Reaction of Iodine Atoms with Submicrometer Squalane and Squalene Droplets: Mechanistic Insights into Heterogeneous Reactions

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ABSTRACT: The gas-phase reaction of iodine atoms with hydrocarbon molecules is energetically unfavorable, and there is no direct evidence for iodinated product formation by either H abstraction or I addition reactions at ambient temperature. Here we consider the possible heterogeneous reaction of I atoms with submicrometer droplets composed of a saturated alkane, squalane (Sq), and an unsaturated alkene, squalene (Sqe). The investigations are performed in an atmospheric pressure photochemical flow tube reactor in conjunction with a vacuum ultraviolet photoionization aerosol mass spectrometer and a scanning mobility particle sizer. Squalane, a branched alkane, is unreactive toward I atoms within the signal-to-noise, and an upper limit of the effective reactive uptake coefficient is estimated to be $\chi^\text{sq}_I \leq 8.58 \times 10^{-7}$. In contrast, the reaction of I atoms with unsaturated submicrometer squalene droplets results in observable iodinated squalene products. The effective reactive uptake coefficient of I atom with squalene particles is determined to be $\chi^\text{sqa}_I = (1.20 \pm 0.52) \times 10^{-4}$ at an average I concentration of $1.5 \times 10^{14}$ molecules·cm$^{-3}$.

1. INTRODUCTION

Free radicals play key roles as initiators, propagators, or reactive intermediates of the complex chemical reactions involved in combustion, atmospheric chemistry, polymerization, plasma chemistry, biochemistry, and many other chemical processes.1–5 Although several aspects of homogeneous gas and liquid phase chemistry of free radicals are quite well understood, there remains significant uncertainty about the heterogeneous reactivity of free radicals with organic surfaces composed of long chain hydrocarbons.

The hydrocarbon’s free radical chemistry might considerably change with hydrocarbon phase (gas, liquid, or solid) or hydrocarbon degree of unsaturation (number of C= C bonds). The reactivity of organic particles is often considered within the context of known gas-phase reaction mechanisms in an effort to establish key similarities and differences between heterogeneous and homogeneous chemistry. However, rate coefficient predictions based upon gas phase structure–activity relationships cannot always be applied to heterogeneous reaction of free radicals with large hydrocarbons at the surface of a liquid droplet. Moise et al. showed that many types of chemical reactions that are slow in the gas phase are enhanced by several orders when they occur at an organic surface.6

Depending on the hydrocarbon degree of unsaturation the initial steps in hydrocarbon free radical chemistry might be H atom abstraction or free radical addition reactions. While the saturated hydrocarbon free radical reaction is initiated solely by the abstraction of a H atom by the radical species, the unsaturated hydrocarbon free radical reaction occurs either by H atom abstraction or by addition of radical species to the C= C double bond. These reactions result in the formation of alkyl radicals whose subsequent reactions are key steps in radical chain reactions in combustion chemistry, in the photochemical production of smog, or in heterogeneous aging of organic aerosols.

The halogens vary greatly in their reactivity toward hydrocarbons, considering a free radical initiation and possible chain reaction mechanism. In this context, it is of considerable interest to investigate the reactive behavior of I atoms, since iodine atoms and molecules are recognized to be the least reactive in the halogen activity series. For initiation, the I$_2$ molecule possesses intense absorption spectra in the visible, a region where few organic species absorb.7–8 The simplicity of iodine photolysis in conjunction with I atom attack on specific organic sites makes photolysis of iodine in the presence of organic compounds a potential source of structurally specific R$_2$, RO$_2$, and RO$_2$ radicals. Because of their unique chemistry, I atoms are frequently used as an initiator in polymerization reactions.

The gas-phase reaction of I atoms with a variety of alkanes10–13 and alkenes14–17 has been investigated experimentally as well as theoretically. While many aspects of the
homogeneous gas-phase chemistry of I atoms are well understood, little is known about the heterogeneous reactivity of I atoms with organic surfaces composed of long chain saturated and unsaturated hydrocarbons.

The reaction of squalane submicrometer particles with another halogen atom, Cl, has been investigated in this group in the absence and presence of oxygen.\textsuperscript{18} In the presence of O\textsubscript{2}, chlorinated reaction products are formed and an alkyl radical chain reaction propagates because of the presence of Cl\textsubscript{2}. In the presence of O\textsubscript{2} (as much as 20%) the chain reaction is observed to effectively be shut off, as the Cl\textsubscript{2}-propagated reaction instead produces oxygenated products.

The reactions of both squalane and squalene submicrometer particles with NO\textsubscript{3} have been studied as models for aerosol aging.\textsuperscript{19} The reactive uptake coefficient measured for squalene was found to be about 2 orders of magnitude larger than the one measured for squalane. In addition, the reactive uptake measured for squalene increases with the squalene degree of oxidation from an initial value of 0.18 on fresh squalene aerosols to 0.82 for an average of more than three NO\textsubscript{3} reactions per squalene molecule in the aerosol.\textsuperscript{19}

Recently, the kinetics and products of the heterogeneous OH-initiated oxidation of squalene submicrometer particles have been measured.\textsuperscript{20} In the presence of as much as 10% O\textsubscript{2} in the flow reactor the effective uptake coefficient for squalene was found to be 2.34 ± 0.07, and the reactive uptake increases with [O\textsubscript{2}] in the reactor. The effective reactive uptake coefficient of squalene (which is dominated by secondary radical chain reactions) is also in this case about 1 order of magnitude larger than the one measured in the case of squalane under similar experimental conditions but only 5% O\textsubscript{2}.\textsuperscript{21}

Here the heterogeneous reaction of iodine atoms with submicrometer particles composed of squalane or squalene is investigated. These hydrocarbons represent simple model systems that can also successfully reproduce the reactive behavior of other molecules that are important chemical constituents of biofuel droplets or organic aerosols. The molecular structures of both squalane and squalene are shown in Figure 1.

The heterogeneous reaction of I atoms with submicrometer squalane and squalene droplets is measured in a photochemical flow tube reactor with detection by a vacuum ultraviolet photoionization time-of-flight aerosol mass spectrometer (VUV-AMS). Aerosol mass spectrometry is used to measure heterogeneous reaction rates and product distributions in order to formulate a detailed reaction mechanism. The heterogeneous reactions of I atoms with submicrometer organic droplets composed of squalane or squalene are quantified to determine how the heterogeneous reaction rate of I atoms compares with previous measurements of Cl.

The main goal of the present studies is to evaluate simple model systems that are useful in developing a fundamental understanding of the mechanisms of heterogeneous reactions between small gas phase reactive species and submicrometer hydrocarbon particles. This paper also focuses on the reactive behavior of two molecules with similar structure but different reactivity due to the presence of C≡C double bonds.

2. METHODOLOGY

2.1. Experimental Setup. The experimental setup employed in the present study, schematically presented in Figure 2, consists of an atmospheric pressure photochemical flow tube reactor in conjunction with a scanning mobility particle sizer (SMPS) and an aerosol mass spectrometer (AMS). Details concerning the experimental technique are given elsewhere.\textsuperscript{21} Briefly, submicrometer droplets of pure squalane and squalene are generated by homogeneous nucleation of the organic vapor in a 0.350 standard liter per minute (slm) flow of N\textsubscript{2} (high purity, 99.9999%) through a 45 cm long Pyrex tube containing the liquid samples. The Pyrex tube is heated in a tube furnace to 135 °C. Vapors of the organic compound become supersaturated upon leaving the heated region and nucleation occurs, yielding a log--normal particle size distribution with a geometric standard deviation of ~1.3 and a mean surface-weighted diameter of ~180 nm (mean surface-weighted diameter defines the average particle size based on the number distribution weighted by the surface area of the distribution). Gas phase I\textsubscript{2} vapor is generated by a flow of N\textsubscript{2} (high purity, 99.9999%) over iodine crystals (Sigma-Aldrich, 99.999%) placed in a 6 mm copper U-shaped tube located just before the flow tube reactor and surrounded by a heating tape, so the I\textsubscript{2} vapor pressure and consequently the I\textsubscript{2} concentration in the flow stream are maintained at a constant value for the duration of each experiment. I\textsubscript{2} is mixed with the particle
stream in a balance of N₂ prior to entering the flow tube reactor. An O₂ scrubber (Sigma-Aldrich, 0.75 L Supelpure O₂ trap) is placed in the N₂ lines to reduce the residual oxygen level to less than 0.5 ppm.

The mixed gases and particles are then introduced into a 100 cm long, 2.5 cm inner diameter Pyrex reaction cell, mounted vertically with the injection port at the top end. Iodine atoms are generated along the length of the reaction cell by I₂ photolysis using 10 continuous output halogen lamps, each 20 cm in length, placed in pairs along the length of the reactor. The atomic I concentration is controlled by adjusting the molecular iodine, I₂, concentration in the flow tube; i.e., the concentration of I₂ is increased by increasing the N₂ flow through the iodine cell at a constant I₂ flow cell temperature.

In order to prevent heating of the reaction flow due to the proximity of the halogen photolysis lamps, the flow tube is surrounded by a jacket through which thermostated water is circulated, enabling regulation of the reactor temperature. The reactor temperature was kept at 40 °C. The flow tube was monitored using K-type thermocouples and is essentially constant (<0.5 °C variation) over the duration of the experiment. The reactor and the surrounding halogen lamps are located in an opaque enclosure in order to prevent any I₂ dissociation due to ambient light.

Gas flows are regulated by calibrated mass flow controllers (MKS) before being transferred to the reaction system via copper tubing. The total flow rate through the flow tube was fixed to 1 slm, which, based on the illuminated section of the flow tube of 100 cm, corresponds to a reaction time of 30 s.

The chemical composition of the droplets is monitored using a custom built, soft-ionization aerosol mass spectrometer (VUV-AMS) as described by Gloaguen et al. Briefly, a time-of-flight mass spectrometer is coupled to an aerodynamic lens for the purpose of focusing the particle beam and reducing the gas phase molecular concentration by means of pinhole plates described by Li et al. The transmitted particles impinge on a heated copper block (~110 °C) where they are flash vaporized. The VUV-AMS measures droplet composition by thermally vaporizing the droplets followed by VUV photoionization of the gas phase species. Tunable VUV radiation is produced at the Chemical Dynamics Beamline of the Advanced Light Source. A part of the remaining flow exiting the reactor is sampled by a scanning mobility particle size spectrometer (SMPS, TSI model 3936) to simultaneously determine the particle size distribution and number concentration.

2.2. Data Evaluation. To assess the experimental data, in particular, with respect to the reaction probability, the reaction models underlying the measured process are briefly outlined in this section. Details of the kinetic evaluation procedure are described elsewhere.

The rate of a heterogeneous reaction with a gas phase reactive species is expressed as a reaction probability or reactive uptake coefficient (\( \gamma_X \) where X = reactive species), which is defined as the fraction of collisions with a surface that results in a reaction. The uptake coefficient can be directly extracted by measuring the loss of a gas phase species exposed to a hydrocarbon surface. Often the gas phase reactive species are in very low concentrations and consequently very difficult or sometimes impossible to detect. An alternative approach to obtain an effective heterogeneous reaction probability (\( \gamma^{RH} \)) is to monitor the reaction via the condensed phase loss of the hydrocarbon (RH).

By use of a formalism developed by Smith et al. for the reaction of OH radicals with squalane particles, the reaction probability for the reaction of an I atom with submicrometer hydrocarbon droplets RH (here RH = squalane or squalene) can be written as

\[
\frac{\langle \rho_I \rangle}{\langle \rho_I \rangle_0} = \frac{4k_{RH}D_{surf}N_A}{6\bar{\gamma}M_{RH}}
\]

where \( k_{RH} \) represents a second order heterogeneous rate constant for the reaction of I atoms with the submicrometer hydrocarbon droplets, \( D_{surf} \) is the mean surface-weighted particle diameter, \( \rho_0 \) is the initial hydrocarbon molecule density (0.810 g cm⁻³ for squalane and 0.858 g cm⁻³ for squalene), \( N_A \) is Avogadro’s number, \( \bar{\gamma} \) is the mean speed of gas phase I, and \( M_{RH} \) is the molar mass of the hydrocarbon molecule (422.81 g mol⁻¹ for squalane and 410.72 g mol⁻¹ for squalene).

A significant difficulty associated with the present study was the lack of a suitable choice of a reference gas phase reaction to obtain the I atom concentration. Instead, to quantify the concentration of gas-phase I atoms, the chemical kinetics program Kinetica2003 developed by Richardson et al. is employed, together with the photolysis rate of I₂. Here, the program is used to model the time dependent I atom concentrations.

Because of the absence of appreciable side reactions of either I atoms or I₂ molecules with gas phase RH, or with O₂, H₂O, CO, or CO₂ that may be present in very small amounts in the reactor (as residual impurity in the N₂ carrier gas cylinder), the mechanism used to determine the I atom concentration in the flow tube consists of primarily three reactions (eqs 2a–2c).

\[
\begin{align*}
I_2 + h\nu \xrightarrow{k_2} I + I & \quad (2a) \\
I + I + N_2 \xrightarrow{k'} I_2 + N_2 & \quad (2b) \\
I & \xrightarrow{k''} \text{reactor walls} & \quad (2c)
\end{align*}
\]

The first reaction considered is the photodissociation of the I₂ molecule (eq 2a). \( k_2 \) denotes the I₂ photolysis rate constant, and its value is calculated as described in subsection 2.3. In addition to the photodissociation step, the I atom recombination (eq 2b) and I atom reaction with the reactor wall (eq 2c) are considered. \( k' \) represents the third order rate constant for the I atom recombination reaction and was set to a value of \( k' = 10^{32} \text{ cm}^3\text{molecule}^{-2}\text{s}^{-1} \) (N₂ as collision partner, used in the present study) as reported by Jenkin et al., while \( k'' \) is a rate constant for the I atom loss by reaction or recombination on the reactor wall. These three reactions primarily govern the I atom concentration in the flow tube reactor. The reaction and recombination of I atoms with the aerosol particles are not taken into account here because it accounts for less than 1% of the loss compared to the walls and the three-body recombination.

The loss of OH radical through reaction with the reactor wall was determined previously in this group by comparing the concentration of OH radical measured experimentally with that predicted from the concentration and photolysis rate of the OH
precursor without the wall. In this context, a value of 53 s\(^{-1}\) for the reaction constant of the loss of OH at the reactor wall was determined. A similar value was obtained for the case of Cl atom wall loss. Because of the poor reactivity of I atoms with tracer compounds, such as butane or hexanal, typically used in the experiments, the I atom wall loss rate could not be determined in a similar manner. Therefore, in the simulation \(k'\) was set to a value of 53 s\(^{-1}\), similar to that obtained for the OH wall loss. Possible errors that might result in determining the I atom concentration caused by the assumed \(k'\) value are discussed in section 3.3.

In addition the program requires the input of the I\(_2\) initial concentration and the concentration of the N\(_2\) carrier gas. The concentration of I\(_2\) is calculated from the N\(_2\) flow rate through the iodine cell, regulated by means of a mass flow controller, and the I\(_2\) vapor pressure at a certain temperature. Possible errors that might result in determining the I atom concentration caused by the assumed \(k'\) value are discussed in section 3.3.

**2.3. I\(_2\) Rate of Photolysis.** Ten halogen lamps (see section 2.1 for more details), which emit light between 380 and 1100 nm (as measured by an optical spectrometer, Ocean Optics HR4000), are used in the present experiment to produce atomic iodine via I\(_2\) photodissociation.

The iodine molecule is known to absorb light in the visible region of the spectrum. Two photodissociation channels can be accessed by means of visible light: (i) direct photodissociation via excitation of the A band, which leads to the formation of iodine atoms in the ground state, i.e., I\(_2^\text{A}\), and (ii) indirect dissociation via excitation of the B state. The B state is a bound state of I\(_2\) and has a radiative lifetime on the order of microseconds. However, the I\(_2\) molecule when excited into the B state can decompose to iodine atoms via collisional predissociation due to the coupling of the B state with three repulsive states. The predissociation rate of the B state increases with the pressure, i.e., number of collisions. Under our experimental conditions where the total pressure in the flow tube is \(\geq 760\) Torr, the relaxation of the excited I\(_2^\text{B}\) state molecule leads entirely to dissociation to I atoms (I\(_2^\text{B}\) \(\rightarrow\) I\(_2^*\) \(\rightarrow\) \(\text{I} + \text{I} + \text{M}\)).

The rate at which I\(_2\) is removed from the reaction cell as a result of photolysis can be calculated from the relation

\[
k_f(\lambda) = \int F(\lambda) \cdot \sigma(\lambda) \cdot \varphi(\lambda) \, d\lambda
\]

where \(F(\lambda)\) is the actinic flux (photons cm\(^{-2}\) s\(^{-1}\) nm\(^{-1}\)), \(\sigma(\lambda)\) represents the absorption cross-section (cm\(^2\) molecule\(^{-1}\)) while \(\varphi(\lambda)\) is the quantum yield for photodissociation. The actinic flux can be determined from the relation

\[
F(\lambda) = \frac{P}{S \cdot E}
\]

where \(P\) is the power emitted by the photodissociation lamps, measured by a optical power meter (Coherent, FieldMate), \(S\) is the irradiated surface area, and \(E\) is the photon energy. In the present calculation of the actinic flux we considered only the photon flux that passes once through the reactor. We estimate that the light back-reflected by the opposite lamps (see Figure 2) has a contribution of less than 10% to the total light flux that illuminates the cell.

The value of the absorption cross-section over the wavelength range from 390 to 750 nm, reported with 1 nm resolution, is available from the literature. For our calculations, at a value of 1 across the whole 390–750 nm spectrum.

Figure 3 displays the wavelength dependence of the I\(_2\) photolysis rate constant calculated at 1 nm resolution.

**3. RESULTS AND DISCUSSION**

The present investigations are designed in part to understand the pertinent difference between two organic systems with similar structure but different reactivity and mechanistic pathways as a result of carbon–carbon single bonds versus carbon–carbon double bonds. In this context, two sets of experiments were performed to investigate the heterogeneous reaction of saturated squalane and unsaturated squalene submicrometer hydrocarbon particles with I atoms.

As discussed above, the reactions of squalane and squalene submicrometer particles with I atoms are investigated in the photochemical flow tube reactor. In these experiments it is assumed that the reactant (squalane or squalene particles) is removed solely because of the reaction with iodine atoms. To verify this assumption, an initial set of experiments are performed to investigate the reaction of submicrometer droplets of squalane and squalene with molecular I\(_2\) (photolysis lamps off). No evidence of product formation is observed within the signal-to-noise of the photoionization mass spectrometer after the combination of I\(_2\) alone with either squalane or squalene particles over the typical residence time (30 s) used in the present work, even after increasing the I\(_2\) concentration to \(5 \times 10^{16}\) molecules cm\(^{-3}\). For experimental reasons, \(5 \times 10^{16}\) molecules cm\(^{-3}\) was the highest I\(_2\) concentration that could be achieved inside of the flow tube reaction for all experiments presented here. Furthermore, to test for the possible photolysis of the reactants with the lamps used in the present work, mixtures of the reactants (squalane or squalene particles) in nitrogen in the absence of iodine are irradiated; no photolysis of any of the reactants is observed.

**3.1. Reaction of I Atoms with Squalane Submicrometer Droplets.** Squalane (C\(_{30}\)H\(_{62}\)), a branched alkane, containing 8 primary, 16 secondary, and 6 tertiary carbon atoms, represents an ideal model system to mimic the variety of reactive carbon sites typically found on hydrocarbon surfaces and that may also occur in ambient organic aerosols. Abstraction is the only path available for the reaction of I.
atoms with alkanes. For this purpose the photoionization mass spectrum of unreacted squalane (no I atoms in the flow tube, photolysis lamps off) is recorded and compared to the photoionization mass spectrum of squalane reacted with I atoms for 30 s in the flow tube (I$_2$ in the flow tube, photolysis lamp on). No changes in the recorded product mass spectrum, compared to that of unreacted squalane, are observed by turning the halogen photolysis lamps on in the presence of I$_2$. The measurements were repeated several times at different I atom exposures by keeping the flow rate (i.e., reaction time) constant and varying the concentration of I atoms. No evidence for product formation is observed, even when increasing the concentration of I atoms in the flow tube up to $2.5 \times 10^{14}$ molecules-cm$^{-3}$, by increasing the amount of I$_2$ precursor molecule up to $5 \times 10^{16}$ molecules-cm$^{-3}$. For experimental reasons, $2.5 \times 10^{14}$ molec-cm$^{-3}$ was the highest I atom concentration that could be achieved inside the flow tube reactor. Under these experimental conditions, it appears that squalane submicrometer droplets react slower with I atoms than the experimental time scale allows.

On the basis of the background noise signal of the AMS (a maximum of 1 count per s) in the present experiment, peaks smaller than 0.02% of the unreacted squalane molecular ion intensity cannot be detected using the AMS technique. The absence of any type of reaction products in the reacted squalane mass spectrum indicates that either no products are formed or the amount of formed products is below the detection limit.

To consider the possibility of whether any alkyl radicals that are formed could be trapped chemically because of the reaction with an adjacent squalane molecule, i.e., by abstracting an H atom and re-forming squalane rather than reacting with an I atom or I$_2$ molecule, the reaction of squalane particles with iodine atoms was investigated in the presence of 20% O$_2$. If formed by an initial H atom abstraction by the I atom, an alkyl radical can react with O$_2$ to form peroxy radical. The peroxy radical self-reaction will then result in the production of higher molecular weight products with one oxygenated functional group. However, no evidence for oxygenated species is observed in the recorded mass spectra within the signal-to-noise ratio of the present experiment. This suggests that H abstraction followed by alkyl radical reactions is not occurring for the experimental conditions and with this signal-to-noise.

3.2. Reaction of I Atoms with Squalene Submicrometer Droplets. The reaction of squalene with I atoms is investigated using the same procedure as described for squalane particles. Squalene (C$_{30}$H$_{50}$) is an alkene that has 6 double bonds and 8 primary, 16 secondary, and 6 tertiary carbon atoms, making it an ideal proxy to represent the different kinds of reactive carbon sites typically found on hydrocarbon surfaces. In general, the course of reaction between an I atom and an alkene might include simultaneous contributions from addition, direct abstraction, and addition—elimination channels in contrast to the reaction of an I atom with alkane, where abstraction is the only path available. Addition is expected to be the major channel in the reactions of I atoms with squalene particles, which will proceed by I atom adding to one of the squalene double bonds and simple C−I bond formation. The formed adduct is an unstable radical and will further react. A second channel, consisting of direct abstraction of a hydrogen atom, might be also possible. The occurrence of this channel is more likely in the case of I atom reactions with alkenes than for the reactions with alkanes. This can be explained by the lower value of the C−H bond dissociation energy for allylic hydrogen compared with that in alkanes and also by the possibility of an indirect addition—elimination process in the H atom abstraction.

Figure 4 displays the photoionization mass spectra of squalene particles (a) before and (b) after reaction with I atoms, recorded at a photon energy of 9.5 eV after evaporation of the reacted particles. A higher photon energy is observed to yield larger ion signals, but those photon energies also increase the fragmentation of the parent ions. Therefore, scans of multiple photon energies were performed (from 7.4 to 11 eV photon energy) and a compromise between overall signal strength and product fragmentation was made.

Before the reaction (photolysis lamps off), the photoionization mass spectrum consists of a main peak at m/ z = 410 corresponding to the squalene molecular ion. No peaks with masses higher than squalene are observed in the absence of I atoms. Smaller peaks at masses lower than the mass of squalene are also observed, which correspond to squalene fragmentation due to dissociative photoionization or thermal dissociation (Figure 4a). When the photolysis lamps are turned on in the presence of I$_2$, the squalene peak decreases in intensity and the formation of reaction products is clearly observed (see Figure 4b). The reaction products appear in clusters of peaks separated by 126 amu. The first cluster of peaks is centered at m/z = 536, while the second cluster of peaks has the highest intensity peak located at m/z = 662. On the basis of the mass difference from the squalene molecular ion peak, the peaks located at m/z = 536 and 662 are consistent
with products in which H atom(s) is(are) replaced by I atom(s) during the reaction. On the basis of the mass spectrum of reacted squalene particles shown in Figure 4b, no peaks are detected above the background signal at masses larger than \( m/z = 664 \).

There is neither evidence for products containing more than two I atoms nor products that involve polymerization after the I atom addition to the squalene molecule in the experiment. This indicates that within the signal-to-noise ratio of the present experiment either higher molecular weight products are not formed or they undergo dissociative photoionization during the ionization. Another explanation for the lack of higher molecular weight products \( (m/z > 664) \) in the squalene product mass spectrum might be due to their decomposition during the thermal vaporization of the aerosol prior to ionization. However, a closer look in the fragments region of the reacted squalene particle mass spectrum (Figure 4b) reveals that apart from three new peaks at \( m/z = 127, 128, \) and 254 corresponding to an I atom, HI molecule, and \( I_2 \) molecule, the mass spectrum remains essentially unchanged compared to the same region of unreacted squalene mass spectrum. The origin of atomic I and HI in the product mass spectra will be discussed later in this section. The \( I_2 \) molecule detected in the mass spectra can be explained by the recombination of the I atoms in the particle, because no \( I_2 \) signal is detected in the mass spectrum when the photolysis lamps are off (no I atoms are produced in the flow tube reactor). Therefore, we can rule out the possibility that \( I_2 \) signal observed in the product mass spectrum originates from \( I_2 \) dissolved into the hydrocarbon particle and subsequently liberated from the particle during the vaporization on the copper block. Consequently, the observed reaction is exclusively initiated by the I atoms formed in the gas-phase.

Finally, the formation of polymerized products cannot be excluded. It might be possible that such products are formed, but they are not effectively detected because of the fact that their volatility might be much lower than the squalene volatility. As mentioned above, the temperature of the copper block was kept at 110 °C in our experiments to preserve the squalene molecular ions.

Apart from the iodinated products, several peaks at masses slightly lower than the mass of squalene, such as \( m/z = 409, 408, \) etc., are also observed (see Figure 4c, left panel). These peaks correspond to dehydrogenated squalene. As can be seen in Figure 4c middle and right panels, dehydrogenated iodinated products are also observed. The possible origin of these peaks is discussed later in this section.

Figure 5 displays the 402–416 amu region of the squalene mass spectra recorded before (Figure 5a) and after (Figure 5b–f) exposure to different amounts of I atoms. The concentration of I atoms in the flow tube is (b) 5.007 × 10^{13} \text{molecules-cm}^{-3}, (c) 8.212 × 10^{13} \text{molecules-cm}^{-3}, (d) 1.450 × 10^{14} \text{molecules-cm}^{-3}, (e) 1.951 × 10^{14} \text{molecules-cm}^{-3}, and (f) 2.445 × 10^{15} \text{molecules-cm}^{-3}.

Figure 5. The 402–416 amu region of the squalene mass spectra recorded (a) before and (b–f) after the exposure to different amounts of I atoms. The concentration of I atoms in the flow tube is (b) 5.007 × 10^{13} \text{molecules-cm}^{-3}, (c) 8.212 × 10^{13} \text{molecules-cm}^{-3}, (d) 1.450 × 10^{14} \text{molecules-cm}^{-3}, (e) 1.951 × 10^{14} \text{molecules-cm}^{-3}, and (f) 2.445 × 10^{15} \text{molecules-cm}^{-3}.
atoms. Thus, the peaks observed at \( m/z = 411, 412, 537, 538, 663, 664 \) correspond to \(^{13}\text{C}\) isotopes of squalene and its iodinated products for singly and doubly \(^{13}\text{C}\) substituted molecules. The relative intensities of isotopic peaks are consistent with the carbon chain length given the instrumental uncertainty.

Figure 6 shows a tentative reaction mechanism for the I atom reaction with submicrometer squalene droplets based on reaction products observed in Figure 4. Multiple isomers are expected to be formed for each of the suggested pathways. In addition some of the indicated reactions can occur both at the particle surface as well as within the particle. The initial reactions involve either the addition of an I atom to one of the six squalene double bonds (Figure 6, step 1) to form RI-(SqeI·) or the H atom abstraction by the I atom (Figure 6 step 1’) to form R·-(Sqe-H) alkyl radicals.

Since direct addition of the I atom to a double bond yields a radical species (as already mentioned above, to form Sqel·), it is likely that hydrogen abstraction takes place immediately after, resulting in the formation of the (Sqe-H)I reaction product (Figure 6, step 2). The addition of the I atom to a squalene double bond might also be followed by HI elimination and consequently to form a Sqe-H· radical species (Figure 6, step 1a), in an addition–elimination reaction fashion.

Figure 6. Proposed mechanism for the reaction of squalene with I atoms. The rounded rectangles indicate products observed in this study. Red colored items ((Sqe-H)I and (Sqe-2H)I₂) are the first and second generation squalene products, while in blue color (Sqe-H, Sqe-2H, (Sqe-2H)I and (Sqe-3H)I) are the products formed as a result of dissociative photoionization of (Sqe-nH)I, products, which result in the elimination of HI or I atom (see text for more details). The bold arrows indicate the reaction path that is expected to have the major contribution to the formation of the detected products. The mechanism as proposed is relevant for any of the squalene \( \text{C} \equiv \text{C} \) double bonds.
The SqeI· alkyl radical might also gain a hydrogen atom from an adjacent hydrocarbon molecule and could form SqeHI. Previous studies by Lee et al. on NO oxidation of squalene particles showed that a H atom is abstracted from an adjacent squalene molecule by the nitroalkyl radical, which is formed by the initial NO addition to the C=C double bond of the squalene particle. However, on the basis of the signal-to-noise ratio of the present experiment, no evidence for the SqeHI product formation is observed. The absence of SqeHI in the product mass spectrum may also be due to its dissociative photoionization to yield HI and a squalene molecule. Since no other products that might originate from SqeHI are present in the recorded mass spectra (as observed by Lee et al.), this path is not considered in the reaction scheme in Figure 6.

As already mentioned above, direct hydrogen atom abstraction by the I atom might also be possible (Figure 6, step 1'). Although the results observed here for squalene particles suggest that the I atom is essentially inert toward saturated bonds, the presence of C(sp^3)–H bonds in the squalene structure that are allylic in character and, in consequence, have a significantly lower bond energy makes those bonds more reactive than their equivalent in squalane. The H-abstraction at these bonds is more exothermic and has considerably lower activation energy. Thus, the enthalpies of abstraction from allylic carbons are typically between −86 kJ·mol^−1, while abstraction reactions from saturated hydrocarbons have enthalpy values between −56 and −43 kJ·mol^−1, depending on the type of abstracted H. The resulting radical species Sqe· further reacts either with an I atom or with an I₂ molecule to form (Sqe·H) iodinated product (Figure 6, step 2'). Both reaction channels (Figure 6, step 1') and (Figure 6, step 1') in Figure 6 lead to the formation of (Sqe·H)I via step 2 and step 2' in Figure 6. However, because of the experimental limitations, we cannot distinguish between the contributions of channels 1 and 1' (cf. Figure 6) that lead to the observed (Sqe·H)I reaction product. Nevertheless, the H abstraction channel (Figure 6, step 1') is expected to have a minor contribution to the observed reaction products with respect to the addition channel (Figure 6, step 1).

Dissociative photoionization of (Sqe·H)I leads to the formation of multiple fragment ion peaks (see peaks below mass m/z = 410 in Figure 4c). For instance, the blue colored species in Figure 6 (Sqe-H and Sqe-2H) might correspond to Sqe-H (m/z = 409) and Sqe-2H (m/z = 408) generated via dissociative photoionization of (Sqe·H)I via I and HI loss, respectively. The exact origin of the peaks observed at masses m/z = 407–403 in Figure 4c, left panel, is not fully understood. The peaks corresponding to odd masses, e.g., m/z = 407, 405, 403 are radicals, and therefore, they should be exclusively formed via dissociative photoionization of higher molecular mass species formed in the flow tube during the reaction. The peaks corresponding to even masses might be produced by dissociative photoionization, but they could also be a result of some complex chemical reactions, such as particle phase chemistry as already mentioned above.

A similar scenario as described above for (Sqe·H)I product formation is assumed to be followed in the case of the (Sqe-2H)I₂ product. Iodine atom attack on the first generation product (Sqe·H)I (Figure 6, step 3) followed by hydrogen abstraction would result in the formation of the second iodinated product, (Sqe-2HI)I, observed in the experiment. A second channel, contributing to the formation of (Sqe-2HI)I, species, participating in an addition–elimination scenario, might also be possible (Figure 6, steps 3 and 3a). A minor contribution to the formation of (Sqe-2HI)I₂ might be via hydrogen atom abstraction from (Sqe·H)I by the I atom followed by reaction either with the I atom or with the I₂ molecule (Figure 6, steps 3' and 4').

As mentioned above, peaks corresponding to (Sqe·nHI)I (n = 2, 3, 4, 5) are detected in the mass spectrum (Figure 4c, middle panel). These peaks might be attributed to dissociative photoionization of (Sqe-2HI)I₃. For example, (Sqe-2HI)I and (Sqe-3HI)I could be attributed to dissociative photoionization of (Sqe-2HI)I₂ (cf. reaction steps 4a and 4b in Figure 6).

Extremely small peaks corresponding to (Sqe·nHI)I, (n = 3, 4), are also observed (cf. Figure 4c, left panel). They suggest, considering a similar formation mechanism as discussed above, that (Sqe-3HI)I is also potentially formed. However, (Sqe-3HI)I is not detected in this experiment, most likely because of the low amount of this product that is formed and the fact that it also subsequently undergoes dissociative photoionization.

According to the reaction mechanism proposed in this work, double bonds can be re-formed during the reaction, e.g., species like (Sqe-2HI)I₂ in Figure 6, and consequently more than six addition reactions could occur if the reaction sequence reaches completion. The main reaction mechanism that leads to the formation of (Sqe·H)I and (Sqe-2HI)I₂ is attributed to the I atom addition mechanism. Therefore, in Figure 6 this path is indicated by bold arrows.

The tentative reaction mechanism proposed in this work can explain the formation of iodinated squalene reaction products as observed in the experiment. However, the mechanism of squalene reaction with the iodine atom might proceed in a more complex manner.

The kinetic evolution of squalene and its first two groups of higher molecular weight iodinated reaction products, (Sqe·H)I and (Sqe-2HI)I₂, as a function of iodine atom exposure is shown in Figure 7. In all the results reported here, where kinetics is measured, the I atom exposure is varied by keeping the flow rate, i.e., reaction time, constant and varying the I atom concentration. For comparison, Figure 7a displays the normalized decay of squalene ([Sq]/[Sq]₀) and squalene ([Sqe]/[Sqe]₀) as a function of I exposure. The kinetic evolution of squalene iodinated products is displayed in Figure 7b and Figure 7c. As can be seen in Figure 7b and Figure 7c, both (Sqe·H)I and (Sqe-2HI)I₂ iodinated products exhibit a complex dependence on the I atom exposure. Each product is formed followed by a decay at higher I exposures in the case of (Sqe·H)I (see Figure 7b). The (Sqe-2HI)I₂ signal is observed to reach a plateau at higher I exposure (see Figure 7c).

3.3. Effective Uptake Coefficient. The normalized decays of squalene ([Sq]/[Sq]₀) and squalene ([Sqe]/[Sqe]₀) as a function of I exposure (Figure 7a) are further used to determine the effective reactive uptake coefficients.

Upon the basis of the estimated sensitivity of the experimental setup, we derive an upper limit for squalane reactive uptake coefficient. Thus, the normalized squalane signal ([Sq]/[Sq]₀) plotted versus (I)ᵗ (red circles in Figure 7a) is fitted to a single exponential decay function (red solid line in Figure 7a) to determine k_sq. The effective uptake coefficient is subsequently computed using eq 1, and an upper limit for k_sq ≤ 8.58 × 10⁻¹⁶ cm³·s⁻¹·molecule⁻¹ with the I atom exposure, is obtained from a single exponential decay fit to the squalane trace (black solid line in

dx.doi.org/10.1021/jp5085247| J. Phys. Chem. A 2014, 118, 10688–10698
Figure 7. Kinetic evolution of (a) squalane (open red circles), squalene (open black diamond), and (b, c) squalene first two iodinated products ((Sqe-H)I and (Sqe-H2)I2) measured as a function of iodide exposure. The symbols correspond to the measured data, while the solid lines in panel a correspond to a fit using single exponential function. The solid lines in panels b and c represent the fit to the experimental data assuming that the (Sqe-H)I and (Sqe-H2)I2 reaction products are formed with equal probability; see text for more details.

Figure 7a). The effective uptake coefficient is then computed using the same eq 1. A value of $\gamma_{Sqe}^{1e} = (1.20 \pm 0.52) \times 10^{-4}$ for the squalene reactive uptake coefficient is obtained at an average I concentration of $1.5 \times 10^{14}$ molecules-cm$^{-3}$.

An additional source of error is due to the error from the determination of the I atom absolute concentration. In order to see how the effective uptake coefficient explicitly depends on the absolute I atom concentration, the value of the reactive uptake coefficient was calculated for two extreme cases. The concentration of I atoms in the flow tube is assumed to be 5 times lower or 5 times higher than obtained using the Kinetica2003 software. The obtained effective uptake coefficients are $(6.20 \pm 2.66) \times 10^{-4}$ and $(2.50 \pm 1.07) \times 10^{-5}$, respectively. As expected, the value of the reactive uptake coefficient changes by assuming different I atom concentrations in the flow tube reactor (5 times lower and 5 times more than the I atom concentration calculated using Kinetica2003 software), but even in these extreme cases the values for the effective uptake coefficient are still very small, corresponding to a very slow reaction.

The kinetic evolution of (Sqe-H)I and (Sqe-H2)I2 reaction products displayed in Figure 7b and Figure 7c are fit using eq 5 to approximate the rates of formation of these products. Equation 5 describes a sequential oxidation mechanism in a particle that is well mixed on the time scale of the reaction.

$$\frac{[(\text{Sqe-nH})I]_n}{[\text{Sqe}]_0} = B_n \frac{(k \cdot \langle I \cdot t \rangle)^n}{n!} \exp(-k \cdot \langle I \cdot t \rangle)$$

In eq 5 $k$ represents the second order heterogeneous rate constant while $n$ denotes the product generation. In the case of unreacted squalene $n = 0$. $B_n$ is an adjustable parameter to account for differences in isotope abundance, VUV photoionization efficiency, and fragmentation patterns of the oxidation products and squalene. For simplicity, $k$ in eq 5 is fixed to the value obtained from the exponential fit of the squalene decay curve shown in Figure 7a. The kinetic evolution of the squalene products ((Sqe-H)I and (Sqe-H2)I2) is then modeled to fit the data, using eq 5, by exclusively adjusting the $B_n$ parameter, with $n$ characterizing the iodination generation. Model fits to the (Sqe-H)I and (Sqe-H2)I2, experimental data are displayed in Figure 7b and Figure 7c as solid lines. Within the experimental error, the overall kinetic evolution of the parent compound and its first two generations of reaction products can be reasonably fit using a single rate constant. This indicates that for the squalene + I reaction $\gamma_{Sqe}^{1r} = (1.20 \pm 0.52) \times 10^{-4} \approx \gamma_{Sqe}^{1e} \approx \gamma_{Sqe}^{2e}$. However, as double bonds are consumed, the reactive collision probability will decrease.

The calculated value of the effective uptake coefficient $(\gamma_{Sqe}^{1e} = (1.20 \pm 0.52) \times 10^{-4})$ indicates that on average only one squalene molecule is reacted in a particle per 10 000 collisions between I atoms and the squalene particles. Therefore, I atoms might diffuse within the particle instead of directly reacting with or scattering from the particle surface. Slow diffusion into the particle and fast reactive loss would suggest that reactions occur close to the gas-surface interface, whereas fast diffusion and slow reactions would result in reactions throughout the bulk of a particle. The value of reacto-diffusive length $(D_{\text{RD}})$ might give some indication about the nature of this reaction. $D_{\text{RD}}$ represents the typical distance an atom or a molecule will diffuse before reacting, and it is consequently a measure of the concentration gradient of the reactant in the aerosol.

$$D_{\text{RD}} = \sqrt{\frac{D_p}{k^2}}$$

where $D_p$ (cm$^2$-s$^{-1}$) represents the diffusion coefficient of the I atom in the particle, $k^2$ (s$^{-1}$) is the pseudo-first-order rate constant for reaction of I in the particle, and $k^2 = k^0[Sqe]$. $k^0$ is the second-order heterogeneous rate constant for the reaction of an I atom and squalene particle and is here equal to $k_{Sqe}$. $D_p$ is estimated to be $10^{-3}$ cm$^2$-s$^{-1}$.

By use of eq 6, the reacto-diffusive length is then estimated to be $\sim 69$ nm, which suggests that I atoms might significantly diffuse inside the particle prior to reaction. However, this value is just an estimate, since it does not take into account the loss of I atoms, possibly by recombination on the surface of the droplet or inside the droplet, which competes with the I atom addition to the double bond of squalene.

For comparison we calculate the reacto-diffusive length in the case of Cl atoms with squalane particles in the presence of 20% O$_2$ with O$_2$ acting as scavenger for phase-phase secondary chain chemistry propagation. This is known as a very reactive system and has a much larger value of the reactive uptake coefficient $(\gamma_{Sqe}^{1e} = 0.65 \pm 0.07)$ than the one measured in the case of the I + squalene reaction. By use of the same procedure as described above to calculate the reacto-diffusive...
length, a value of $l_{\text{diff}} \approx 0.66$ nm is estimated for the Cl + squalane reaction in the oxygen presence. This length is on the order of a few particle layers and is consistent with the reaction occurring very close to the surface of the particle rather than inside the particle. Unlike more reactive systems where the reaction occurs in a near surface layer, we would expect that I atoms may diffuse significantly into the particle before reacting because the reacto-diffusive length is near the value of the particle radius.

The gas-phase reaction between different alkenes and I atoms has been investigated for a variety of reaction temperatures by a number of groups.14,15,38 No iodinated product formation has been observed for temperatures slightly above ambient temperature. In addition, the presence of catalytic amounts of I atom, in the alkenes + I atom reaction, results in a positional isomerization of the investigated alkenes.

Compared to the gas phase reactions, there are potentially more highly coupled reactions taking place between reactants, products, and intermediates in the case of heterogeneous reactions. As a consequence, reactions that normally do not take place in the gas phase are found to have an enhanced surface reaction probability. Various studies revealed that the surface reactions induced by small radicals or atoms can be enhanced by several orders of magnitude over analogous bimolecular reactions in the gas phase. Moise et al.6 measured large surface enhancement factors for gas phase reactions that are slow ($k < 10^{-15}$ cm$^3$·mole$^{-1}$·s$^{-1}$) because of high activation energies, in contrast with fast reactions ($k < 10^{-12}$ cm$^3$·mole$^{-1}$·s$^{-1}$), which exhibit minimal surface enhancement. The exact mechanism for this reaction enhancement remains currently ambiguous but may originate, in part, from multiple scattering or trapping of reactant species at the interface, thus increasing the overall chance of a reactive encounter.56−38 Furthermore, Wanda et al. also observed an enhancement in the case of ozone reaction with a phospholipid film compared to a typical gas phase ozone−alkene reactions.39 Their results in conjunction with molecular dynamics simulation suggest that the observed enhanced reactivity compared to the one in the gas phase may be due to trapping of ozone in the porous hydrocarbon layer, which increases the probability of an encounter of O$_3$ with a double bond.

On the basis of the calculated value of reacto-diffusive length of iodine into squalene particles, which has a value comparable to the squalene particle radius, it is likely that the observed reaction between squalene submicrometer droplets and I atoms is due to the long time trapping of reactive I atoms in the particle, which considerably increases the probability of an encounter of the I atom with a C=C double bond. The trapping and consequently the multiple interactions of the reagents, i.e., 10,000 collisions or more, in the aerosol particles are the key to why iodinated hydrocarbons are formed at room temperature just in the aerosol particle and not in the gas phase.

4. CONCLUSION

The present investigations reveal new insights into a category of reactions with low probability in the gas phase but that present an enhanced probability in the heterogeneous phase. The reactions of I atoms with submicrometer squalane and squalene droplets were investigated here by monitoring the heterogeneous reaction, using VUV photoionization aerosol mass spectrometry. While the unsaturated squalane particles appear to be unreactive toward I atoms under our experimental conditions, the saturated squalene particles slowly react with I atoms and iodinated hydrocarbon molecules are formed and detected. Despite the fact that similar gas-phase reactions are expected to have extremely low probability, the reaction of I atoms with squalene particles takes on a probability of $10^{-4}$ due to the trapping and consequently the multiple interactions of the I atoms into the squalene aerosol particles. The effective reactive uptake coefficient and the distribution of the iodinated products are measured. Within experimental error, the chemical evolution of squalene and its iodinated products can all be represented by a single kinetic rate parameter. The effective uptake coefficient measured for unsaturated hydrocarbon squalene is found to be about 3 orders of magnitude larger than the one estimated for saturated hydrocarbon squalane.

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The authors declare no competing financial interest.

ACKNOWLEDGMENTS

This work was supported by the Director, Office of Energy Research, Office of Basic Energy Sciences, Chemical Sciences Division of the U.S. Department of Energy under Contract DE-AC02-05CH11231. The authors acknowledge constructive discussions with Dr. M. D. Ward and the technical assistance from Dr. D. J. Taube. In particular, D.M.P.-V. is grateful to the Alexander von Humboldt Foundation for a Feodor Lynen fellowship.

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