO_2 -initiated reaction of myrcene.

 3×10^{15} cm³ molecule⁻¹) was less than 0.08% of the NO₂ levels. At these concentrations and assuming the same reactivity, the dimer would not impact the chemistry occurring in our experimental system. The absolute method did not enable us to make a difference between the reactivity of the ocimene isomers (cis and trans) as the samples used in our

study contained both isomers and from the IR analysis we could not distinguish between these two species. Hence, we performed further measurements using the relative rate method and GC-PID as analytical tool. However, a significant overlap of both chromatographic peaks was observed. Therefore, the quantification has been performed using the peak height. Assuming the same response factors for both isomers, the ratios cis- and trans-ocimene in the reagent were found to be 28% and 72%, respectively, which was in agreement with the composition stated by the supplier. The initial concentrations of ocimene, myrcene, and NO₂ used were (in 10¹³ molecules cm⁻³) 4.9, 6.2, and 6.4, respectively. The initial concentration of NO (as impurity) in the gas mixture was about 2.5×10^{11} molecules cm⁻³. The presence of NO may lead to the possible formation of OH radicals through the reaction NO + HO_2 \rightarrow $OH + NO_{2}$, which could contribute to an additional consumption of the organic reactants, resulting in an extra uncertainty on the measured rate coefficients. Therefore, di-nbutyl ether was added in excess $(3.8 \times 10^{14} \text{ molecules cm}^{-3})$ to the gas mixture to suppress OH radicals formation (k(di-nbutyl ether + OH) = 2.8×10^{-11} cm³ molecule⁻¹ s⁻¹).¹⁴ The plot of $(\ln([\text{ocimene}]_0/[\text{ocimene}]_t) - k_L(\text{ocimene})_t)$ versus $(\ln([myrcene]_0/[myrcene]_t) - k_L(myrcene)_t)$ is shown in the Supporting Information. $k_{\rm L}(cis/trans-ocimene)$ and $k_{\rm L}$ (myrcene) represent the first-order decay rates of *cis/trans*ocimene and myrcene in the absence of NO2, which were estimated to be $\sim 1.4 \times 10^{-5}$ s⁻¹. Under these experimental conditions, the reaction of ocimene and myrcene with NO₂



Figure 2. Plot of absolute rate data for the NO₂ reactions with isoprene (a), myrcene (b), ocimene (c), and 1,3-cyclohexadiene (d).

Table 1. Summary of the Rate Coefficients of Studied Alkenes by Dark Reaction with NO_2 in the Gas Phase and Compared to the Literature^{*a*}

alkenes	<i>T</i> (K)	no. of runs	$k (\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$	ref
isoprene	298 ± 2	6	$(1.1 \pm 0.2) \times 10^{-19}$	this work, b
	NI	5	$(1.15 \pm 0.08) \times 10^{-19}$	3, b
	NI	1	$(1.8 \pm 0.3) \times 10^{-19}$	15, c
	AT	7	$(1.8 \pm 0.3) \times 10^{-19}$	4, c
	295 ± 2	NI	$(1.63 \pm 0.13) \times 10^{-19}$	16, d
	295 ± 2	NI	$(1.03 \pm 0.03) \times 10^{-19}$	5, b
	296	8	$(1.14 \pm 0.07) \times 10^{-19}$	6, d
myrcene	298 ± 1	7	$(2.5 \pm 0.3) \times 10^{-19}$	this work, b
	NI	NI	$(2.9 \pm 0.3) \times 10^{-19}$	10
	297 ± 2	NI	2.9×10^{-19}	11
	294 ± 2	1	$(2.6 \pm 0.2) \times 10^{-19}$	17, b
ocimene	299 ± 2	7	$(8.5 \pm 1.2) \times 10^{-19}$	this work, b
	297 ± 2	1	1.0×10^{-18}	11
	294 ± 2	NI	$(8.9 \pm 0.4) \times 10^{-19}$	17, d, f
cis-ocimene	297 ± 1	1	$(7.3 \pm 1.3) \times 10^{-19}$	this work, e
trans-ocimene	297 ± 1	1	$(7.3 \pm 1.8) \times 10^{-19}$	this work, e
1,3-cyclohexadiene	298 ± 1	7	$(1.5 \pm 0.1) \times 10^{-18}$	this work, b
	296	5	$(1.75 \pm 0.15) \times 10^{-18}$	8, b
			$(1.7 \pm 0.2) \times 10^{-18}$	4, e
			$(1.9 \pm 0.3) \times 10^{-18}$	4, c
			$(1.78 \pm 0.22) \times 10^{-18}$	5, b

^{*a*}AT: ambient temperature. NI: not indicated. ^{*b*}Pseudo-first-order conditions (loss rate of the alkene). ^{*c*}Second-order conditions (loss rates of both reactants). ^{*d*}Pseudo-first-order conditions (loss rate of NO₂). ^{*c*}Relative rate conditions. ^{*f*}Rate coefficients of isomers cis and trans identical within \sim 7%.

accounted, respectively, for 84% and 63% of their total decay. The values of the rate coefficient ratios obtained are k(trans-ocimene + NO₂)/k(myrcene + NO₂) = 2.9 ± 0.2 and k(trans-ocimene + NO₂)/k(myrcene + NO₂) = 2.9 ± 0.4. These results were placed on the absolute basis by multiplying the ratios by k(myrcene + NO₂) = (2.5 ± 0.3) × 10⁻¹⁹ cm³ molecule⁻¹ s⁻¹. The rate coefficients derived for the reactions of NO₂ with ocimene isomers are (in cm³ molecule⁻¹ s⁻¹):

cis-ocimene + NO₂
$$\rightarrow$$
 products $k = (7.3 \pm 1.3) \times 10^{-19}$

trans-ocimene + NO₂
$$\rightarrow$$
 products $k = (7.3 \pm 1.8) \times 10^{-19}$

Errors on the rate coefficients originated from the slopes of the linear fit adding the error to the rate coefficient of the reference compound. Absolute error on the rate coefficient is equivalent to two standard deviations (2σ) .

The rate coefficients compared to the literature data are summarized in Table 1. For isoprene, the reported values are in the range $(1.0-1.8) \times 10^{-19}$ molecules cm⁻³ s⁻¹ at 297 ± 2 K and $P \sim 760$ Torr and could be considered in fair agreement regarding the low reactivity and the difficulties to extract a precise rate coefficient value for this reaction under these conditions.

The rate coefficient value for the reaction of myrcene with NO₂ measured in this work is in good agreement with those obtained by Orlando et al.,¹⁰ Reissell et al.,¹¹ and Atkinson et al.¹⁷ Reissell et al.¹¹ indicated that in their work during the first 15–20 min of the reaction, the decay rate of myrcene was fast and decreased over the course of the reaction time. These authors suggested the OH radicals formation due to the presence of NO (NO + HO₂ \rightarrow NO₂ + OH) either originally present in the gas mixture or introduced with NO₂ might contribute to additional myrcene consumption.

For ocimene, the rate coefficient obtained in this work is in good agreement with those obtained by Reissell et al.¹¹ and Atkinson et al.¹⁷ as shown in Table 1. In addition, we have conducted one run to estimate the reactivity of ocimene isomers toward NO₂ and obtained k(cis-ocimene) = $k(transormene) \approx 7 \times 10^{-19}$ cm³ molecule⁻¹ s⁻¹.

Finally, the rate coefficient value for the reaction of NO₂ with 1,3-cyclohexadiene is also in good agreement with the previous measurements.^{4,5,8}

Examination of the reactivity of NO₂ with the four studied unsaturated VOCs shows the trend: k(1,3-cyclohexadiene) > k(ocimene) > k(myrcene) > k(isoprene). The four species possess a group of conjugated double carbon bonds (-C=C-C=C-), which are the key reactive sites. The different reactivities observed mainly depend on the molecular structure and sites of their substitution groups to the conjugated dienes, revealing that the reactivity increases with the substitution degree of -C=C-C=C- groups which has been observed in this study and elsewhere.^{5,6} The main difference between these four conjugated dienes is the degree of substitution. The difference in the reactivity of NO₂ with ocimene and myrcene shows that the reaction occurs mainly on the conjugated double bonds and the isolated $-C=C(CH_3)_2$ does not play an important role.

4.2. Oxidation Products. A set of three to five experiments have been performed for each reaction. Reactants were monitored using the same procedure as mentioned in the kinetic section (section 4.1).

FT-IR spectra from NO_2 -initiated reactions of myrcene and ocimene have been first subtracted from HNO_3 and HONO, showing the presence of acetone and formic acid among the oxidation products. Examples of IR spectra obtained are presented in the Supporting Information. Plots of acetone versus the consumption of myrcene and ocimene are displayed



Figure 3. Acetone formation versus the consumed concentration of myrcene (a) and ocimene (b) during the reactions with NO_2 .

accounted for $22 \pm 3\%$ and $26 \pm 7\%$, respectively. HCHO, HCOOH, and CO have been quantified at 2.0 \pm 0.8%, 0.7 \pm 0.2%, and 19 \pm 8%, respectively, for the reaction of NO₂ with myrcene and $2.1 \pm 1.6\%$, $1.1 \pm 0.5\%$, and $11 \pm 5\%$, respectively, for the reaction of NO₂ with ocimene. HONO formation yields have been estimated to $3.5 \pm 1.3\%$ for myrcene and $4.1 \pm 1.2\%$ for ocimene. Previous work conducted on the NO₂ initiated oxidation of myrcene and ocimene reported only acetone as the reaction product. The formation yield of acetone from myrcene oxidation obtained in the present work is in disagreement with the reported ones in the literature estimated at $11 \pm 3\%^{10}$ and ~37\%.¹¹ The acetone formation yield for ocimene oxidation is in agreement with that reported by Reissell et al.¹¹ (~28%). However, it has to be mentioned that the data of Reissell et al.¹¹ were derived from a single experiment and the use of a preconcentration system on adsorbent cartridge (Tenax) to collect gaseous compounds during analysis. Indeed, this method could lead to an overestimation of reaction products in the terpene- NO_X -air system.¹⁸ The difference observed with the reported value of Órlando et al.¹⁰ remains unexplained so far. We have not observed the formation of nitrogen-containing compounds as products of the reactions NO_2 + myrcene and NO_2 + ocimene. This may indicate that these types of products are not formed

at all or formed at concentration levels below the detection limit of our analytical system.

FT-IR analyses of isoprene/NO₂/air and 1,3-cyclohexadiene/NO₂/air mixtures have shown the formation in time of nitro-organic compounds, as shown the IR spectra on Figure 4a,b. In the case of isoprene, we have observed an increase of IR



Figure 4. FT-IR spectra of products recorded in the range 1900 -700 cm⁻¹ from the NO₂-initiated reaction of isoprene (a) and 1,3-cyclohexadiene (b). Panels A are the recorded spectra of isoprene and 1,3-cyclohexadiene, before NO₂ addition on the Figure 4a,b, respectively. Panels B have been recorded at 27 and 8 min after NO₂ addition for isoprene and 1,3-cyclohexadiene, respectively. Panels C have been recorded 2 and 2 h and 35 min after NO₂ addition for isoprene and 1,3-cyclohexadiene, respectively. Spectra in panels B and C have been subtracted from isoprene (Figure 4a) and 1,3-cyclohexadiene (Figure 4b) and known products (nitrous acid, nitric acid, and formic acid for isoprene/NO₂ system).

absorption bands centered at 1721, 1296, and 791 cm⁻¹ assigned to $-OONO_2$ and at 1373 cm⁻¹ assigned to $-NO_2$ groups.^{19,20} An example of the recorded spectra resulting from the NO₂ reaction with isoprene in 760 Torr of air is shown in Figure 4a. IR band features of identified product species have been subtracted from these spectra. Major IR bands from spectra B and C have been assigned to nitrogen-containing compounds, $-O_2NO_2$ and $-NO_2$ groups. However, in the absence of FT-IR reference spectra, their quantifications were not possible. Figure 5 displays an example of the temporal



Figure 5. Time concentration profiles of R-OONO₂ and R-NO₂ type products from NO₂-initiated reaction of isoprene in the dark.

profiles of the IR bands intensities attributed to the -OONO₂ and $-NO_2$ groups, obtained from the subtraction of the residual spectra assigned as reference spectra, versus the consumed fraction of isoprene over the course of the experiment. Figure 5 shows a rapid formation at an earlier stage of the reaction and a decrease over the reaction time for the R-OONO₂ compound while bands assigned to R-NO₂ show a linear increase before reaching a plateau against the consumed fraction of isoprene. HONO has been also observed at low concentrations, and its formation has been estimated to be $3.5 \pm 1.8\%$. HCHO, HCOOH, and CO have been identified among the oxidation products but are mainly formed as secondary products. Their primary formation yields have been estimated to be 1.3 \pm 0.6%, 0.7 \pm 0.5%, and 0.9 \pm 0.3% for HCHO, HCOOH, and CO, respectively. Analysis of NO2initiated oxidation of 1,3-cyclohexadiene data has shown intense peaks that can also be assigned to nitrogen-containing groups (Figure 4b). Panel B shows three strong bands located at 1721, 1296, and 790 cm⁻¹ characteristic of $-O_2NO_2$ groups, which could be attributed to the formation of nitrocyclohexenvlperoxy nitrate.⁸ From panel C, it can be seen that these compounds exhibit a strong reaction time variability. For a longer reaction time, peaks assigned to $-O_2NO_2$ groups decreased while the IR bands at 1280 and 822 cm⁻¹ increased. HONO was also identified from the reaction of NO₂ with 1,3cyclohexadiene with a yield $2.4 \pm 0.6\%$, in agreement with that reported by Jenkin et al.,⁸ estimated to be around 2%. In addition, Jenkin et al.⁸ have identified benzene among the oxidation products from the reaction of 1,3-cyclohexadiene with NO₂ at 0.41 \pm 0.02%. HCHO, CO, and HCOOH have been identified among the oxidation products but are mainly

formed from secondary sources. Their primary formation yields have been estimated to be <1%.

4.3. Reaction Mechanism. The suggested mechanistic schemes of the reaction of myrcene and ocimene with NO₂ are displayed in Figures 6 and 7, respectively. The reaction proceeds by addition of NO2 on conjugated double bonds >C=C-C=C<. For both compounds, the carbon in the sixth position is supposed to be unfavored due to a steric hindrance and a low electron delocalization. The expected addition site for NO₂ is that on carbons 5 and 8 for ocimene and on carbons 8 and 10 for myrcene, leading to the formation of the allylic C= $C-C^{\bullet}$ radical. Although leading to a less stable radical (C= $C-C-C^{\bullet}$) than the allylic radical, addition of NO₂ at position 7 on ocimene is also considered as a pathway leading to the observed acetone production. This leads to the formation nitroalkyl radicals (R–NO₂) through reaction 1. These addition sites are expected to occur due to the high stability of the nitroalkyl radical formed. It either reacts directly with O₂ to form a nitroperoxyalkyl radical (ROONO₂) and further with RO via reaction 5 or undergoes electron delocalization, leading to two other nitroalkyl radical intermediates as suggested through the equilibrium pathways 2.1. To lead to acetone formation, we have proposed an intramolecular cyclization of these intermediates (reaction 2.2 for myrcene and reactions 1.1 and 2.2 for ocimene), which would react quickly to form a nitroalkoxyl radical (RONO₂) through reaction 3 for myrcene and reaction 5 for ocimene. Acetone might be formed through the decomposition of the nitroalkoxyl radical (reaction 4 for myrcene and reaction 6 for ocimene). From the NO_2 reaction with myrcene, the decomposition of the nitroalkoxyl radical originated from the direct addition of O2, without any resonance may lead to the formation of a carbonyl compound and OCH₂NO₂ radical for myrcene + NO₂ reaction. This latter decomposes to form NO2 and formaldehyde. The low formation yield of formaldehyde (<1%) shows that this reaction occurs to a minor extent.

The reaction of NO₂ with 1,3-cyclohexadiene proceeds via NO₂ addition on the double bond, leading to the formation of a nitrohexenyl radicals.^{4,8} The addition of NO₂ on the terminal bond is favored due the stability—resonance of the radical formed. An unpaired electron is removed by electronic delocalization. A rapid reaction with oxygen leads to the formation of a nitroperoxyhexenyl radical, which further reacts with NO₂.

The mechanism proposed for the reaction of isoprene with NO2 is based on the studies of the reaction of 1,3-butadiene $(CH_2=CHCH=CH_2)$ with NO₂ proposed by Calvert et al.,² Atkinson et al.,⁵ and Niki et al.,¹⁹ which proceeds by electrophilic addition on the double bond >C=C<. A proposed scheme for the reaction of isoprene with NO₂ is given in the Supporting Information. As shown in the FT-IR spectra (Figure 4a), formation of $-O_2NO_2$ and $-ONO_2$ compounds have been identified from the beginning of the reaction. NO2 will add predominantly to C1 due to the formation of the most substituted nitroalkyl radical. NO2 addition on C4 is expected to lead to the formation of a less stable secondary alkyl radical, which may, however, also occur to some extent. Additions to C2 and C3 appear to be unlikely due to a low stability of the radical formed. The nitroalkyl radical, $R-NO_2$, will form a nitroperoxy radical (RNO_2-OO), which in turn reacts with NO₂, leading to a dinitroperoxy nitrate (RNO₂-OONO₂). The alkyl peroxynitrate may decompose back to the peroxy radical. The experiments



Figure 6. Proposed mechanism for the reaction of NO2 with myrcene in the dark.

showed that despite the presence of a relatively high NO₂ concentration, the infrared peroxynitrate band disappears when the reaction goes on (Figure 4a). This behavior may indicate an instability of this peroxynitrate, which decomposes followed by a self-reaction of the peroxy radicals. This pathway leads to the formation of an alkoxy radical which in turn reacts with NO₂ to form R–ONO₂, as observed by FT-IR measurement. The disappearance of peroxynitrate may be due to the following equilibrium ROONO₂ + NO₂ + M \leftrightarrow ROONO₂–NO₂ + M,⁵ or/and further reaction of RO₂ with residual NO formed over the course of the reaction, which turns into RO and NO₂. However, the peroxynitrate RO₂–NO₂ formed is thermally unstable and may promptly redissociate back to RO₂. The peroxy radicals will react rapidly with NO, leading to a rapid disappearance of the peroxynitrates:⁵

$$ROONO_2 + NO \rightarrow RONO_2 + NO_2$$

or the self-reaction of $R-O_2NO_2$, leading to the formation of $RONO_2$ and O_2 .

It should be noticed that the formation of oxidation products formed from NO₂ + alkenes in the absence of NO arise after RO₂ + RO₂ and RO₂ + HO₂ reactions. Therefore, the nature of the products formed and their yields may be dependent on the presence of mix of specific RO₂ radicals present in gas mixture and $[RO_2]/[HO_2]$ ratios, which then differ from laboratory experiments to ambient atmospheres.^{21,22} The presence of HONO among the products revealed that H-abstraction mechanism also may occur in the NO₂ reaction with the studied alkenes but to a minor extent compared to the addition mechanism. $^{\rm 8}$

5. ATMOSPHERIC IMPLICATIONS

The rate coefficients for the reactions of NO₂ with a selected series of important atmospheric alkenes (myrcene, ocimene, isoprene, and 1,3-cyclohexadiene) have been determined and found to be in the range $(0.1-1.5) \times 10^{-18}$ cm³ molecule⁻¹ s⁻¹. These values indicate that the reactivity of these species toward NO₂ is rather slow compared to that with other atmospheric oxidants such as OH, O₃, and NO₃ ($k_{\rm OH} = (1-2.5) \times 10^{-10}$, $k_{\rm O_3}$ = (0.1-12) × 10⁻¹⁶, and $k_{\rm NO_3}$ = (0.07-2.2) × 10⁻¹¹ cm³ molecule⁻¹ s⁻¹).^{2,23,24} However, high levels of NO₂ have been measured in different locations in the world, in particular in rapid economic development region such as China and India as reported by satellite measurements.²⁵⁻²⁷ These observations indicate that the contribution of this process to the atmospheric oxidation of the studied compounds, especially in anthropogenically influenced rural air areas,²⁸ may be not so negligible. This process may occur under dark as well as under light conditions and indoor or outdoor. On the other hand, assuming the concentrations for the following atmospheric oxidants: $[OH] = 2 \times 10^6$ molecules cm⁻³ for a 12 h daytime average, NO₃ = 5 × 10⁸ molecules cm⁻³ for a 12 h nighttime average, and NO₂ = 2×10^{13} molecules cm⁻³ (800 ppb, a very high level that could conceivably be found in a very heavily trafficked area under stagnation conditions). The NO₂ reactions with ocimene and 1,3-cyclohexadiene could, under



Figure 7. Proposed mechanism for the reaction of NO_2 with ocimene in the dark.

those conditions, be of some, albeit minor, importance. This may also be the case at the edges of fossil-fueled power plant plumes in polluted urban air masses, where mixing of NO2-rich power plant plumes interact with the VOCs of polluted air masses. The NO₂ level is often found to be higher in indoor environments than in outdoor ones due to internal sources originating from combustion processes such as cooking, cigarettes, unvented heaters, leading to a NO_2 concentration up to ppm level.^{29–31} Besides, secondary NO_2 can be produced from the dark conversion of NO by O_3 . Therefore, the NO₂ reaction is likely to occur in an indoor atmosphere with a higher level of alkenes. Hence, this process could constitute a non-negligible atmospheric source of the nitro-containing compounds in some specific areas. Indeed, the present work has shown that the reactions of NO2 with the investigated alkenes may lead to the formation of a series of nitrogencontaining oxygenated VOCs such as RNO₂ and ROONO₂, though this assumption was not confirmed through the use of authentic samples. This observation is of importance because the nitro-containing compounds are known to be toxic to humans.^{32,33} Hence, this process may contribute to the nitrocompounds budget even with a low reactivity. Furthermore, possible formation of nitrous acid from the reactions alkene + NO₂ has been observed in this study with formation yields in the range 2.4-4.1% for the investigated alkenes, indicating that this process may not contribute significantly to the HONO budget. The reaction of other organic compounds, having a larger reactivity toward NO₂, such as α -terpinene and α - phellandrene, may have higher contributions to both nitroorganic compounds and HONO.

ASSOCIATED CONTENT

Supporting Information

Summary of the experimental conditions and the results of the absolute rate study from the reaction of NO_2 with alkenes (isoprene, myrcene, ocimene, and 1,3-cyclohexadiene) is available. A graph obtained of the relative rate measurements from the NO_2 -initiated reaction of ocimene using myrcene as reference organic compound in the presence of an excess of di*n*-butylether can be also found. In addition, FT-IR spectra of products recorded in the range 1900–700 cm⁻¹ from the NO_2 -initiated reaction of the NO_2 -initiated reaction of the NO_2 -initiated reaction of the spectra of products recorded in the range 1900–700 cm⁻¹ from the NO_2 -initiated reaction of myrcene (a) and ocimene (b) are provided. A proposed mechanism of the NO_2 -initiated reaction of isoprene in the dark is also included. Complete author lists for refs 7 and 28 are provided. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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