Application of positive matrix factor analysis in heterogeneous kinetics studies utilizing the mixed-phase relative rates technique

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Abstract:

The mixed-phase relative rate approach for determining aerosol particle organic heterogeneous reaction kinetics is often performed utilizing mass spectral tracers as a proxy for particle phase reactant concentration. However, sometimes this approach may be influenced by signal contamination from oxidation products during the experiment. In the current study, the mixed-phase relative rates technique has been improved by combining a Positive Matrix Factor (PMF) analysis with electron ionization Aerosol Mass Spectrometry (unit mass resolution), thereby removing the influence of m/z fragments from reaction products on the reactant signals. To demonstrate the advantages of this approach, the heterogeneous reaction between OH radicals and citric acid (CA) was investigated using a photochemical flow tube coupled to a compact time-of-flight aerosol mass spectrometer (C-ToF-AMS). The measured heterogeneous rate constant \( k_2 \) of citric acid toward OH was \((3.31 \pm 0.29) \times 10^{-12} \text{cm}^3 \text{molecule}^{-1} \text{s}^{-1}\) at 298 K and (30±3) % RH and was several times greater than the results utilizing individual m/z fragments. This phenomenon was further evaluated for particulate-phase organophosphates (TPhP, TDCPP, and TEHP), leading to \( k_2 \) values significantly larger than previously reported. The results suggest that heterogeneous kinetics can be significantly underestimated when the structure of the products is highly similar to the reactant and when a non-molecular tracer is measured with a unit mass resolution aerosol mass spectrometer. The results also suggest that the heterogeneous lifetime of organic aerosol in models can be overestimated due to underestimated OH uptake coefficients. Finally, a comparison of
reported rate constants implies that the heterogeneous oxidation of aerosols will be
dependent upon a number of factors related to the reaction system, and that a single
rate constant for one system cannot be universally applied under all conditions.
1. Introduction

Reaction kinetics data provide key parameters for both air quality and climate models. They are required to compute the trace gas and particle matter (PM) content of the atmosphere (Kolb et al., 2010) and to evaluate the atmospheric lifetime and fate for individual species. Organic particles make up 10-90% of the global submicron particle mass in the lower troposphere (Zhang et al., 2011), and is comprised of various reactive organic species, which are subject to atmospheric heterogeneous oxidation. Previous studies have found that heterogeneous reactions with OH in particular, can lead to an increase in density, CCN activation (George and Abbatt, 2010) and optical extinction (Cappa et al., 2011) of organic particulate matter. Therefore, there is a growing interest in understanding not only the mechanism of PM transformation through heterogeneous reactions including oxidation, but also determining the rates at which organic aerosols are chemically transformed in the atmosphere.

To this end, Donahue et al. (Donahue et al., 2005) and Hearn and Smith (Hearn and Smith, 2006) developed a mixed-phase relative rate technique for measuring organic PM component heterogeneous reaction kinetic rate constants. In this method, the rate constant of the compound of interest is determined from the decrease of its particle phase relative concentration as a function of oxidant exposure. The oxidant levels are simultaneously estimated via the measured loss of a gas phase reference compound after applying the known second-order rate constant \( k_2 \) toward the oxidant. In this approach the rates of chemical change are given by,
\[
\frac{dc_A}{dt} = k_{2,A} c_A c_{Ox} \quad (1)
\]
\[
\frac{dc_R}{dt} = k_{2,R} c_R c_{Ox} \quad (2)
\]
where \( c_A \), \( c_R \) and \( c_{Ox} \) are the particle phase concentration of the compound of interest \((A)\), the gas phase concentration of the reference compound \((R)\) and oxidant \((\text{molecules cm}^{-3})\) respectively, while \( k_{2,A} \) and \( k_{2,R} \) are the second-order rate constant of \( A \) and \( R \) to the oxidant \((\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})\). A relative rate constant \( (k_r) \) \((\text{ie: particle phase reaction rate of } A, \text{relative to the gas phase reaction rate of } R)\) can be derived by dividing Eq. (1) by Eq. (2). The derivation of \( k_r \) provides a means to obtain heterogeneous kinetic data without the need to know the absolute concentration of the oxidant. The differential and integral forms for the relative rates technique are shown as Eq. (3) and (4),
\[
\frac{dc_A}{c_A} = \frac{k_{2,A}}{k_{2,R}} \frac{dc_R}{c_R} = k_r \frac{dc_R}{c_R} \quad (3)
\]
\[
\log \frac{c_A}{c_{A,0}} = k_r \log \frac{c_R}{c_{R,0}} \quad (4)
\]
from which the relative rate constant \( (k_r) \), is the slope of the line derived by plotting the logarithmic relative concentration of \( A \) against that of \( R \) (relative to initial conditions; \( c_{A,0} \)). The second-order heterogeneous rate constant of the compound of interest \( (k_{2,A}) \) towards the oxidant may then be calculated using the obtained \( k_r \) and the known \( k_{2,R} \) \((\text{ie: } k_{2,A} = k_r \times k_{2,R})\).

Using this method, a number of studies have quantified the uptake coefficients of \( O_3, OH, Cl, \) and \( NO_3 \) on various organic particles, and the corresponding second order rate constants for the degradation of organic compounds (Hearn and Smith, 2006; George et al., 2007; Lambe et al., 2007; McNeill et al., 2007; McNeill et al.,...
Although gas chromatograph mass spectrometry (GC-MS) has been widely used in the kinetics studies (Weitkamp et al., 2008a; Weitkamp et al., 2008b; Lambe et al., 2009; Isaacman et al., 2012), quantifying the particle phase loss of an organic compound in such studies often relies upon aerosol mass spectrometry techniques to monitor specific particle phase reactant ions of interest in semi-real time. Aerosol mass spectrometry instruments utilizing high resolution detection and soft ionization techniques, such as chemical ionization (Aerosol CIMS) (Hearn and Smith, 2006; McNeill et al., 2007; McNeill et al., 2008; Renbaum and Smith, 2011; Sareen et al., 2013) and vacuum ultraviolet photo-ionization (VUV-ATOFMS) (Liu et al., 2012), have been utilized to measure the concentration of the target organic compounds in particles. However, Aerosol Time-of-Flight or Quadrupole Mass Spectrometry (ToF-AMS or Q-AMS) employing electronic ionization (EI; 70 eV) as an ion source remains the prevalent instrument used in such organic particle experiments. In utilizing this approach, a specific fragment (usually the fastest-decaying ions in the spectrum) is often chosen as a tracer for the particle phase compound of interest. For example, m/z 297 has been selected as a tracer for bis(2-ethylhexyl) sebacate (BES) (George et al., 2007), m/z 71 for hexacosane (Lambe et al., 2007), m/z 113 for squalane (Smith et al., 2009), m/z 104 and 144 for erythritol and levoglucosan (Kessler et al., 2010), and m/z 152, 68 and 98 for 1,2,3,4-butaneetettracarboxylic acid, citric acid and tartaric acid (Kessler et al., 2012) respectively.
However, the use of EI in conjunction with a particle vaporizer in the AMS results in heavy fragmentation for organic compounds due to the high energy associated with the EI source (70 eV) and the high temperature (~873 K) of the vaporizer (Jayne et al., 2000; Allan et al., 2003). Under such conditions, the tracer m/z fragment is prone to interferences due to (1) the fragmentation of larger ions and/or molecules and (2) fragments from particle phase oxidation products. Both can contribute to the tracer m/z signal, sometimes leading to an insensitive or nonlinear response of the tracer m/z to the concentration of the target reactant during oxidation. The same may also be true for the m/z for the molecular ion should one exist. In particular, it is true if the structure of the product is highly similar to the reactant and when the tracer is measured with a unit-mass resolution (UMR) aerosol mass spectrometer. Although it is often assumed that the chosen tracer ion does not contribute significantly to the mass spectra of any possible oxidation products or vice versa (Kessler et al., 2010), this is not always the case. In our previous work, we observed that the magnitude of the second order heterogeneous rate constant ($k_2$) increases as a function of increasing m/z of the fragment chosen as a tracer of the parent molecule (Liu et al., 2014). The same trend has also been observed for the OH oxidation of ambient biogenic secondary organic aerosol (SOA) (Slowik et al., 2012). This suggests an interference from the fragments selected, and points to the necessity to separate the signals of the compound of interest, from other compounds (products and/or fragment) for kinetic studies.

In the current study, we improve the mixed-phase relative rate technique used for
studies of the heterogeneous oxidation of organic aerosol (OA) using positive matrix
factorization (PMF) analysis of UMR-AMS derived kinetic data. Heterogeneous
kinetics of citric acid (CA) toward OH oxidation was studied in a photo-chemical
flow tube coupled to an Aerodyne C-ToF-AMS and an Ionicon Analytik High
Resolution Proton Transfer Reaction Mass Spectrometer (PTR-ToF-MS). As it was
applied to heterogeneous oxidation of ambient biogenic SOA (Slowik et al., 2012),
PMF analysis was used to successfully deconvolve the full mass spectra of the
reactant from the potential oxidation products, hence allowing proper accounting of
the time evolution of reactant concentrations during photochemical oxidation.

2. EXPERIMENTAL DETAILS

2.1 Flow tube experiments. A detailed schematic representation of the experimental
system utilized in this study has been described elsewhere (Liu et al., 2014). Briefly,
organic particles (citric acid) were generated via atomization (model 3706, TSI), dried
through a diffusion drier and size-selected with a differential mobility analyzer (DMA)
(model 3081, TSI). The dried, monodispersed CA particles were introduced into the
flow tube reactor and exposed to differing OH concentrations. OH radicals were
produced by the photolysis of O$_3$ at 254 nm in the presence of water vapor. O$_3$ was
generated by passing zero air through an O$_3$ generator (OG-1, PCI Ozone Corp.). The
O$_3$ concentration in the reactor was measured using an O$_3$ monitor (model 205, 2B
Technologies) and ranged from 0-1000 ppbv. Relative humidity (RH) in the reactor
was held constant (30±3) % by varying the ratio of wet to dry air used as an air source,
and was measured at the exit of the flow tube reactor. The temperature was held
constant at 298 K by circulating a temperature controlled fluid through the outer
jacket of the reactor. The residence time in the flow reactor was 52 s. The steady-state
OH exposures were varied from 0 to ~7.0×10^{11} molecules cm^{-3} s which was estimated
on the basis of the decay of methanol from (as a reference compound) its reaction
with OH. The decay of methanol from its reaction with OH was measured using the
PTR-ToF-MS. The $k_2$ of methanol, 9.4×10^{-13} cm^{3} molecule^{-1} s^{-1}, was used for the OH

OH radical reactions were performed in a custom-made reactor consisting of two
electro-polished stainless steel cylinders with inner diameter of 7.3 cm. The first stage
contained static mixing elements (StaMixCo) to ensure that particles and gas phase
species were well mixed prior to entering the reaction region (second stage). Fluid
dynamics simulations of the flow tube confirmed that particles and gas phase species
were well mixed in the reactor, with a uniform initial velocity profile. The size and
composition of the particles exiting the reactor were measured by a scanning mobility
particle sizer (SMPS, TSI) and an Aerodyne C-ToF-AMS (Drewnick et al., 2005)
Control experiments demonstrated that O₃ or 254 nm light exposure did not lead
to the decomposition of CA. Analytic grade CA (EM, Germany) was used as received.
18.2 MΩ water was used as solvent.

2.2 PMF analysis and kinetics calculation.

PMF is a multivariate factor analysis tool that decomposes a matrix of speciated
sample data into two matrices, namely, factor contributions and factor profiles
(Paatero and Tapper, 1994), such that
\[ x_{ij} = \sum_p g_{ip} f_{pj} + e_{ij} \quad (5) \]

Where \( i \) and \( j \) refer to row and column indices in the matrix, respectively, \( p \) is the number of factors in the solution, \( x_{ij} \) is an element of the \( m \times n \) matrix \( X \) of measured data elements to be fit, and \( e_{ij} \) is the residual. Results are constrained so that no sample can have a negative source contribution. The PMF solution minimizes the object function \( Q \) (Eq.6), based upon the uncertainties \( u \) (Norris and Vedantham, 2008).

\[ Q = \sum_{i=1}^{n} \sum_{j=1}^{m} \left( \frac{e_{ij}}{u_{ij}} \right)^2 \quad (6) \]

Its ability to separate the signals of a multi-component matrix has been well established. PMF analysis has been widely used for source apportionment of ambient particles in field measurements (Song et al., 2006; Yuan et al., 2006; Viana et al., 2008; Ulbrich et al., 2009; Liggio et al., 2010; Schwartz et al., 2010). Three secondary organic aerosol factors (SOA1, SOA2, SOA3) have been identified for OH initiated oxidation of laboratory SOA (George and Abbatt, 2010). Similarly, SOA factors have also been successfully isolated in OH oxidation of ambient biogenic SOA (Slowik et al., 2012). Therefore, the use of PMF for separating the reactants from the products in laboratory studies aimed at using the relative rates method for heterogeneous kinetic studies would seem to be a reasonable approach.

The AMS data for CA oxidation from all experiments combined were used as input into the PMF Evaluation Toolkit (PET) v2.05 (Paatero, 1997; Paatero and Tapper, 1994; Ulbrich et al., 2009) to separate the signals of CA and the corresponding oxidation products. In the AMS data, the \( m \) rows of \( X \) are ensemble average mass
spectra (MS) of typically tens of thousands of particles measured over each averaging period (typically 2 min) and the $n$ columns of $X$ are the time series (TS) of each m/z sampled.

PMF analyses were done in the robust mode. The default convergence criteria were not modified. The Q values as a function of FPEAK from -1 to +1 were examined (Reff et al., 2007). For the variables with signal-to-noise ratio (SNR) less than 0.2 ("bad" variables) and downweight variables with SNR between 0.2 and 2 ("weak" variables), their error estimates were increased by a factor of 10 and 3, respectively, as recommended by Paatero and Hopke (Paatero and Hopke, 2003). In this study, the SNR of all m/z fragments are larger than 0.2. The error values for m/z 44, 18, 17 and 16 were multiplied by sqrt(4).

The extracted factor profiles (mass spectra for CA and the oxidation products) were compared with the NIST mass spectrum of pure CA and that measured with the C-ToF-AMS directly via atomization. The temporal concentration profiles (factor contributions) of CA were further confirmed via comparison to the known experimental conditions used for kinetics calculations (ie: zero OH exposure should result in a CA factor contribution of 100 %). For comparison with the PMF results, the kinetic rate constants ($k_r$) were also calculated using specific individual tracers of CA at m/z 87, 129 and 147, separately. The $k_r$ of CA toward methanol was calculated according to Eq. (4). The $k_2$ of CA was further calculated with the known $k_2$ of methanol and $k_r$.

The reactive uptake coefficient of OH (γOH) with CA was calculated using the
following formulation (Kessler et al., 2010; Worsnop et al., 2002; Kessler et al., 2012; Liu et al., 2012):

\[ \gamma = \frac{2D_p \rho_{CA} N_A}{3v_{OH} M_{CA}} \varphi k_2 \]  

Eq (7)

Where \( D_p \) is the surface-weight average particle diameter of unreacted particles (cm), \( \rho_{CA} \) is the density of CA (g cm\(^{-3}\)), \( N_A \) is Avogadro’s number, \( v_{OH} \) is the average speed of OH radicals in the gas phase (cm s\(^{-1}\)), \( M_{CA} \) is the molecular weight of CA (g mol\(^{-1}\)), and \( \varphi \) is a correction factor for diffusion of OH from the gas phase to particle phase.

3.0 Results

3.1 PMF analysis of AMS data.

To ensure that oxidation of CA in particles does not result in a PTR-ToF-MS response for methanol in the gas phase (thus compromising the OH radical reference measurement), the oxidation of pure CA was performed in the absence of methanol, with no gas phase methanol signal detected by the PTR-ToF-MS. The mass concentration of the OA measured with the AMS during oxidation is shown in Figure 1A, which was constant. The results of Figure 1A demonstrate that the aerosol source is adequately stable for kinetic studies to be performed.

A two factor solution from the PMF analysis accounts for 99.98 % of the variance of the data. When the number of factors is greater than 2, none of the obtained factors resembles that of pure CA, whose contribution should be approximately 100 % when OH is absent in the reactor. Figures 1B and C represent the temporal variations of the typical 2-factor PMF solution of AMS data when CA is
exposed to varying OH concentrations. The error bars indicate the rotational uncertainty in the PMF analysis. Three independent experiments were performed to test the response of CA signal to OH exposure (determined by O₃ concentration). In the first and the third experiments, OH exposure was stepped downwards (high to low OH) by changing the power of O₃-generator with the same flow rate and RH, while the inverse sequence was performed in the second experiment.

As demonstrated in Figure 1B and C, in the absence of OH radical (labeled “0”), factor 1 (Figure 1B) accounts for (94.7±0.9) % of the OA mass, while factor 2 (Figure 1C) contributes (6.2±0.7) % of OA. This is consistent with the experimental conditions of zero OH radical (ie: no oxidation), and suggests that factor 1 should be assigned to the citric acid reactant. Impurities in the CA or the water used to atomize CA likely contributed to factor 2. When OH exposure was decreased in a step-wise manner in the first and the third experiment (Figure 1B), the extracted factor representative of CA (factor 1) increased synchronously, and is accompanied with a decrease in factor 2. Therefore, factor 2 is interpreted as the OH oxidation products of CA. This is consistent with the second experiment, where the inverse trend was observed with a step-wise OH exposure increase. Based upon this evidence, we conclude that changes in the time series of factors 1 and 2 extracted by PMF are consistent with the expected response to OH exposures, namely, that higher OH exposure resulted in a decrease in CA (factor 1) and an increase in the oxidation products (factor 2).

The factor profiles (ie: mass spectra) extracted by PMF analysis are shown in
Figure 2. The main fragments of CA including m/z values 129 (C$_3$H$_5$O$_4^+$) and 87 (C$_3$H$_3$O$_3^+$) are present in factor 1 (Figure 2A). These fragments are in good agreement with the NIST mass spectra of pure CA and the mass spectrum of pure CA particles measured with the C-ToF-AMS (Figure 3). Figure 4 further compared the normalized mass spectra of factor 1 and pure CA directly measured with the C-ToF-AMS. The relative intensities for all ions of pure CA are linearly correlated with that of factor 1 with a slope of 0.985 and R of 0.9999. This further confirmed that factor 1 should be assigned to unreacted CA. Figure 2C shows the difference mass spectra (factor 2 – factor 1). Consumption of m/z values 147 (C$_5$H$_7$O$_5^+$), 129, 87, 85 (C$_4$H$_5$O$_2^+$) and 60 (C$_2$H$_4$O$_2^+$) can be observed, which is consistent with the assignment that factor 2 belongs to oxidation products of CA. However, small changes in the relative intensities of these peaks suggest that the structure of the oxidation products of CA are likely similar to that of CA. For example, as shown in Figure 2A and B, the intensity of m/z 129 and 87 in factor 2 are 0.0125±0.0046 and 0.0218±0.0013 compared to 0.0141±0.0046 and 0.0235±0.0013 in factor 1.

The changes of the relative concentrations of gas phase methanol and particle phase CA are shown in Figure 5. The signal of CA extracted by PMF analysis also responded to OH exposure as expected, when methanol was present in the gas phase, which is similar to that of Figure 1. The relative intensities of the typical tracers of CA at m/z 87, 129 and 147 are shown in Figure 5C. As shown in Figure 5 B and C, the drop in the PMF product factor is substantially greater than that of any of the individual ions. In addition, the consumption of the smaller tracer (m/z 87) is
substantially lower than that of the larger ones (m/z values 129 and 147). For example, the maximum consumption of CA extracted with PMF analysis is approximately 80 %, in comparison to ~30 %, ~10 % and 5 % for m/z values 147, 129 and 87, respectively. These results support the small differences in the mass spectra between the unreacted CA and its oxidation products as shown in Figure 2. Furthermore, it suggests that the measured loss of these fragments, which were supposedly only derived from CA, had in fact contributions from the fragmentation of the products of CA oxidation. This ultimately would lead to an underestimation of the second order heterogeneous rate constant (or OH uptake coefficients) if these fragments were chosen as the proxies for the particle phase concentration of CA. In particular, the difference between the PMF-factor decay rate and the marker-ion decay rate is mathematically possible only when the two factors (reactants and products) are extremely similar.

3.2 Reaction kinetics. The saturation vapor pressure of CA at 298K is 1.6×10^{-7} Pa (Huisman et al., 2013), thus 99.9% of CA should be present in the particle phase under the current experimental conditions according to a partition model (Kroll and Seinfeld, 2008; Pankow, 1994). Although new particle formation was observed with a CPC in the experiments (at the exit of the reactor), it has no influence on the measured mass concentration of OA due to the small particle size of the new particles. This is well supported by the constant mass concentration of OA measured with the AMS during oxidation experiments (Figure 1A). In addition, as pointed out in our previous work (Liu et al., 2014), evaporation of CA from particles could potentially contribute to the decreases in particle phase CA concentration observed as a function of OH
exposure. If CA evaporation was at play, the derived reaction rates will be overestimated using either the present approach or the simpler methods of using single fragments. The evaporation of CA from the particle phase under these experimental conditions from control experiments is less than 0.005% based upon an evaporation model (Jacobson, 2005). This implies that the observed changes in CA concentration in the particle phase were due to the particle phase reaction.

The relative rates (relative to initial conditions) for CA and methanol in these experiments are shown in Figure 6. The logarithmic c/c₀ of CA both measured with the tracers and extracted with PMF analysis linearly correlated to that of methanol with $R^2 > 0.95$. The derived relative rate constant based upon PMF analysis is $3.01 \pm 0.27$, while it is $0.72 \pm 0.05$ and $0.22 \pm 0.01$ for m/z values 147 and 129, respectively. Applying the $k_2$ value of methanol towards OH of $9.4 \times 10^{-13}$ cm³ molecule⁻¹ s⁻¹ at 298 K (Atkinson and Arey, 2003), the $k_2$ of CA is calculated as $(2.83 \pm 0.25) \times 10^{-12}$ using the PMF approach, or $(6.77 \pm 0.47) \times 10^{-13}$ using the single tracer at m/z 147 and $(2.02 \pm 0.01) \times 10^{-13}$ cm³ molecule⁻¹ s⁻¹ using the single tracer at m/z 129, respectively. The reaction between methanol and OH radicals occurs in the gas phase, while for the CA oxidation it occurs in particle phase. Thus it is necessary to correct for OH diffusion from the bulk gas phase to the particle phase. Applying a diffusion correction utilizing a previously developed empirical formula (Fuchs and Sutugin, 1970; Worsnop et al., 2002; Widmann and Davis, 1997), the diffusion-corrected $k_2$ is $(3.31 \pm 0.29) \times 10^{-12}$ using the PMF approach, and $(7.92 \pm 0.55) \times 10^{-13}$ and $(2.36 \pm 0.01) \times 10^{-13}$ cm³ molecule⁻¹ s⁻¹ using single tracers at m/z
The diffusion-corrected $\gamma_{OH}$ is calculated as $2.74\pm0.24$ using the PMF approach, and $0.66\pm0.05$ and $0.20\pm0.01$ using single tracers at m/z 147 and 129, respectively. The $\gamma_{OH}$ for a number of different organic particles have previously been measured, and are in the range of $0.3 - 2.0$ (Kessler et al., 2010; George et al., 2007; Hearn and Smith, 2006; Lambe et al., 2007; Smith et al., 2009; Kessler et al., 2012). As pointed out by Hearn and Smith (2006), the large $\gamma$ implies that secondary radical reactions within the particles could play a significant role in heterogeneous chemistry, particularly since gaseous citric acid in these studies is insignificant.

It should be pointed out that oxidant diffusion in the particle phase should lead to a concentration gradient of oxidant and a negative impact on reaction kinetics (Donahue et al., 2005). However, as shown in Figure 6, this effect is negligible under the current experimental conditions. Based upon the measured c/c$_0$ and the initial diameter of the CA particles, the maximum OH diffusion depth is approximately 25 nm. Given the residence time ($\tau$) in this study (52 s), a significant OH concentration gradient will exist in particle phase if the $D_{OH}$ in CA particles is smaller than $1.2\times10^{-17}$ m$^2$ s$^{-1}$ ($D=\frac{l^2}{\tau}$) (Donahue et al., 2005). At the present time, the $D_{OH}$ in CA particles is unavailable. However, Price et al. (Price et al., 2014) have reported the diffusion of D$_2$O in several organics (sucrose and levoglucosan) to be larger than $\sim1\times10^{-16}$ m$^2$ s$^{-1}$ even under dry condition. This implies that a gradient in OH concentration in the CA particles is negligible under the current conditions.

4. Discussion
Kessler et al. (Kessler et al., 2012) have reported the $k_2$ of CA toward OH to be $(4.3\pm0.8)\times10^{13}$ cm$^3$ molecule s$^{-1}$ at 308 K and 30 % RH with an Aerodyne HR-AMS. In their work, the diameter of particles and RH were equivalent to the current work, while their experimental temperature was 10 K higher. In addition, a m/z fragment of 68 was used as a tracer for CA in their work to derive the heterogeneous rate constant. Conversely, no significant consumption of m/z 68 was observed in the current study. The lack of a m/z 68 fragment consumption here may be explained by the choice of reaction conditions. In the work of Kessler et al., OH concentration exposure $(0-7\times10^{12}$ molecule cm$^{-3}$ s) was approximately an order of magnitude higher than that reported here. Recent evidence suggests that the product distribution during OA oxidation greatly depends upon OH exposure levels (Wilson et al., 2012). Hence, it is possible that more oxidized products formed via multi-generational chemistry at high OH were formed, which may have less of an influence on the signal of the chosen tracer (m/z 68), and result in product AMS spectra which are significantly different than that of the reactant thus mitigating the use of PMF (which was not the case here). In studies of the OH oxidation of squalane (Sq) (Smith et al., 2009; Wilson et al., 2012), the first generation product (SqO) was the primary contributor to the products when the OH exposure was the same as the highest OH level in the current study $(7.0\times10^{11}$ molecules cm$^{-3}$ s), while higher generation products were predominant at OH exposures was greater than $\sim2\times10^{12}$ molecules cm$^{-3}$ s. Although not directly comparable, it is reasonable to assume that lower OH exposure in the current work should lead primarily to the first generation products, which are highly similar to CA.
However, the formation of multi-generation products cannot be completely ruled out. Secondly, the difference in OA and oxidant concentrations as well as timescale may also have an influence on the product distribution. Thirdly, a high resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) was used in their work, while a C-ToF-AMS with unit-mass resolution was used in this study. The higher mass resolution of the HR-ToF-AMS relative to a C-ToF-AMS, may further reduce the influence of product fragments on m/z 68 (or others). Finally, differences in temperature or other reaction conditions between experiments may also have led to differences in the morphology of CA and subsequent differences in the reactivity of CA.

Given the above discrepancy in the consumption of m/z 68, the fragments at m/z 129 and 147 were used as tracers in this work. The measured $k_2$ of CA utilizing m/z 129 and 147 in this study is of the same order of magnitude as that reported in Kessler et al. (2012). However, as shown in Figure 3C, the consumption of m/z 87 is much lower than that of m/z 129. The apparent $k_2$ of CA based on m/z 87 is $(9.9\pm0.8)\times10^{-14}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, and the diffusion-corrected $k_2$ is $(1.16\pm0.09)\times10^{-13}$ cm$^3$ molecule$^{-1}$ s$^{-1}$. This suggests that the derived rate constant greatly depends upon the size of the tracer fragment, with larger fragments resulting in larger values of $k_2$ in this study. This is consistent with previous work investigating the OH oxidation of tris-phenyl phosphate (TPhP) (Liu et al., 2014) and ambient biogenic SOA (Slowik et al., 2012). The $k_2$ of CA based upon PMF analysis is approximately an order of magnitude larger than the Kessler result measured with the tracer at m/z 68, and 4.2 times greater than
that calculated based upon m/z 147 in this study. The differences in product
distributions that may arise between this and the work of Kessler et al., consistent
with the fact that m/z 68 is not consumed in the current study, suggests that the PMF
approach was likely required in this work to separate similar product and reactant
spectra ultimately caused by lower OH exposure.

A number of factors may be responsible for the discrepancy between derived rate
constants. It has previously been observed that the presence of O₃ can inhibit the rate
of OH reaction, perhaps by reacting with OH radicals or by O₃ or intermediate species
blocking surface active sites (Renbaum and Smith, 2011). A Langmuir-Hinshelwood
mechanism has been observed for the reaction of O₃ on organic surfaces (Pöschl,
2005). It has also been demonstrated that a higher concentration of gas phase reactant
often leads to a lower uptake coefficient due to surface saturation (Ma et al., 2010 ;Li
et al., 2002). Differences in $k_2$ may also arise from the competition between reaction
products and reactants for available OH, or via the blocking or coating of the reactant
by products which would require liquid phase diffusion of OH to degrade the original
CA. In experiments with higher OH exposures (Kessler et al 2012) it is possible that
significantly more product mass is mixed and/or coated onto the original particle thus
decreasing the perceived $k_2$. Different timescales and concentrations of reactants
might also lead to different rate constants (Che et al., 2009). Finally, as pointed out
above, the differing reaction conditions may have led to a different CA morphology
and subsequent differences in the reactivity towards OH. The significant difference
between the reported rate constants highlights an important issue in heterogeneous
reactions of the atmosphere, and in the experiments trying to derive such kinetics. It implies that the particle composition and/or morphology as determined by the reaction conditions in the laboratory or the ambient atmosphere will have a large effect on the OH kinetics.

Citric acid is a hydroxyl substituted poly carboxyl acid. Scheme 1 summarizes its possible fragmentation pathways. The typical mass peaks including m/z 147, 129, 87, 85 and 68 would result from this scheme and were indeed observed. The fragments at m/z 129, 87, 85 and 68 are also likely from CA oxidation products fragments, and hence their signal intensities may be highly influenced by products and/or larger fragments, in particular, when the oxidized products are highly similar to the reactant. In some instances, oxidation products can exhibit similar fragmentation pathways as the reactants. This is likely the case for the smaller fragments of CA. For example, scheme 2 illustrates the possible fragmentation pathways of 2,3-dihydroxypropane-1,2,3-tricarboxylic acid, which is one of the possible products from the OH oxidation of citric acid in terms of the general chemistry described by Atkinson et al. (Atkinson, 1986). As observed in Scheme 2, there are several pathways leading to the fragment at m/z 87, implying that the decrease in the signal of m/z 87 due to CA oxidation is likely to be compensated by fragments from the oxidation products. In addition, it is also possible to form fragments with the same m/z as the parent citric acid if the dehydration reaction (the 6\textsuperscript{th} path in Scheme 2) takes place initially. Other possible reaction products might also play similar roles in the fragments. This is highly possible when the product distribution contains products.
which are structurally similar to the reactant under low oxidant exposure conditions.

Scheme 1. Possible fragmentation pathways for citric acid.
Scheme 2. Possible fragmentation pathways for an oxidation product of citric acid in terms of the general chemistry described by Atkinson et al. (Atkinson, 1986).

The current relative rate method based upon PMF analysis was used to re-analyze the heterogeneous oxidation kinetics of three organophosphate flame retardants found in ambient particles and reported previously (Liu et al., 2014). The $k_2$ values for TPhP, tris-1,3-dichloro-2-propyl phosphate (TDCPP) and tris-2-ethylhexyl phosphate (TEHP) utilizing the tracer and PMF approaches are summarized in Table 1. For TPhP, its molecular ion peak ($M^+$;326) was chosen as a tracer; while the largest detectable fragments, i.e. m/z 381 and 323, were chosen for TDCPP ($M^+$; 431) and TEHP ($M^+$;
435), respectively. The typical evolution of the PMF factors of TPhP, TDCPP, and TEHP are shown in Figures S1-3. For TPhP, the measured $k_2$ values derived by both methods are comparable within the experimental uncertainties, while $k_2$ of TDCPP and TEHP based upon PMF analysis is 1.5 and 1.6 times larger than that using the chosen tracers. The good agreement between methods for TPhP is likely due to the fact that the molecular ion peak ($M^+$) is measurable for TPhP with the AMS, while it is not observable for TDCPP, TEHP and CA. Therefore, the influence of secondary fragmentation from larger fragments has little influence on the signal of $M^+$ for TPhP. These results also demonstrate that a substantial underestimation of rate constants could result when a non-molecular ion tracer is used to monitor the particle phase concentration of organic matter with UMR-AMS for heterogeneous kinetic studies. The discrepancy between the tracer and PMF based methods for other compounds will depend upon a number of factors including: structure of products, OH exposure level, particle morphology, and organic species competing OH reactions.

5.0 Implications and Conclusions

The measured $k_2$ for citric acid toward OH is $(3.31\pm0.29) \times 10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ at 298 K and 30 % RH. This value is at least 4.2 times greater than that calculated on the basis of atypical tracer m/z 147. Although the tracer and PMF approaches can at times agree (Kroll, 2014) and the tracer and molecular-ion approaches can sometimes agree as well (Smith et al., 2009), our results suggest that the heterogeneous kinetics of OA is underestimated when a non-molecular ion peak is
used as the tracer to measure the particle phase concentration of OA based on UMR-AMS. In model simulations, the reactive uptake coefficient of OH or other radicals, which are calculated based upon $k_2$, is an important parameter in evaluating the fate of OA during transport. The current results suggest that the lifetime of OA estimated in models due to heterogeneous oxidation might be overestimated for a reaction system where the products are highly similar to the reactant and the kinetic data are derived by individual non-molecular m/z tracers of OA. The results also suggest that it may be necessary to revisit the kinetic data of other organic aerosol components (and OH uptake coefficients) which have been derived using the relative rates technique (George et al., 2007; Lambe et al., 2007) based on UMR-AMS. Finally, these results imply that the heterogeneous oxidation of aerosols will be dependent upon a number of factors related to the reaction system, and that a single rate constant for one system cannot be universally applied under all conditions. Future work is thus required to elucidate the chemical and physical parameters which control the OH heterogeneous reaction kinetics and the associated need to apply PMF for a variety of chemical systems. This may be best accomplished through systematic application of the PMF approach to species with differing mass spectral characteristics, such as linear/branched alkanes, monocarboxylic acids and other oxygenates. As illustrated in this study, the kinetics derived with PMF may differ from that derived with tracer ions, with both based upon EI-AMS approaches. This highlights the usefulness of measurements from CIMS, GC-MS and VUV-AMS, for the determination of heterogeneous loss rates, since these instruments are more likely to retain the reactant
molecular information.

Supporting Information

Supplementary material related to this article is available online at:

Acknowledgements

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Literature Cited:


George, I. J., and Abbatt, J. P. D.: Chemical evolution of secondary organic aerosol from OH-initiated


Table 1. Comparison of the measured $k_2$ values utilizing PMF and select m/z tracers, for organophosphate compounds and CA.

<table>
<thead>
<tr>
<th>OA</th>
<th>Mean $k_{r, PMF}$</th>
<th>$k_2 \left(10^{12}\right)$ cm$^3$ molecule$^{-1}$ s$^{-1}$</th>
<th>$k_2, \text{obs}_{PMF}$</th>
<th>$k_2, t, PMF$</th>
<th>$k_2, t, \text{Tracer}$</th>
<th>$k_2, \text{Tracer}$</th>
<th>M$_{\text{Tracer}}$/M$^+$</th>
</tr>
</thead>
<tbody>
<tr>
<td>TPhP</td>
<td>1.58±0.33</td>
<td>1.48±0.31</td>
<td>1.95±0.43</td>
<td>2.10±0.19</td>
<td>2.10±0.19</td>
<td>0.9</td>
<td>326/326</td>
</tr>
<tr>
<td>TDCPP</td>
<td>1.20±0.31</td>
<td>1.13±0.29</td>
<td>1.35±0.35</td>
<td>0.92±0.09</td>
<td>0.92±0.09</td>
<td>1.5</td>
<td>381/431</td>
</tr>
<tr>
<td>TEHP</td>
<td>3.52±0.65</td>
<td>3.31±0.61</td>
<td>4.25±0.78</td>
<td>2.70±0.63</td>
<td>2.70±0.63</td>
<td>1.6</td>
<td>323/435</td>
</tr>
<tr>
<td>CA</td>
<td>3.01±0.27</td>
<td>2.83±0.25</td>
<td>3.31±0.29</td>
<td>0.79±0.06</td>
<td>0.79±0.06</td>
<td>4.2</td>
<td>147/192</td>
</tr>
</tbody>
</table>

a.(Liu et al., 2014)
Figure captions

Figure 1. Changes in (A) total organic mass concentration, (B) the fraction of unreacted citric acid derived by PMF and (C) products of citric acid oxidized by OH derived by PMF, as a function of relative experimental time. The values 1 – 6 represent a step-wise O₃ concentration decrease, corresponding to decreased OH exposure; 0 represents an O₃ concentration of zero. Experimental conditions are $D_m$: 200 nm, RH: 30±3 %, T: 298 K.

Figure 2. Normalized mass spectra of (A) citric acid (PMF factor 1), (B) citric acid oxidation products (PMF factor 2), and (C) the difference mass spectrum (Factor 2 – Factor 1). The numbers in the upper two rows are the intensities of m/z 87 and 129, while negative values are shown in the bottom row. The red and green lines indicate a negative and positive value, respectively.

Figure 3. Mass spectra (A) of CA from NIST database, (B) of pure CA measured with the C-ToF-AMS.

Figure 4. Comparison between the mass spectra of factor 1 from PMF analysis and pure CA directly measured by the C-ToF-AMS. The inset graph is the correlation of their corresponding signal intensities.

Figure 5. Changes in the relative concentration of (A) methanol, (B) citric acid extracted with PMF analysis and (C) specific tracers measured with the AMS during the OH initiated oxidation of citric acid. Experimental conditions are $D_m$: 140 nm, RH: 30±3 %, T: 298 K. The values in the top row represent the OH exposures.

Figure 6. Relative concentration of citric acid ($c/c_0$) as a function of the relative concentration of methanol based upon (A) PMF analysis, (B) m/z=129 and (C) m/z
147. Experimental conditions are $D_m$: 140 nm, RH: 30 ± 3 %, T: 298 K.
Figure 1.
Figure 2.
Figure 3.
Figure 4.
Figure 5.
Figure 6.