Role of ozone in SOA formation from alkane photooxidation

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Abstract. Long-chain alkanes, which can be categorized as intermediate volatility organic compounds, are an important source of secondary organic aerosol (SOA). Mechanisms for the gas-phase OH-initiated oxidation of long-chain alkanes have been well documented; particle-phase chemistry, however, has received less attention. The δ-hydroxycarbonyl, which is generated from the isomerization of alkoxy radicals, can undergo heterogeneous cyclization and dehydration to form substituted dihydrofuran. Due to the presence of C=C bonds, the substituted dihydrofuran is predicted to be highly reactive with OH, and even more so with O3 and NO3, thereby opening a reaction pathway that is not usually accessible to alkanes. This work focuses on the role of substituted dihydrofuran formation and its subsequent reaction with OH, and more importantly ozone, in SOA formation from the photooxidation of long-chain alkanes. Experiments were carried out in the Caltech Environmental Chamber using dodecane as a representative alkane to investigate the difference in aerosol composition generated from “OH-oxidation-dominating” vs. “ozonolysis-dominating” environments. A detailed mechanism incorporating the specific gas-phase photochemistry, together with the heterogeneous formation of substituted dihydrofuran and its subsequent gas-phase OH/O3 oxidation, is used to evaluate the importance of this reaction channel in dodecane SOA formation. We conclude that (1) the formation of δ-hydroxycarbonyl and its subsequent heterogeneous conversion to substituted dihydrofuran is significant in the presence of NOx; (2) the ozonolysis of substituted dihydrofuran dominates over the OH-initiated oxidation under conditions prevalent in urban and rural air; and (3) a spectrum of highly oxygenated products with carboxylic acid, ester, and ether functional groups are produced from the substituted dihydrofuran chemistry, thereby affecting the average oxidation state of the SOA.

1 Introduction

Alkanes are important constituents of gasoline and vehicle emissions (Hoekman, 1992; Zielinska et al., 1996; Kirchstetter et al., 1999; Gentner et al., 2012; Jathar et al., 2013), accounting for ∼50 % of volatile organic compounds (VOCs) in the urban atmosphere (Fraser et al., 1997; Schauer et al., 1999 and 2002). The unresolved complex mixture (UCM) of organics, which is potentially a significant source of secondary organic aerosol (SOA) formation in the atmosphere, has recently been shown to comprise many long-chain alkanes (Isaacman et al., 2012), which are expected to contribute to SOA formation (Robinson et al., 2007).

Laboratory chamber investigations of SOA formation from long-chain alkanes (Lim and Ziemann, 2005, 2009a, b, c; Presto et al., 2009, 2010; Miracolo et al., 2010, 2011; Craven et al., 2012; Lambe et al., 2012; Tkacik et al., 2012; Yee et al., 2012; Loza et al., 2013) provide a framework for understanding chemical mechanisms and determination of SOA yields (Jordan et al., 2008; Aumont et al., 2012 and 2013; Cappa et al., 2013; Zhang and Seinfeld, 2013). Particle-phase products from OH oxidation of alkanes contain a number of functional groups: organonitrate (–ONO2), hydroxyl (–OH), carbonyl (–C=O), ester (–C(O)O–), and hydroxyperoxide (–OOH). Ambient measurements of organic aerosol composition have shown, in addition, that the carboxylic acid functional group (–C(O)OH) is closely associated with products from fossil fuel combustion sources (Liu et al., 2011; Russell et al., 2011), of which alkanes are a principal component.

Atmospheric alkanes react in daytime exclusively with OH, producing an array of peroxy radicals (RO2). In the presence of sufficient NO, the alkoxy radical (RO) is the key product of the subsequent RO2 reaction with NO: RONO2...
is also produced, with a branching ratio of 0.1–0.3 (Arey et al., 2001). For alkanes with carbon number ≥ 5, the 1,5-H shift isomerization is the dominant reaction pathway for RO, producing a δ-hydroxy carbonyl, the primary fate of which is reaction with OH (Jenkins et al., 2003; Saunders et al., 2003; Bloss et al., 2005), with a lifetime of ~11.5 h at room temperature and a typical ambient OH concentration of \(1 \times 10^6\) molecules cm\(^{-3}\). One particular δ-hydroxy carbonyl, 5-hydroxy-2-pentanone, was found to cyclize to form the cyclic hemiacetal. The cyclic hemiacetal can subsequently lose water to form 4,5-dihydro-2-methylfuran, with an overall lifetime as short as ~1.1 h at 298 K (Cavalli et al., 2000; Martin et al., 2002). The presence of water vapor can, in principle, serve to convert the 4,5-dihydro-2-methylfuran back to 5-hydroxy-2-pentanone, leading to an equilibrium between these two species within several hours (Martin et al., 2002; Baker et al., 2005; Holt et al., 2005; Reisen et al., 2005):

![Diagram of chemical reaction]

This conversion to 4,5-dihydro-2-methylfuran is not unique to 5-hydroxy-2-pentanone and also occurs for other C\(_5\)-C\(_7\)δ-hydroxy carbonyls (Limm and Ziemann, 2005, 2009a, b, c) at a rate that increases with the length of the carbon chain (Holt et al., 2005; Ziemann and Atkinson, 2012). The substituted dihydrofuran is highly reactive towards OH, O\(_3\), and NO\(_3\) in the gas phase, owing to the presence of a C=C double bond. For typical ambient concentrations of OH, O\(_3\), and NO\(_3\), i.e., 2 × 10\(^6\) molecules cm\(^{-3}\) (12 h average), 30 ppb (24 h average), and 5 × 10\(^8\) molecules cm\(^{-3}\) (12 h average), respectively, the lifetimes of the substituted dihydrofuran with respect to reactions with these oxidants in the gas phase are 1.3 h, 7 min, and 24 s, respectively (Martin et al., 2002; Ziemann and Atkinson, 2012). Based on these estimates, ozonolysis of substituted dihydrofurans may dominate OH oxidation during daytime in the urban atmosphere.

SOA formation from long-chain alkanes involves multiple generations of OH oxidation that include functionalization (O-atom addition in forms of a variety of moieties) and fragmentation of the parent carbon backbone. These two routes can eventually lead to highly oxygenated fragments that partition into the particle phase. Two recent modeling studies of C\(_{12}\) alkanes suggest that particle-phase chemistry might play a potentially important role in the chemical composition of alkane SOA; this is inferred from the fact that simulations driven solely by gas-phase chemistry can successfully reproduce the particle-measured SOA yield, but these fail to replicate the observed particulate O : C and H : C ratios in the absence of a particle-phase chemistry channel (Cappa et al., 2013; Zhang and Seinfeld, 2013). The extent to which particle-phase chemistry is important in alkane SOA formation has not been clearly established.

We address here the heterogeneous formation of substituted dihydrofurans and their subsequent gas-phase reaction with ozone in SOA formation from alkanes. We present the results of a series of chamber dodecane photooxidation experiments under two reaction regimes, i.e., “OH-dominant”, in which over 70% of substituted dihydrofurans are oxidized by OH, vs. “O\(_3\)-dominant”, in which 80–90% of substituted dihydrofurans react with O\(_3\). Gas-phase products that are unique to the substituted dihydrofuran chemistry are identified. The impact of ozonolysis of substituted dihydrofurans on the particle-phase product distribution from the photooxidation of dodecane is investigated by comparing intensities of certain ions that are indicative of ozonolysis chemistry. We also develop a detailed mechanism with the incorporation of substituted dihydrofuran chemistry and simulate the effect of this reaction channel on SOA yield from the photooxidation of dodecane.

2 Experimental

Experiments were conducted in the new Caltech dual 24-m\(^3\) Environmental Chamber, in which the temperature (\(T\)) and relative humidity (RH) are automatically controlled. Prior to each experiment, the Teflon chambers were flushed with clean, dry air for 24 h until the particle number concentration was < 10 cm\(^{-3}\) and volume concentration <0.01 µm\(^3\) cm\(^{-3}\). Seed aerosol was injected into the chamber by atomizing 0.015 M aqueous ammonium sulfate solution to provide sufficient surface area for the partition of semivolatile products. Hydrogen peroxide (H\(_2\)O\(_2\)) was used for the OH source by evaporating 85 and 226 µL of 50 wt % aqueous solution into the chamber with 5 L min\(^{-1}\) of purified air for ~110 min, resulting in an approximate starting H\(_2\)O\(_2\) concentration of 1.5 and 4 ppm, respectively, under high- and low-NO\(_x\) conditions. The 4 ppm H\(_2\)O\(_2\) concentration employed in low-NO\(_x\) experiments creates a RO\(_2\)+HO\(_2\) dominant reaction regime. We replaced H\(_2\)O\(_2\) with nitrous acid (HONO) as the OH source for one experiment in order to minimize the formation of ozone. HONO was prepared by dropwise addition of 15 mL of 1 wt % NaNO\(_2\) into 30 mL of 10 wt % H\(_2\)SO\(_4\) in a glass bulb and introduced into the chambers with 5 L min\(^{-1}\) of purified air for ~40 min. To minimize the vapor-phase wall loss along the injection line, 60 µL of dodecane (Sigma-Aldrich, 98 % purity) was injected into a glass bulb, which was connected directly to the Teflon chamber via a 1/4 in. inside diameter (i.d.) Swagelock to NPT fitting located on a Teflon plate. Heated 5 L min\(^{-1}\) of purified air flowed through the glass bulb into the chamber for 30 min, introducing ~200 ppb dodecane into the chamber. After ~1 h of mixing, photooxidation was initiated by irradiating the chamber with black lights with output wavelength ranging from 300 to 400 nm.

Experiments were carried out under conditions in which the peroxy radicals formed from the initial OH reaction with dodecane react either essentially exclusively with NO (so-called high NO\(_x\)) (Exp. 1, 2, 3, 4, and 5) or essentially...
exclusively with HO2 (so-called low NOx) (Exp. 6, 7, and 8). For each condition, O3-dominant vs. OH-dominant environments were generated by varying the OH source, initial NO, NO2, and O3 concentrations, as well as the additional NO injection rate during the irradiation period (Table 1). The ozonolysis vs. OH oxidation of substituted dihydrofuran is calculated to be equally competitive when the ratio of O3 to HO concentration is 6.3 × 10^4 : 1 (approximately 2.6 ppb O3 vs. 1.0 × 10^6 molecules cm^-3 OH). Under low-NOx conditions, NO, NO2, NOx, and O3 were below detection limits, and no ozone formation was observed over the course of 20 h experiments. OH oxidation of dihydrofuran is dominant with average OH concentrations of 1.8 × 10^6 and 1.5 × 10^6 molecules cm^-3, respectively, in Exp. 6 and 8. OH concentration is calculated by optimal fitting of the exponential decay of the GC-FID (gas chromatograph with flame ionization detection)-measured dodecane temporal profile. In order to evaluate the effect of the ozonolysis of dihydrofuran chemistry on the dodecane SOA formation under low-NOx conditions, additional ~30 ppb of ozone was injected into the chamber before the onset of irradiation (Exp. 7). Under high-NOx conditions, HONO was used as the OH source for the OH-dominant environment (Exp. 1). Double HONO injection was carried out in order to fully consume dodecane (before the onset and after ~3 h of irradiation). The average OH concentration is ~5.0 × 10^6 molecules cm^-3, whereas O3 peaks at 8 ppb after 60 min of photooxidation and rapidly decays to ~0 ppb within 2 h. In this case, >73% of dihydrofuran is oxidized by OH over the course of Exp. 1. An O3-dominant environment was generated by injecting NO (38–287 ppb) and NO2 (12–156 ppb) before the onset of irradiation. Continuous NO injection with a certain flow rate (20–100 ppb h^-1) was conducted over the course of experiments to reach a sufficient amount of O3 and to maintain high-NOx levels. Under experimental conditions in Exp. 2, 3, 4, and 5, the average OH concentrations are calculated to be 1.7 × 10^6, 2.0 × 10^6, 1.4 × 10^6, and 1.3 × 10^6 molecules cm^-3, respectively, and the maximum O3 mixing ratios are 20, 380, 150, and 250 ppb, respectively. As a result, ~82, ~96, ~97, and ~98% of dihydrofuran reacts with O3 when dihydrofuran peaks after ~3 h of photooxidation. High-humidity experiments were also carried out as a set of control experiments (Exp. 4, 5, and 8) addressing the role of water vapor in the heterogeneous interconversion between δ-hydroxycarbonyls and substituted dihydrofurans.

A suite of instruments was used to investigate gas- and particle-phase chemistry. T, RH, NO, NOx, and O3 were continuously monitored. Dodecane concentration was monitored by taking hourly samples at ~0.13 L min^-1 of chamber air for 3 min onto a Tenax adsorbent, which was loaded into the inlet of a GC-FID (Agilent 6890N), desorbed at 275°C for 13 min, and then injected onto an HP-5 column (15 m × 0.53 mm i.d. × 1.5 µm thickness, Hewlett-Packard) held at 30°C. The oven was ramped from 30 to 275°C at 10°C min^-1 and held at 275°C for 5 min. The retention time for dodecane is ~27.5 min. The gas-phase species were monitored using a custom-modified Varian 1200 triple-quadrupole chemical ionization mass spectrometer (CIMS) (Crounse et al., 2006; Paulot et al., 2009). In negative mode operation, CF3O^- was used as the reagent ion to cluster with an analyte such as hydroperoxide or acid [R], producing [R-CF3O^-] or m/z [M + 85]^-, where M is the molecular mass.

Table 1. Experimental conditions for the photooxidation of dodecane.

<table>
<thead>
<tr>
<th>Exp.</th>
<th>T0 (K)</th>
<th>RH0 (%)</th>
<th>HC0 (ppb)</th>
<th>NOx0 (ppb)</th>
<th>NOx20 (ppb)</th>
<th>O30 (ppb)</th>
<th>Initial seed vol. (µm^3 m^-3)</th>
<th>Additional NO inj. (ppb h^-1) × (h)</th>
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<td>High NOx</td>
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<td>1</td>
<td>~300</td>
<td>~3</td>
<td>208</td>
<td>430</td>
<td>576</td>
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<td>2</td>
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<td>69</td>
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<td>~2</td>
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<td>Low NOx</td>
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<td>~55</td>
<td>216</td>
<td>&lt; DL</td>
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<td>&lt; DL</td>
<td>58</td>
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</table>

a Under high-NOx conditions, the simulated NO concentration (>5 ppb) is at least 4 orders of magnitude higher than RO2 (<5 × 10^7 molecules cm^-3). Over 99% of RO2 is predicted to react with NO.

b Under low-NOx conditions, the simulated HO2 concentration (~1 × 10^10 molecules cm^-3) is ~20 times higher than RO2 (<5 × 10^6 molecules cm^-3). RO2 + HO2 dominates the fate of RO2.
c NO source was controlled at 100 ppb h^-1 for the first 1 h of reaction and then 25 ppb h^-1 for the remainder of the reaction.
d NO source was controlled at 25 ppb h^-1 over the course of the experiment.
e NO source was controlled at 30 ppb h^-1 for the first 3 h of reaction and then 100 ppb h^-1 for the next 1 h of reaction and then back to 30 ppb h^-1 for the remainder of the reaction.
f NO source was controlled at 50 ppb h^-1 over the course of the experiment.
g Detection limits (DL) for O3, NO, and NO2 are 0.5 ppb, 0.4 ppb, and 0.4 ppb, respectively. H2O2 has an interference on the O3 detection, increasing the O3 monitor readout by ~2–3 ppb in the current study.
weight of the analyte. For more strongly acidic species [H-X], the transfer product, [H·X·F]− or m/z [M + 19]−, is formed during ionization. Carboxylic acids tend to have contributions to both the transfer and cluster product, in which case the overall signal of a compound is considered as the sum of the two product signals. In positive-mode operation, an analyte [R] can undergo proton transfer reaction, producing an ion in the form of [R·H]+, and/or react with n positively charged water clusters to form a cluster in the form of [(H2O)n·R·H]+. Positive mode is employed in this study for tracking less polar compounds, such as the substituted dihydrofuran.

Particle size distribution and number concentration were measured by a cylindrical differential mobility analyzer (DMA, TSI Model 3081) coupled to a condensation particle counter (TSI Model 3010). The protocol for applying particle wall loss correction to DMA measured SOA growth data is described in the Supplement. Real-time particle mass spectra were collected continuously by an Aerodyne high-resolution time-of-flight aerosol mass spectrometer (DeCarlo et al., 2006; Canagaratna et al., 2007). The AMS switched...
Fig. 1b. Schematic mechanism for the photooxidation of dodecane under low-NOx conditions. Note that only the first three generations of products are shown here. The boxes indicate compounds and associated reaction pathways incorporated in the model simulation.

once every minute between the high-resolution “W mode” and the lower resolution, higher sensitivity “V mode”. The V mode was utilized for quantification, as the higher \( m/z \) values exhibit a more favorable signal-to-noise ratio. The W mode was used for ion identification and clarification. Detailed AMS data analysis protocols can be found in the Supplement.

3 Chemical mechanism

A photochemical mechanism was used to simulate the gas-phase photochemistry of NOx, HOOx, and O3 corresponding to the experimental conditions. Reactions and corresponding reaction rate constants are listed in Tables S1–S3 in the Supplement. Photolysis rate constants are calculated using the irradiance spectrum measured for the chamber UV lights as well as absorption cross sections and quantum yields from Sander et al. (2011).

The kinetic scheme for the gas-phase OH-initiated oxidation of dodecane and the further OH oxidation of multi-generational products was developed primarily based on the Master Chemical Mechanism v3.2 (MCM v3.2, http://mcm.leeds.ac.uk/MCM/). Products identified in Lim and Ziemann (2005; 2009 a, b, c) that are not in MCM are
also included here. The kinetic scheme was incorporated in the photochemical model to estimate yields of particle-phase products generated in the chamber. A simplified flow chart illustrating the mechanism for the multigeneration gas-phase chemistry is shown in Fig. 1a, b. In general, the OH-initiated oxidation of dodecane leads to RO2, the fate of which controls the distribution of further generation products. When sufficient NOx is present (the concentration of NO, i.e., > 5 ppb, is at least 4 orders of magnitude higher than RO2, i.e., < 5 × 10^3 molecules cm^-3), over 99% of RO2 reacts with NO, leading to RO or alkyl nitrate (RONO2). The branching ratios and vapor pressures of the production of RO and RONO2 are obtained from Jordan et al. (2008). RO can isomerize through a 1,5-H-atom shift to a δ-hydroxyalkyl radical, react with O2, or undergo fragmentation. The alkyl nitrate formed either undergoes photolysis or reacts with OH via H-atom abstraction from a C atom or reacts with OH via H-atom abstraction from a C atom with a –ONO2 group attached to produce a –C=O group. The δ-hydroxyalkyl radical reacts with O2 and then NO and undergoes another isomerization to produce a δ-hydroxyacetyl. At sufficiently low NOx concentrations, the simulated HO2 concentration (≈1 × 10^10 molecules cm^-3) is ≈20 times higher than RO2 (≈5 × 10^8 molecules cm^-3). RO2 + HO2 dominates the fate of RO2, producing a hydroperoxide (ROOH). Further oxidation of ROOH involves the photolysis of the –OOH group, H-atom abstraction, and the OH oxidation of a C atom with a –OOH group attached to produce a –C=O group. Products through eight generations of oxidation are included in the mechanism, although only the formation of the first three generations of products is illustrated in Fig. 1. The reaction rate constants are obtained from MCM v3.2. In the absence of specific data, the photolysis rate constants of the –OOH, –C=O, and –ONO2 groups on the carbon backbone are assumed to be the same as those for methyl peroxide (CH3OOH), 2-butanal (C3H7CHO), and n-propyl nitrate (n-CH3O2NO2).

Cyclization and subsequent dehydration of δ-hydroxyacetyl to substituted dihydrofuran is a heterogeneous process, including (1) gas-phase diffusion and reactive uptake of δ-hydroxyacetyl to particles, (2) cyclization of δ-hydroxyacetyl to cyclic hemiacetal, and (3) dehydration of cyclic hemiacetal to substituted dihydrofuran. (Atkinson et al., 2008; Lim and Ziemann, 2009c). The extent of this heterogeneous process occurring on/in particles has been predicted to predominate over chamber walls (Lim and Ziemann, 2009 c). In this study, we represent the individual steps of the conversion of δ-hydroxyacetyl to substituted dihydrofuran by an overall first-order decay rate of 3 × 10^-3 s^-1. This value is estimated by monitoring the decay of 5-hydroxy-2-pentanone using CIMS in the presence of 20 μm^3 (NH4)2SO4/H2SO4 seeds at 3% RH. This assumed decay rate is consistent with those rates measured in previous studies (Cavalli et al., 2000; Martin et al., 2002, Holt et al., 2005; Lim and Ziemann, 2009 c). The equilibrium constant K (K = [4,5-dihydro-2-methylfuran]/[5-hydroxy-2-pentanone]) is estimated to be ≈7 at 3% RH from the CIMS measured 5-hydroxy-2-pentanone decay curve, based on the assumption that the decrease in the 5-hydroxy-2-pentanone concentration is accompanied by stoichiometric formation of the corresponding 4,5-dihydro-2-methylfuran. Note that although the proportion of the heterogeneous conversion occurring on the chamber walls is unknown, the potential contribution of chamber walls to the particle-phase production of substituted dihydrofuran has been accounted for by employing the measured overall conversion rate in the mechanism.

The substituted dihydrofuran formed evaporates rapidly due to its high volatility and undergoes reactions with OH, O3, and NO3 in the gas phase (Ziemann and Atkinson, 2012), with reaction rate constants of 2.18 × 10^-10, 3.49 × 10^-15 cm^3 molecule^-1 s^-1, and 1.68 × 10^-13 cm^3 molecule^-1 s^-1, respectively (Martin et al., 2002; Atkinson et al., 2008). Reaction with NO3 is not important under the conditions of this study. In general, the OH addition to an alkyl-substituted dihydrofuran produces either an alkyl-substituted tetrahydrofuran or a carbonyl ester (see Fig. 2a) (Martin et al., 2002, Lim and Ziemann, 2005, 2009a, b, c, Jordan et al., 2008). The mechanism for the O3 reaction with an alkyl-substituted dihydrofuran, as shown in Fig. 2b, was developed following the ozonolysis of 4,5-dihydro-2-methylfuran (Martin et al., 2002), alkyl vinyl ethers (Thiault et al., 2002; Klotz et al., 2004; Sadezky et al., 2006) and monoterpenes (Jenkin et al., 2000; Jenkins, 2004). The reaction of alkyl-substituted dihydrofuran with O3 involves the addition of O3 to the C=O double bond to produce an energy-rich primary ozonide, which rapidly decomposes into two excited Criegee intermediates. The energy-rich Criegee intermediates are either collisionally stabilized or decompose to yield OH (or OH+CO) and an additional α-carbonyl peroxy radical (or peroxy radical). The resulting α-carbonyl peroxy radical (or peroxy radical) can undergo the well-established reactions available for peroxy radicals; see Fig. 2b. The stabilized Criegee intermediates are predicted to react primarily with water (Martin et al., 2002), leading to 3-propoxy-nononal (C12H22O3) and 3-propoxy-nononic acid (C12H22O3), with molar yields of 90 and 10%, respectively. The total yield of 3-propoxy-nononal is predicted to be 18%, which is close to the yield of succinaldehydeic acid methyl ester (23%) from the ozonolysis of 4,5-dihydro-2-methylfuran (Martin et al., 2002).

We consider SOA formation by dodecane photooxidation, including heterogeneous formation of substituted dihydrofuran and its subsequent reaction with OH/O3. Gas–particle equilibrium partitioning of semivolatile products is assumed. The branching ratio and vapor pressure (P^i atm^-1) at 300 K (predicted by SIMPOL.1; Pankow and Asher, 2008) of each product from the OH/O3-initiated oxidation of alkyl-substituted dihydrofuran are labeled in Fig. 2. The corresponding effective saturation concentrations...
(C^n = 10^6 \rho_{L_i} \gamma_i \tilde{M}_w / RT) range from 7.0 × 10^1 \mu g \cdot m^{-3} to 4.7 × 10^4 \mu g \cdot m^{-3}, for which approximately 0.1 % ~ 41.7 % of these products are in the particle phase at an organic loading of \( \sim 50 \mu g \cdot m^{-3} \). Compounds with the lowest volatility (\( \sim 10^{-6} \) atm) are produced mostly from the \( \text{RO}_2 + \text{NO} \rightarrow \text{RONO}_2 \) reaction, the branching ratio of which ranges from 0.11 to 0.28. As a result, the total amount of organic nitrates in the particle phase is relatively high; see Fig. 9c. Compounds generated from the \( \text{RO}_2 + \text{NO} \rightarrow \text{RO} + \text{NO}_2 \) reaction have higher molar yields (0.72–0.89). But they are too volatile (\( \sim 10^{-6}–10^{-7} \) atm) to partition significantly into the particle phase. The stabilized Criegee intermediate reaction with water is predicted to dominate over reaction with NO/NO\(_2\) at RH > 3 %. The ester containing carboxylic acid (\( \text{C}_{12}\text{H}_{22}\text{O}_4 \)) is predicted to be present in the particle phase due to its sufficiently low volatility (\( \sim 10^{-8} \) atm); see Fig. 9c.

### 4 Results and discussion

In this section, we seek to evaluate the impact of substituted dihydrofuran chemistry on dodecane SOA formation as follows: (1) predict the yield of alkyl-substituted dihydrofuran from the photooxidation of dodecane under both high- and low-NO\(_x\) conditions, and the fraction of alkyl-substituted dihydrofuran that reacts with O\(_3\) in the O\(_3\)-dominant regime (Sect. 4.1); (2) measure the time-dependent evolution of cyclic hemiacetal and alkyl-substituted dihydrofuran at 3 %, 10, 20, and 50 % RH (Sect. 4.2); (3) propose gas-phase products that are unique to dihydrofuran chemistry based on the CIMS speciation (Section 4.3); (4) compare particle-phase chemical composition under OH-dominant vs. O\(_3\)-dominant environments via identifying AMS measured ions representative of dihydrofuran oxidation products (Sect. 4.4); and (5) estimate the change in SOA yield and elemental composition by incorporating the complete substituted-dihydrofuran formation and removal pathways into the dodecane SOA prediction model (Sect. 4.5).

#### 4.1 Predicted substituted dihydrofuran formation and reaction with ozone in the dodecane system

To what extent is the formation of substituted dihydrofuran and its subsequent chemistry important in the formation of dodecane SOA? A mechanism simulation was conducted, with initial conditions similar to those employed in the chamber (Table 1), i.e., 200 ppb dodecane, 1.5 ppm H\(_2\)O\(_2\), 100 ppb O\(_3\), and 100 ppb + 30 ppb h\(^{-1}\) NO for high-NO\(_x\) conditions and 200 ppb dodecane, 4 ppm H\(_2\)O\(_2\), 100 ppb O\(_3\), and 0 ppb NO\(_x\) for low-NO\(_x\) conditions. Figure 3 shows the predicted mass distribution of \( \delta \)-hydroxycarbonyl, alkyl-substituted dihydrofuran, together with other products of the same generation. \( \delta \)-hydroxycarbonyl is predicted to account for > 90 % total organic mass of first-generation products under high-NO\(_x\) conditions (see Fig. 3a (H)), but < 5 % to the total organic mass of the third-generation products under low-NO\(_x\) conditions (see Fig. 3a (L)).
heterogeneous conversion pathway from $\delta$-hydroxycarbonyl to alkyl-substituted dihydrofuran is predicted to result in a rapid consumption of $\delta$-hydroxycarbonyl under dry conditions, as opposed to a slow decay from OH oxidation. Alkyl-substituted dihydrofuran accounts for up to 70% of the decay of $\delta$-hydroxycarbonyl, as shown in Fig. 3b (H). As discussed earlier, an ozonolysis-dominant environment occurs for O$_3$ > 3 ppb at a typical OH concentration of $1.0 \times 10^6$ molecules cm$^{-3}$. In the high-NO$_x$ simulation case, ozone is predicted to lead to 82–98% of the total alkyl-substituted dihydrofuran loss over the course of photooxidation; see Fig. 3d (H).

The alkyl-substituted dihydrofuran is not the only product that contains a dihydrofuran structure in the dodecane photooxidation system. The first-generation product 3-dodecyl nitrate under high-NO$_x$ conditions is an example. Three pathways exist that could produce semivolatile compounds containing a dihydrofuran structure from the further photochemical reaction of 3-dodecyl nitrate: (1) photolysis of the –ONO$_2$ group leads to an alkyl-substituted dihydrofuran, (2) abstraction of an H atom by OH potentially leads to an...

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Fig. 2b. Proposed mechanism for the ozonolysis of alkyl-substituted dihydrofuran under high-NO$_x$ conditions. The boxes indicate compounds or reaction pathways incorporated in the model simulation. CIMS monitored species have m/z noted in red. Estimated vapor pressure (atm) of each compound is indicated in blue.
organonitrate-substituted dihydrofuran, and (3) H-atom abstraction from a C atom with the –ONO₂ functional group attached and further H-atom abstraction from a C atom potentially leads to a carbonyl-substituted dihydrofuran. It is worth noting that the gas-phase ozonolysis of alkyl-substituted dihydrofuran under high-NOₓ conditions contributes to most of the ozonolysis reactions because the alkyl-substituted dihydrofuran is a major first-generation product, as shown in Fig. 3b (H).

4.2 Formation of cyclic hemiacetals and alkyl-substituted dihydrofuran

CIMS measurement at (+)m/z 183 in positive mode represents the ion C₁₂H₂₂O·H⁺ generated from the proton transfer reaction with alkyl-substituted dihydrofuran (M_w = 182). An additional source of (+)m/z 183 is the proton transfer reaction followed by dehydration of hydroxyl dodecanone (M_w = 200), which is also detected as a fluoride cluster product at (-)m/z 285 (C₁₂H₂₄O₂·CF₃O⁻) in negative mode.
mode. Figure 4 shows temporal profiles of (+)m/z 183 and (−)m/z 285 monitored under both high- and low-NOx conditions (Exp. 2 vs. Exp. 6). The (+)m/z 183 ion was detected in each of the two experiments, but with distinct time-dependent patterns. Under high-NOx conditions, (+)m/z 183 peaks during the first 3 h of irradiation, indicating fast formation of alkyl-substituted dihydrofuran, and decays during the next 15 h of photooxidation. Under low-NOx conditions, however, the (+)m/z 183 signal eventually reaches a plateau, which is more consistent with the pattern of (−)m/z 285. This behavior is consistent with the mechanism prediction that the alkyl-substituted dihydrofuran is formed in significant amounts only under high-NOx conditions (Fig. 3b (H)). An alternative explanation for the temporal profiles of (+) m/z 183 under low-NOx conditions is that this ion is formed by dehydration of the cyclic hemiacetal following protonation in CIMS. The occurrence of a plateau could be indicative of an absence of dehydration in the particle phase due to the lack of an acid catalyst (no HNO3 formation from NO2+OH reaction as in the high-NOx cases). However, based on the model prediction, cyclic hemiacetal is not likely to accumulate since the formation of its precursor, δ-hydroxycarbonyl, is a minor pathway when the RO2+HO2 reaction is dominant and the photolysis of the resulting peroxide is not important.

The AMS measured m/z 183 (C12H23O+) is the major characteristic ion for cyclic hemiacetal. This ion is produced by the neutral loss of OH (Mw = 17) from the 2 protonation in the cyclic hemiacetal (Mw = 200) during electron ionization (Gong et al., 2005; Lim and Ziemann, 2009c). In addition, the C12H23O+ ion is suggested to be the characteristic fragment of carbonyl-hydroperoxide-derived peroxyhemiacetal (Yee et al., 2012). The temporal profiles of the C12H23O+ ion under both high- and low-NOx conditions (Exp. 2 vs. Exp. 6) exhibit distinct growth patterns (Fig. 4). Under high-NOx conditions, the C12H23O+ ion signal increases rapidly to a maximum during the first 2 h and decays over the next ~14 h. The temporal behavior of the C12H23O+ ion is a result of its rapid formation, i.e., uptake of δ-hydroxycarbonyl onto particles and subsequent cyclization, and relatively slower removal, i.e., dehydration. Under low-NOx conditions, the C12H23O+ ion increases over the course of a 20 h experiment because of the accumulative formation of peroxyhemiacetal. During the first 3 h of irradiation under low-NOx conditions, the organic loading is below the AMS detection limit; thus the C12H23O+ ion signal does not appear in Fig. 4 during this period. The C12H23O+ ion signal under low-NOx conditions therefore potentially represents peroxyhemiacetal, since the formation of peroxides is the major reaction pathway in the RO2+HO2 dominant regime. This is again consistent with the mechanism prediction that the formation of alkyl-substituted dihydrofuran is unimportant under low-NOx conditions (Fig. 3b (L)).

Figure 5 shows temporal profiles of the ion C12H22O⋅H+ detected by CIMS and the ion C12H23O+ detected by AMS.
in the presence of NO<sub>x</sub> under varying RH. In general, the AMS ion C<sub>12</sub>H<sub>22</sub>O<sup>+</sup> peaks ~3 h earlier than the CIMS ion C<sub>12</sub>H<sub>22</sub>O·H<sup>+</sup>. The decay rates of AMS ion C<sub>12</sub>H<sub>23</sub>O<sup>+</sup>, which is calculated by assuming first-order kinetics, are 4.02 × 10<sup>−3</sup>, 3.06 × 10<sup>−3</sup>, 1.71 × 10<sup>−3</sup>, and 1.37 × 10<sup>−3</sup> s<sup>−1</sup> at RH of 3, 10, 20, and 50 %, respectively. Aerosol water content could slow down the formation and removal rate of AMS ion C<sub>12</sub>H<sub>23</sub>O<sup>+</sup>, mainly because (1) water could accelerate the hydration rate, thus changing the equilibrium coefficient of the interconversion between cyclic hemiacetal and dihydrofuran, and (2) the addition of water might neutralize the particle-phase acid (e.g., HNO<sub>3</sub>), which is thought to catalyze the cyclization of δ-hydroxycarbonyl to cyclic hemiacetal (Lim and Ziemann, 2009c). The decay rates of AMS ion C<sub>12</sub>H<sub>23</sub>O<sup>+</sup> are within the same order of magnitude of that measured for 5-hydroxy-2-pentanone at 3 % RH, indicating that the dehydration of cyclic hemiacetal is the rate-limiting step in the overall heterogeneous conversion process. Water vapor influences CIMS sensitivity to certain compounds in negative mode by clustering with the reagent ion CF<sub>3</sub>O<sup>−</sup> to form [H<sub>2</sub>O·CF<sub>3</sub>O]<sup>−</sup>. The decreasing intensity of the ion C<sub>12</sub>H<sub>22</sub>O·H<sup>+</sup> as RH increases is likely indicative of this effect.

4.3 Products from alkyl-substituted dihydrofuran oxidation

Products unique to the alkyl-substituted dihydrofuran oxidation chemistry have been proposed based on CIMS measured m/z signals; see Table 2 for chemical structures. Authentic standards are not commercially available for these products. Compounds from the dodecane photooxidation route that share the same m/z signal with alkyl-substituted dihydrofuran oxidation products are also presented in Table 2. The CIMS signal (−)m/z 346 is composed of 2-nitroxy-2-alkyl-3-hydroxyl-5-heptyl-tetrahydrofuran and 2-carbonyl-5-hydroxyl-2-dodecyl-nitrate. The latter is a third-generation product resulting from the further oxidation of dodecyl nitrate. The branching ratio for the addition of an −ONO<sub>2</sub> group is much lower than that for the formation of the RO radical; therefore the interference of 2-carbonyl-5-hydroxyl-2-dodetyl-nitrate in the (−)m/z 346 signal can be neglected. The CIMS signal (−)m/z 299 represents 3-propoxy-nonanal, which is a primary product from both OH oxidation and ozonolysis of alkyl-substituted dihydrofuran, and 8-hydroxy-3,5-dodecanedione, which is produced from the further oxidation of dodecyl nitrate. As discussed earlier,
Table 2. Proposed structures for CIMS ions unique to the alkyl-substituted dihydrofuran chemistry. C and T indicate the cluster and transfer product, respectively. Commercial standards are not available.

<table>
<thead>
<tr>
<th>Observed m/z</th>
<th>Product</th>
<th>Chemical formula</th>
<th>Proposed structure</th>
<th>Chemical pathway</th>
<th>Interference</th>
</tr>
</thead>
<tbody>
<tr>
<td>183 (+)</td>
<td>T</td>
<td>C_{12}H_{22}O</td>
<td><img src="image1" alt="Proposed Structure" /></td>
<td>Heterogeneous conversion</td>
<td><img src="image2" alt="OH" /></td>
</tr>
<tr>
<td>328 (−)</td>
<td>C</td>
<td>C_{12}H_{21}NO_{4}</td>
<td><img src="image3" alt="Proposed Structure" /></td>
<td>Heterogeneous conversion</td>
<td>—</td>
</tr>
<tr>
<td>346 (−)</td>
<td>C</td>
<td>C_{12}H_{23}NO_{5}</td>
<td><img src="image4" alt="Proposed Structure" /></td>
<td>OH-oxidation</td>
<td><img src="image5" alt="OH" /></td>
</tr>
<tr>
<td>299 (−)</td>
<td>C</td>
<td>C_{12}H_{22}O_{3}</td>
<td><img src="image6" alt="Proposed Structure" /></td>
<td>OH-oxidation, Ozonolysis</td>
<td><img src="image7" alt="OH" /></td>
</tr>
<tr>
<td>249 (−)</td>
<td>T</td>
<td>C_{12}H_{22}O_{4}</td>
<td><img src="image8" alt="Proposed Structure" /></td>
<td>Ozonolysis</td>
<td>—</td>
</tr>
<tr>
<td>315 (−)</td>
<td>C</td>
<td>C_{12}H_{22}O_{4}</td>
<td><img src="image9" alt="Proposed Structure" /></td>
<td>Ozonolysis</td>
<td>—</td>
</tr>
<tr>
<td>332 (−)</td>
<td>C</td>
<td>C_{11}H_{21}O_{3}N</td>
<td><img src="image10" alt="Proposed Structure" /></td>
<td>Ozonolysis</td>
<td>—</td>
</tr>
<tr>
<td>348 (−)</td>
<td>C</td>
<td>C_{11}H_{21}O_{3}N</td>
<td><img src="image11" alt="Proposed Structure" /></td>
<td>Ozonolysis</td>
<td>—</td>
</tr>
<tr>
<td>301 (−)</td>
<td>C</td>
<td>C_{11}H_{20}O_{4}</td>
<td><img src="image12" alt="Proposed Structure" /></td>
<td>Ozonolysis</td>
<td>—</td>
</tr>
</tbody>
</table>

the formation and transformation of dodecyl nitrate is predicted to be a minor pathway, and thus the (−)m/z 299 is dominated by 3-propoxy-nonanal.

A distinct feature of proposed products from the dihydrofuran oxidation by either OH or O₃ is the formation of an ester group (−C(O)O−) or an ether group (−O−), both of which cannot be accessed from photochemical reaction pathways initiated by the OH attack on the aliphatic hydrocarbon. The OH oxidation channel leads solely to the formation of tetrahydrofuran and carbonyl ester. In the ozonolysis pathway, on the other hand, the reaction of stabilized Criegee intermediates with water produces one ester group containing aldehyde (3-propoxy-nonanal) and carboxylic acid (3-propoxy-nonanic acid); see proposed structures in Table 2. From the intact C₁₂ skeleton and highly oxygenated nature, these two products are consistent with CIMS signals in
(-)m/z 299 and (-)m/z 249 and 315 in negative mode with little interference from other products.

Figure 6 shows the temporal profiles of (-)m/z 299, as well as (-)m/z 249 and 315, under “O3-limiting” (Exp. 2) and “O3-rich” (Exp. 3) conditions. The decay rates of their precursor (+)m/z 183, as calculated by assuming first-order kinetics, are $1.19 \times 10^{-5}$ s$^{-1}$ and $1.44 \times 10^{-5}$ s$^{-1}$, respectively. Under O3-rich conditions, both species increase and eventually level off. When O3 reaction is less competitive, the CIMS signals start to decrease after 10 h of photooxidation. Note that these three ions were also detected under low-NOx conditions. No significant increase in their signals was observed in the O3-rich environment (Exp. 7), compared with the O3-limiting case (Exp. 6). If they were assigned the same chemical structures as those proposed under high-NOx conditions, their growth pattern would suggest a minor formation of their precursor alkyl-substituted dihydrofuran. This can be attributed to the slowdown or even termination of the dehydration of cyclic hemiacetal in the absence of strong acid catalysis (Atkinson et al., 2008; Lim and Ziemann, 2009c).

### 4.4 Ozone vs. OH oxidation

Carboxylic acids produce significant signals at m/z 60 and 73, specifically C$_2$H$_4$O$_2^+$ and C$_3$H$_5$O$_2^+$, in the AMS measurement (Aiken et al., 2008). The formation of these two ions, as a function of the time-dependent decay of the ion C$_{12}$H$_{23}$O$_7^+$, is shown under different RH conditions in Fig. 7. The largest slope of either $\Delta C_{24}H_{45}O_2^+$/$\Delta C_{12}H_{23}O_7^+$ or $\Delta C_{35}H_{60}O_2^+$/$\Delta C_{12}H_{23}O_7^+$ is associated with the highest RH, i.e., 50%. However, the production of C$_2$H$_4$O$_2^+$ and C$_3$H$_5$O$_2^+$ is not significant under dry conditions. Also, the changes in slopes along with changes in RH values are consistent for both ions. This indicates that the formation of the carboxylic acid functional group detected in particles is associated with the water vapor concentration in the gas phase, consistent with the reaction of the stabilized Criegee intermediate with water in the substituted dihydrofuran oxidation system.

Two dominant oxygen-containing ions, m/z 44 (mostly CO$_2^+$) and m/z 43 (C$_3$H$_7^+$ and C$_2$H$_3$O$^+$), have been widely used to characterize organic aerosol evolution in chamber and field observations. Previous studies have shown that CO$_2^+$ results mostly from the thermal decarboxylation of an organic acid group (Alfarra, 2004). The $f_{44}$ (ratio of m/z 44, mostly CO$_2^+$, to total signal in the component mass spectrum) axis is also considered to be an indicator of photochemical aging (Alfarra et al., 2004; Aiken et al., 2008; Kleinman et al., 2008). It has been found that increasing OH exposure increases $f_{44}$ and decreases $f_{43}$ (ratio of m/z 43, mostly C$_3$H$_7^+$, to total signal in the component mass spectrum) for SOA generated from gas-phase alkanes (Lambe et al., 2011). The C$_3$H$_7$O$_2^+$ ion at m/z 43 is assumed predominantly due to nonacid oxygenates, such as saturated carbonyl groups (Ng et al., 2011). The evolution of dodecane SOA from four experiments (Table 1), characterized by different OH and O$_3$ exposure, and different RH levels, is shown in $f_{CO_2^+} - f_{C_3H_7O^+}$ space in Fig. 8. Overall, high-NOx dodecane SOA lies to the lower left of the triangular region derived for ambient SOA. The relatively high organic loading (~200 ppb dodecane) employed in this study favors partitioning of less oxidized species, which would remain in the gas phase under atmospheric conditions. For each experiment, $f_{CO_2^+}$ decreases and $f_{C_3H_7O^+}$ increases with increasing SOA at the beginning of irradiation. After several hours of photooxidation, the trends reverse, resulting in increasing $f_{CO_2^+}$ and decreasing $f_{C_3H_7O^+}$. Curve in $f_{CO_2^+} - f_{C_3H_7O^+}$ space has been also observed in other chamber/flow reactor studies (Kroll et al., 2009; Ng et al., 2010; Chhabra et al., 2011; Lee et al., 2011; Lambe et al., 2011). When the organic loading is small, only the highly oxygenated and least volatile species partition to the particle phase. As particles grow, more volatile and less oxidized species are able to...
Fig. 8. SOA evolution as a function of OH exposure vs. O₃ exposure from dodecane photochemistry under high-NOₓ conditions in the $f_{44}$ vs. $f_{43}$ space. The top-left graph shows the combination of all the data, and the bottom-left graph shows an expanded version. The other graphs show the specific behavior of each experiment: Exp. 1 corresponds to a regime in which OH oxidation of dihydrofuran is dominant. Exp. 3, 4, and 5 correspond to a regime in which ozonolysis of dihydrofuran is dominant, at 10, 20, and 50 % RH, respectively.

Fig. 9. (A) SOA average carbon oxidation state as a function of OH exposure from dodecane photochemistry under high-NOₓ conditions. Exp. 1 corresponds to a regime in which OH oxidation of dihydrofuran is dominant. Exp. 3, 4, and 5 correspond to a regime in which ozonolysis of dihydrofuran is dominant, 10, 20, and 50 % RH, respectively. (B) van Krevelen diagram. AMS measured O : C vs. H : C ratios for the four experiments under the OH total exposure ranging from $1.0 \times 10^7$ to $2.0 \times 10^7$ molecules cm$^{-3}$ h. Gray dashed lines denote the average carbon oxidation state.

... participate in equilibrium partitioning, leading to a decrease in $f_{CO_2}$. Progressive oxidation of semivolatile products in the gas phase eventually leads to multifunctionalized species contributing to the increase of $f_{CO_2}$.

Both O₃- and OH-initiated oxidation of substituted dihydrofuran lead to the formation of $f_{CO_2}$, via the thermal decarboxylation of an organic acid group and photochemical aging, respectively. The contribution of each reaction path-way to the intensity of $f_{CO_2}$ can be evaluated based on the $f_{CO_2} - f_{C_2H_5O^+}$ plot. As discussed in Sect. 2.1, Exp. 1 is designed as an OH-dominant case, in which it is estimated that > 73 % of the dihydrofuran reacts with OH over the course of the experiment. Exp. 3, 4, and 5 were designed to be O₃-dominant at 10 %, 20 %, and 50 % RH, respectively. Approximately 96, 97, and 98 % of substituted dihydrofuran is predicted to react with O₃ when it peaks after ~3 h of photooxidation in Exp. 3, 4, and 5, respectively.
Fig. 10. Comparison of CIMS measured ions in positive and negative mode, with chemical structures proposed in Table 2, with model simulations using experimental conditions of Exp. 2.
As shown in Fig. 8, the intensities of $f_{CO_2}$ in Exp. 1, 3, 4, and 5 are 0.019, 0.020, 0.028, and 0.030, respectively, at the same OH exposure, i.e., $2 \times 10^7$ molecules cm$^{-3}$ h. The increase in $f_{CO_2}$ intensities results from the increasing O$_3$ and RH levels in these four experiments. Overall, Exp. 1 exhibits the least intensity of $f_{CO_2}$ at the end of the experiment, i.e., 0.017, although the total OH exposure is the highest, i.e., $3.8 \times 10^7$ molecules cm$^{-3}$ h. The total OH exposures for Exp. 3, 4, and 5 are $3.5 \times 10^7$, $2.2 \times 10^7$, and $2.1 \times 10^7$ molecules cm$^{-3}$ h, respectively, which are less than that in Exp. 1. However, the intensities of $f_{CO_2}$ for these three experiments are 35–82\% higher than that in Exp. 1 at the end of experiments. In addition, the intensity of $f_{CO_2}$ increases along with increasing RH and O$_3$ exposure for these three experiments, varying from 0.023 to 0.031. The highest $f_{CO_2}$ intensity shown in Exp. 5 corresponds to the highest O$_3$ exposure and RH level, but lowest OH exposure.

The difference in elemental composition of organic particles produced from O$_3$-dominant vs. OH-dominant environments is also examined via the AMS measured O : C and H : C ratios in the van Krevelen plot and the time-dependent evolution of the average carbon oxidation state (OS$_C = 2 \times O : C - H : C$; see Fig. 9). In general, the OS$_C$ values calculated fall into a region characterized by oxidized primary organic aerosol and semivolatile oxidized organic aerosol (Kroll et al., 2011). The measured O : C (~0.2) and H : C (~1.7) ratios at an OH exposure on the order of $\sim 10^7$ molecules cm$^{-3}$ h agree with those measured for organic aerosols generated from C$_{10}$–C$_{17}$ alkanes (Lambe et al., 2011, 2012). As mentioned earlier, the four experiments (Exp. 1, 3, 4, and 5) were conducted in such a way that the total OH exposure is decreasing, whereas the total O$_3$ and RH exposure is increasing along with increasing experimental numbers (for example, Exp. 5 has the least
OH exposure but the largest O$_3$ and RH exposure). A clear trend observed from the van Krevelen plot is that O$_2$:C increases, whereas H:C decreases under elevated O$_3$ and RH levels. The effect of ozonolysis of substituted dihydrofuran chemistry on the aerosol chemical composition can be evaluated by comparing OS$_C$ (O$_2$:C vs. H:C) for these four experiments under the same OH exposure. Consistent with our findings in $f_{CO_2}$ $– f_{C_2H_5O^+}$ space, the highest OS$_C$ is observed under the highest O$_3$ exposure and RH level, but lowest OH exposure (98% substituted dihydrofuran reacts with O$_3$ at 55% RH). With the same OH exposure, e.g., $1.5 \times 10^7$ molecules cm$^{-3}$ h, the average carbon oxidation state increases from $–1.36$ in O$_3$-limiting environments (Exp. 1) to $–1.25$ in O$_3$-dominant environments (Exp. 5). In conclusion, ozonolysis of substituted dihydrofuran plays an important role in the formation of highly oxidized aerosol in alkane SOA.

4.5 Experiment and model comparison

Figure 10 shows the temporal profiles of CIMS measured ions in (+/−) mode, with structures proposed in Table 2, together with the corresponding model predictions under conditions of Exp. 2. In general, four time-dependent growth patterns are observed in experiments, which are also captured by model predictions. “Pattern 1” denotes species with rapid removal pathways, e.g., $m/z$ 301 (−), with a proposed structure of δ-hydroxycarbonyl. In the current mechanism, the overall heterogeneous conversion rate of δ-hydroxycarbonyl to substituted dihydrofuran is taken as $3 \times 10^{-3}$ s$^{-1}$. The model output is consistent with the observed time-dependent trend when this rate is used. Up to $\sim$90% of $m/z$ 301 (−) is consumed due to this rapid heterogeneous reaction pathway at 3% RH after 18 h of photooxidation. “Pattern 2” is indicative of a species that also reacts rapidly, but with a much slower consumption rate than species of pattern 1. A typical example here is $m/z$ 183 (+), which represents the alkyl-substituted dihydrofuran. The reaction rate constants of alkyl-substituted dihydrofuran with either OH or O$_3$ are at least an order of magnitude higher than the generic reaction rate constant for the OH abstraction reaction, which is the dominant gas-phase pathway in the dodecane photooxidation mechanism. The simulated peak occurs $\sim$2 h earlier than observations, indicating that the formation rate of alkyl-substituted dihydrofuran might be slower than the decay rate of its precursor, δ-hydroxycarbonyl, considering that fact that the acid-catalyzed dehydration process in the particle phase is the rate-limiting step. The extent to which the formation rate is slower than the decay rate, however, is unknown since the measurement of rate constants for individual steps is infeasible in this study. “Pattern 3” reflects the temporal profiles for a majority of ions here, e.g., $m/z$ 328 (−), 299(−), 249(−), 315(−), and 332(−). Compounds proposed for the above $m/z$ can be categorized as semivolatile products. In the gas phase, they undergo functionalization or fragmentation, or partition into the particle phase as the precursors of SOA. Overall, the temporal profiles of species in pattern 3 are governed by the progressive photochemistry in the gas phase and gas–particle equilibrium partitioning. “Pattern 4”, including $m/z$ 346 (−) and 328 (−) here, represents “non-volatile” species. Owing to their low volatilities, they will immediately partition to the particle phase once formed and the gas-phase photooxidation becomes negligible, although it might still occur via the OH attack on C atoms.

Figure 11a shows the simulated SOA growth (SIM.1) using the initial conditions in Exp. 2, together with the observed total organic aerosol mass as a function of reaction time and OH exposure. The model reproduces the chamber measured SOA yield at 3% RH when the conversion rate of $3 \times 10^{-3}$ s$^{-1}$ is employed to represent the heterogeneous conversion of δ-hydroxycarbonyl to dihydrofuran. A second simulation (SIM.2) was run with the complete dihydrofuran chemistry removed while other parameters were held constant. The total organic mass is $\sim$ 42% higher as a result after 18 h of photooxidation. The formation of alkyl-substituted dihydrofuran from δ-hydroxycarbonyl is accompanied by an increase of vapor pressure from 5.36 $\times$ 10$^{-2}$ to 1.08 $\times$ 10$^{-4}$ atm at 300 K, as predicted by SIMPOL.1, and the total organic mass formed decreases. Although the addition of OH to the C=C double bond in the substituted dihydrofuran introduces an extra OH group, the decrease of vapor pressure owing to the addition of one OH group does not compensate for the heterogeneous conversion of both –C=O and –OH groups in δ-hydroxycarbonyl to an –O– group in

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**Fig. 12.** Regimes of dominance of ozonolysis vs. OH oxidation of substituted dihydrofuran. Daily maximum OH concentrations vary by regions in the troposphere, i.e., $10^3$–$10^6$ molecules cm$^{-3}$ for background air, $10^6$–$10^7$ molecules cm$^{-3}$ for rural air, and $>10^7$ molecules cm$^{-3}$ for urban air, with a global average concentration of $1 \times 10^6$ molecules cm$^{-3}$. Daily average surface ozone mixing ratios in rural areas are between 10 and 50 ppb, but these can exceed 100 ppb in polluted urban areas.
a nonaromatic ring in dihydrofuran. The predicted average carbon oxidation state is \( \sim 7-15\% \) higher than observations. The overprediction is within the uncertainties in the O:C (31\%) and H:C (10\%) measurement by AMS (Aiken et al., 2008). Incorporation of the substituted dihydrofuran formation and removal pathways in the model leads to an increase in the simulated O/C. Compared with compounds produced from dodecane photooxidation under high-NO\(_x\) conditions, products from dihydrofuran chemistry tend to have a higher O:C but lower H:C due to the formation of ether, ester, and carboxylic acid functional groups. As a result, the calculated average carbon oxidation state is higher in the presence of chemical reactions that accelerate the aerosol aging process.

5 Atmospheric implications

The importance of ozone in the SOA formation from the photooxidation of long-chain alkanes under atmospherically relevant conditions depends on two factors: (1) the relative concentration of O\(_3\) vs. OH, and (2) the heterogeneous conversion rate of \( \delta \)-hydroxycarbonyls to substituted dihydrofurans. Figure 12 shows regimes of ozonolysis vs. OH oxidation of substituted dihydrofuran corresponding to ranges of OH and O\(_3\) concentrations. The OH-initiated oxidation of substituted dihydrofuran is predicted to dominate only under remote atmospheric conditions. Most alkane emissions occur in areas where ozone levels exceed 10 ppb, where the ozonolysis of dihydrofuran should be dominant.

Under conditions of the current study, the substituted dihydrofuran chemistry is predicted to account for \( > 95\% \) of the removal pathways of \( \delta \)-hydroxycarbonyls to substituted dihydrofurans. Figure 12 shows regimes of ozonolysis vs. OH oxidation of substituted dihydrofuran corresponding to ranges of OH and O\(_3\) concentrations. The OH-initiated oxidation of substituted dihydrofuran is predicted to dominate only under remote atmospheric conditions. Most alkane emissions occur in areas where ozone levels exceed 10 ppb, where the ozonolysis of dihydrofuran should be dominant.

In summary, two impacts of substituted dihydrofuran chemistry on alkane SOA formation are expected. First, the SOA yield from the photooxidation of long-chain alkanes can be overpredicted without accounting for substituted dihydrofuran formation and removal pathways. Second, a substantial amount of carboxylic acid, ester, and tetrahydrofuran moieties can be produced, leading to higher O:C but much lower H:C ratios, and thus a higher oxidation state of alkane SOA in general. In this manner, the dihydrofuran chemistry can be considered as a “dehydration” channel in alkane SOA formation.

Supplementary material related to this article is available online at http://www.atmos-chem-phys.net/14/1733/2014/acp-14-1733-2014-supplement.pdf.

Acknowledgements. This work was supported by National Science Foundation grant AGS-1057183.

Edited by: A. Laskin

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