Wang et al. presented a study that evaluated the performance of different particle wall loss correction methods for aging experiments of α-pinene ozonolysis products. This paper may potentially be useful to the SOA chamber community. However, there are portions of the manuscript that are vague and confusing, and they need to be addressed before it can be considered for publication.

We address the various comments of the reviewer below. Our responses (regular font) and corresponding changes in the paper follow each comment (in italics).

(1) Page 5, line 141: How many particle wall loss experiments are typically performed during the year? How reproducible are these particle wall loss rates? It would be useful to show the rates obtained during these (1-year worth of) experiments in the SI.

The annual number of wall loss experiments has been variable. During 2016 there were around 15 experiments. After their potential variability became clear due to the maintenance of the simulation chamber, we perform them every week. The reproducibility is clearly related to the status of the chamber. When the chamber is in its disturbed state loss rates can vary by more than a factor of 5 and they can be quite high. When the chamber is in its undisturbed state the loss rates are a lot smaller and they can vary by 0.05 h^{-1} or so. The 4 curves shown in Fig. 3 are representative of the range. We have added some text in the paper to discuss this variability.

(2) Figure 1: Why are the measured k’s only shown for particles of diameters < 400 nm and < 800 nm for the 12 m^3 and 1.5 m^3 reactors? What about the particles with larger diameters? Were those measured? If not, why not?

The particle loss rate constants for a given size can be measured reliably only when there are enough particles of this size available in the system. In the experiments shown, the produced ammonium sulfate particle size distribution included few larger particles. As a result, the k’s at bigger particle sizes had high uncertainty associated with them due to lack of particles, and were thus excluded from the figure. This is now explained in the manuscript.

(3) Page 8, line 226: I am inferring from Fig 1 that if $k_a$ is larger than $k_c$, coagulation is a significant particle loss process. Is this correct? If yes, this should be stated explicitly in the main text. Currently, there is little explanation in the text of what differences between $k_a$ and $k_c$ shown in Fig. 1 means.

The reviewer is correct that coagulation is a significant particle loss process for the size range where $k_a$ is larger than $k_c$. We made corresponding changes in Section 4.1 of the revised manuscript to explicitly state the above fact and address the differences between $k_a$ and $k_c$.

(4) Page 8, line 232: It would be useful to show in the SI how the particle wall loss rate constants look like after correcting for coagulation.
Figure 1 depicts the effect of the coagulation correction. A reference to that figure and some additional discussion have been added at this point.

(5) Page 8, line 234: “The uncertainty of … is significantly higher …” How much higher? Please give a number.
We now state that the uncertainty increases from approximately 10% to 50%.

(6) Page 8, line 236: “This is due to … versus the linear regression that uses the measured.” It is unclear which linear regression fit the author referring to. Is it the linear regression fit to each particle size bin or the linear regression fit of the measured $k_a$’s?
We refer to the linear regression fit of the logarithm of the concentration of the particles in each size bin. This is now clarified in the text.

(7) Figure 2: Why doesn’t $k_c$ correct for the effect of particle wall loss on the number concentration well? This was not explained in the main text.
$k_c$, being the coagulation-corrected particle wall-loss rate constant, intrinsically excludes the impact of coagulation on particle number concentration. Since coagulation reduces particle number (but conserves mass), $k_c$-corrected particle number concentration is lower than the $k_a$-corrected one, with the difference attributed to the coagulation rate. The corresponding explanation has been added to the paper.

(8) Page 9, line 243: Why weren’t the $k_a$’s for $D_p < 50$ nm measured? If they were measured, there would be no need to back extrapolate the data.
The reason for extrapolation of $k_a$’s for $D_p < 50$ nm is that the measured $k_a$’s were extremely uncertain due to lack of particles at those small sizes and thus could not be used. This is now explained in the revised paper.

(9) Page 9, line 249: “As a result, the coagulation effect is almost an order of magnitude higher than the average …” What do you mean by the “coagulation effect”? Are you referring to the rate constants? Please clarify.
We have rewritten the corresponding sentence which refers to the difference between $k_a$ and $k_c$ due to coagulation.

(10) Page 9, line 259: “Our results are consistent with their low seed …” Which results are you referring to? Your SOA mass yields? Your observation that coagulation plays a minor role in some of your experiments? Please clarify.
We have deleted this rather confusing sentence.
(11) Page 9, line 262: What were the particle number concentrations used in these ammonium sulfate-only experiments? Were they somewhat similar? Also, the uncertainties are missing from the figure.

The particle number concentrations used in these experiments varied from approximately 10,000 to 40,000 cm$^{-3}$. The uncertainties of the $k_c$ curves are shown in Figure S1. We made the corresponding changes in the revised manuscript to direct the readers to Figure S1 for the uncertainties.

(12) Page 10, line 283: How did you determine when condensation/evaporation was minimal in your experiments? By using your SMPS data? Or the AMS data?

We used both the SMPS and the AMS measurements. This is now mentioned in the text.

(13) Page 10, line 285 to 293: The authors claimed that $k_a$ and $k_c$ are almost the same except for particles smaller than 100 nm. Can the observed differences at the smaller particle sizes really explain the differences in the size-independent loss rate constant? A more detailed analysis should be presented.

There are two different issues here that should not be confused. First there is the difference between the two rate loss constants which increases for smaller particles. Then there is the size dependence of either constant. The errors introduced during the use of a size-independent rate constant are mainly affected by the size dependence of the “true” rate constant over the size range covered by the aerosol size distribution. We have added a more detailed discussion of this point.

(14) Figure 4: Was the SMPS only scanned up to 300 nm? If yes, why was this the case? It looks like that the entire volume/number concentration distribution was not captured. This may affect the accurate determination of SOA mass concentrations in Fig. 5.

The SMPS measured up to 700 nm for this experiment. Due to the large uncertainty of $k_c$ at diameters above 300 nm, we assumed a constant $k_c$ value for the larger sizes. This is a reasonable assumption because $k_c$ remains relatively constant from 300-500 nm based on our measurements (Fig. 3) when the chamber is undisturbed. Please note that most of the SOA mass in this experiment was in particles smaller than 300 nm (Fig. 4b). Thus assuming a constant wall loss rate constant for particles larger than 300 nm should have only a small effect on the corrected SOA mass concentration. A brief discussion of this point has been added to the paper.

(15) Page 10, line 295 and Figure 5: What is the main conclusion of section 4.3? Should the size-independent method be used given that their corrected SOA mass concentrations are so different?

We conclude the corrected SOA mass concentration can vary by 20-30% depending on which wall-loss rate constants are used. We recommend using the size-dependent wall loss rate constants for the correction. However, when the chamber is undisturbed and the duration of the experiment is a couple of hours, using the size-independent wall loss rate constant derived from the initial 4-
hour seed wall-loss period can give relatively accurate results (errors of 5% or less). We have rewritten part of this paragraph to make our conclusions clear.

(16) Page 11 line 315: “Particles of smaller sizes with larger organic to sulfate ratios…” Loza et al. (ACP 2012) previously discussed in detail the effect of size dependence wall loss on the organic to sulfate ratio. Thus, that paper should be cited.
We have added the citation to the work of Loza et al. (2012) at this point.

(17) Section 4.5: When were the particle wall loss correction applied? Just before the start of SOA formation (time = 0 min) or when the ammonium sulfate seed was first injected into the chamber? The particle wall loss correction was applied when the ammonium sulfate seed was first injected into the chamber, as depicted in Figure 8a. This is now clarified in the paper.

(18) Page 14, line 420: It was recommended that metal gloves be used if it was absolute necessary to touch the chamber walls. Was this recommendation tested? Did using metals gloves really resulted in less electrostatic forces in the chamber and thus smaller differences in the measured \( k \)’s? If yes, the data should be shown to back up this recommendation.
We have rephrased this sentence given that we have only anecdotal evidence to support this recommendation. We now state that “other practices like use of metal gloves can help reduce the build-up of static electricity on the chamber walls”.
Responses to the Comments of Reviewer 2

Wang et al. reviewed and presented a comprehensive study about current methods to evaluate the particle wall-loss rate in CMU smog chambers. Particle wall-loss correction in smog chamber is a very important topic and can be applied by the chamber community in both experimental data interpretation and chamber simulations. This manuscript is well organized and very informative. But I found several parts confusing, which have to be clarified before considering for publication. We address the various comments of the reviewer below. Our responses (regular font) and corresponding changes in the paper follow each comment (in italics).

General comments:

(1) About $k_c$. My understanding of the difference between $k_c$ and $k_a$ is that (correct me if I am wrong): the coagulation-corrected $k_c$ is actually the inherent particle wall-loss rate, which reflects the effect of all physics other than coagulation inside the chamber; while the apparent $k_a$ is the synergistic effects of diffusion, gravity, eddy intensity, coagulation, and even charge, among which coagulation can be isolated to derive $k_c$. Then it is very confusing when the experimental data is corrected by $k_c$: should coagulation effect be counted? For example, to clarify that coagulation is important in particle number concentration decay but not in volume concentration decay in Fig. 2, I think $k_c$-corrected curve does not count coagulation (Eq. (6)), otherwise, it should overlap with $k_a$-corrected curve. How about in other cases? I thought the right way to perform particle wall-loss correction was to insert the derived coagulation-free $k_c$ into the general dynamic equation to get the right particle number concentration. The authors may want to clarify this point in the revised manuscript.

The reviewer’s understanding of $k_a$ and $k_c$ is accurate. The reviewer is also correct about the fact that $k_c$-corrected curve in Fig. 2 shows the evolution of the particle number concentration correcting for their losses to the wall. The change in the number concentration in this case of a non-reacting system is only due to coagulation. This is the goal of these wall corrections, to correct for this process. The method used for the correction here is for all practical purposes equivalent to that proposed by the reviewer because the measured concentration is used for the correction at each time step. This concentration is continuously changing due to condensation and coagulation, but the effects on these processes that are described by the general dynamic equation are “included” in the measurements. A brief explanation has been added to the revised paper.

(2) About SOA correction. How are the particles deposited on wall treated in the SOA correction? Are they still acting as a condensation sink of VOC molecules or just removed from the system during the correction? It looks like that Eq. (7) treats the deposited particles the same as suspended particles. Moreover, what time does $V_s$ refer to? The beginning of the injection of seeds or the beginning of the SOA experiment? These points should be clarified in the revised manuscript.
The interaction of the condensing vapors with the walls (including the particles deposited on the walls) is clearly an important research topic but has been out of the scope of this work. Our approach implicitly assumes that the particles deposited on the walls are removed from the system and stop interacting with the gas phase. $V_s$ refers to the corrected seed volume concentration right before SOA formation. Note that the $V_s$ is for all practical purposes constant during the seed-only periods after particle wall-loss correction. These points are clarified in the revised paper.

Specific comments:

(3) In Section 2, what are sampling rates in both 12 m$^3$ and 1.5 m$^3$ chambers? Are there any significant volume changes during the experiment, especially for 1.5 m$^3$ chamber?
For both experiments, the SMPS was sampling at 0.3 L/min. For a 5-hour experiment, the volume lost due to sampling is 0.09 m$^3$. This is a small change for both the 12 m$^3$ and the 1.5 m$^3$ chambers. This information has been added to the paper.

(4) What is the scanning time of SMPS? How does this reconcile with the coagulation correction algorithm, i.e., is the time step the same as SMPS scanning time? Or is the time step just 15 min as mentioned in Line 196? More details should be included in Section 3.1.1.
The SMPS scanning time was 5 min for each sample. The time step of 15 min mentioned in Line 196 is the result of averaging of 3 consecutive samples and use of the corresponding average particle number distribution as input for the model. We added the corresponding information in Section 3.1.1 in the revised manuscript.

(5) In Section 4.1, notes should be added that the particle number concentrations in Exp. 1 and Exp. 2 have 1 order of magnitude difference, thus coagulation effect is more significant in the small chamber.
We made the corresponding change in Section 4.1 in the revised manuscript.

(6) The authors may want to replace Fig. 3 with Fig. S1. There are no uncertainties in Fig. 3 but Fig. 1S has. Also, I suggest changing the name of legends. Is the first number referring to the experiment times? Over the past four years, is the chamber renewed? If so, is there any effect?
This should be clarified.
We used Fig. 3 in the main text because the uncertainty area (as shown in Fig. S1) can be distracting and may not allow the reader to see the differences of the various measurements. We now direct the readers interested in the uncertainties to Fig. S1. The first number refers to the month when the corresponding experiment was performed. We changed the name of the legends as suggested. The experiments discussed in the paper refer to the same chamber. This is now clarified in Section 2.1 of the revised manuscript.
(7) In Figure 4b, the number distribution may be more straightforward than the volume distribution to explain the difference between $k_1$, $k_2$, and $k_3$. It will be beneficial to mention in the caption that Figure 4c is from Period 1.
Our rationale of showing the volume distribution is that it indicates where the majority of the particle mass are distributed. This aids in our later discussion of the wall-loss corrected SOA mass concentration. We added the recommended information in the caption of Fig. 4c in the revised manuscript.

(8) Figures 6&7 are very similar to Figures 6&7 in Wang et al. (2018, https://doi.org/10.5194/acp-18-3589-2018). I am not sure if this is allowed in the policy of EGU publication. Since Figures 6&7 are from the same authors, I guess it is fine. In addition, it will be beneficial to mention about the conversion from $D_{xa}$ to $D_p$ in Section 4.4 as in Wang et al. (2018).
We have replaced these two figures with similar ones depicting results from another experiment. We have added the information about the diameter conversion in Section 4.4 in the revised manuscript.

(9) In Figure 8, which $k_c$ is more representative of the condition inside the chamber?
Both $k_c$ profiles are representative of the chamber condition, but at their corresponding time periods. When the chamber is under “disturbed” conditions, the wall-loss rate constants were observed to be time-dependent as discussed in Section 4.5. A brief discussion has been added to the paper.

(10) The authors may want to replace Fig. 9 with Fig. S3, or mention in the manuscript that Fig. S3 has uncertainties.
We used Fig. 9 in the main text because the uncertainty areas (as shown in Fig. S3) can be quite confusing for a lot of readers. We are now directing the interested readers to Fig. S3 for the corresponding uncertainties.

(11) The authors may want to pay attention to a just accepted manuscript in AS&T (https://www.tandfonline.com/doi/abs/10.1080/02786826.2018.1474167) on the similar topic as in Sections 4.5 and 4.6.
We have added a reference to the corresponding relevant article in our introduction.
Responses to the Comments of Reviewer 3

The authors present results of an experimental/modeling study aimed at evaluating the effects of particle wall loss on measurements of SOA yields, and providing recommendations for the best approach for correcting for these losses. Data acquired from chamber studies conducted over a period of 3 years were analyzed using a modified version of a model developed by Nah et al. (2017) with SMPS measurements of size-dependent seed particle wall loss at the beginning and end of experiments. Results indicate that the corrections are sensitive to particle size distributions, coagulation, and static charge that can accumulate on Teflon chambers during maintenance or experiments. The results provide quantitative insights into the consequences of wall loss, which are helpful for getting a sense of when and by how much different factors can influence corrections. I think the manuscript is concise and clearly written, and is a useful addition to the growing literature on this topic, which can sometimes be confusing. I recommend it be published in ACP after the following minor comments are addressed.

We address the various comments of the reviewer below. Our responses (regular font) and corresponding changes in the paper follow each comment (in italics).

Specific comments:

(1) It would be useful to be clear in the Conclusions how this study, the results, and the conclusions compare to those of Nah et al. (2017), which seemed to be pretty comprehensive. The inclusion here of the effect of chamber disturbances is new, and the multi-year data, but it is not clear to me what else is. We agree with the reviewer that one of the major contributions of the present work is the conclusion that a Teflon chamber can be in different states and its corresponding wall losses can have different quite different magnitudes, size dependences, and time variability. The second is the evaluation of a range of particle wall-loss correction methods, showing that most of them work reasonably well when the chamber is in its undisturbed state, but that multiple wall loss-characterization periods (before and after an experiment) are needed if the chamber has been disturbed. Finally, it is the realization that some of the differences among results of past studies may be simply due to a change in the state of the chamber that was not noticed. We have added these points in the discussion of the main results of the paper.

(2) There does not seem to be a discussion of how well the sulfate tracer method is expected to work under different conditions. The correction for organic/sulfate ratio seems related, but is not obviously the same as correcting an SOA yield. Since this method is often used, and it is much easier than the one recommended here (as long as one has an AMS), it would help to provide more quantitative comments on this method and the uncertainties that can be achieved for certain types of size distributions.
The main conclusion is that the sulfate tracer method, together with most other methods, performs reasonably well as long as the chamber is in its undisturbed state. In that case the dependence of the wall loss rate constant on particle size is relatively weak in the size ranges used in these SOA experiments and the errors introduced by assuming similar loss rate constants are usually less than 10% for the yields. The corresponding error for a chamber in its disturbed state depends both on the shape of the loss rate constant function as a function of size, on the differences of the size distributions of the organics and sulfate, and on the location of these distributions relative to the loss rate constant curves. As the size distributions are evolving during an experiment so does the error. We have performed a few tests using some experimental scenarios and the corresponding errors were of the order of 20-30%. However, it is difficult to generalize these results. The errors could be higher in some extreme cases or actually lower when the errors cancel each other. A correction similar to the one used here (taking into account the size-dependent losses and the size distributions of OA and sulfate) should be performed in this case of a disturbed chamber as a safeguard against higher errors. We have added a few sentences about this issue in the revised paper.
Response to the Comment of Simon O’Meara

The other reviewers have provided thorough (and I think valid) feedback, I have just one suggestion. It is very beneficial to the reader that $k_c$ values from the disturbed chamber have been quantified before and after an experiment, and that a clear increase is observed afterward (Fig. 8). This is not so for the undisturbed experiment. Could the authors offer some suggestions of the process causing this increase in $k_c$ during the disturbed chamber experiment?

The fact that $k_c$ became drastically time-dependent within the time frame of a 4-h experiment is most likely attributed to the increased electric field within the chamber after the maintenance of the room (Section 4.5). Repeated contact with the Teflon walls during the maintenance probably led to charge build-up on the walls. The change in the loss rates suggests a change in the electric field in the chamber during this experiment. This could be due to additional charge build-up or redistribution of the charges as the experiment progresses (lights are turned on and off, the chamber walls move due to the air motion from the temperature control system, etc.). Our main point in this section is that when there is a strong electric field in the chamber, it can also vary with time making the corrected results of the experiments quite uncertain. We have added a brief discussion of this point in the revised paper.
Particle Wall-loss Correction Methods in Smog Chamber Experiments
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Abstract
The interaction of particles with the chamber walls has been a significant source of uncertainty when analyzing results of secondary organic aerosol (SOA) formation experiments performed in Teflon chambers. A number of particle wall-loss correction methods have been proposed including the use of a size-independent loss rate constant, ratio of suspended organic mass to that of a conserved tracer (e.g., sulfate seeds), size-dependent loss rate constant, etc. For complex experiments such as chemical aging of SOA, the results of the SOA quantification analysis can be quite sensitive to the adopted correction method due to the evolution of the particle size distribution and the duration of these experiments.

We evaluated the performance of several particle wall-loss correction methods for aging experiments of $\alpha$-pinene ozonolysis products. Determining the loss rates from seed loss periods is necessary for this system because it is not clear when chemistry is over. Results from the organic to sulfate ratio and the size-independent correction methods can be influenced significantly by the size-dependence of the particle wall-loss process. Coagulation can also affect the particle size distribution, especially for particles with diameter less than 100 nm, thus introducing errors in the results of the wall-loss correction. The corresponding loss rate constants may vary from experiment to experiment, and even during a specific experiment. Friction between the Teflon
chamber walls and non-conductive surfaces can significantly increase particle wall-loss rates and the chamber may require weeks to recover to its original condition. Experimental procedures are proposed for the characterization of particle losses during different stages of these experiments and the evaluation of corresponding particle wall-loss correction.

1. Introduction

Smog chamber experiments have been an important tool for the study of atmospheric aerosol processes. One major challenge of smog chamber experiments is the particle wall-loss processes. The aerosols inside the chamber are lost to its walls due to Brownian diffusion, convection, electrostatic effects (especially for Teflon chambers) and gravitational sedimentation (Crump and Seinfeld, 1981). The particle wall-loss process is first-order and the particle wall-loss rate constant, $k$, is defined as,

$$\frac{\partial N(D_p, t)}{\partial t} = -k(D_p, t)N(D_p, t),$$

where $N(D_p, t)$ is the number concentration of particles with diameter $D_p$ at time $t$. For an aerosol population, $k$ is in general a function of particle size and time. Smaller-sized particles (less than 50 nm) have a higher loss rate due to diffusion-dominated wall-loss process while particles larger than one micron are lost mainly due to sedimentation for a reactor with low air motion inside. Electrostatic effect can play a major role for intermediate sizes (McMurry and Rader, 1985). Charan et al. (2018) studied the charge effect on the rate of particle wall deposition by parameterizing and optimizing for estimating both the eddy-diffusion coefficient and the average magnitude of the electric field within their chamber, and found it necessary to address the effect for each specific chamber.

Early studies of chamber simulations of secondary organic aerosol (SOA) formation and growth assumed that the particle wall loss is negligible in fairly large chambers (~30 m$^3$) when determining SOA yields (Stern et al., 1987). Several particle wall-loss correction methods have since been developed and adopted in chamber studies. Pathak et al. (2007) proposed a semi-empirical wall-loss correction method that involves determining the first-order particle wall-loss rate constant, $k$, from the SMPS-measured SOA mass concentration after chemical reactions have
been completed. This total mass concentration-based method is based on the assumption that $k$ is independent of particle size for the size range of particles present in the experiment and remains constant during the course of an experiment. The constant $k$ is found as the slope of the linear regression:

$$\ln[C_{SOA}^{sus}(t)] = -kt + Q,$$

(2)

where $C_{SOA}^{sus}(t)$ is the measured SOA mass concentration at time $t$ and $Q$ is an arbitrary constant. The values of $C_{SOA}^{sus}(t)$ used for the fit are taken after the SOA production has finished (condensation/evaporation is minimal). The corrected SOA concentration can be found by:

$$C_{SOA}^{tot} = C_{SOA}^{sus}(t) + k \int_{0}^{t} C_{SOA}^{sus}(t) \, dt - C_{seed}(0),$$

(3)

where $C_{seed}(0)$ is the seed mass concentration when SOA formation begins. This approach is relatively accurate when $k$ remains more or less constant over the size range of the aerosol population inside the chamber, and accounts for the experiment-to-experiment variability of the particle wall-loss rates. However, it requires a period during which no reactions are taking place in the chamber and assumes that the rate constant does not vary during the experiment.

The size-dependent correction method involves determining a first-order $k(D_p)$ through the aforementioned linear fitting of the number concentration of the suspended particles, for each size, usually with the help of a scanning mobility particle sizer (SMPS). Several studies that adopted this method determined the $k(D_p)$ profile for the corresponding chamber through seed experiments where inert (e.g., ammonium sulfate) particles were used (McMurry and Grośjean, 1985; Keywood et al., 2004; Ng et al., 2007; Fry et al., 2014; Nah et al., 2017; Fry et al., 2014). In these studies, an average $k(D_p)$ profile was applied to all experiments. This method takes care of the size dependence of $k$ but not its potential variation from experiment to experiment. Ng et al. (2007) and Wang et al. (2018) determined a $k(D_p)$ profile using the initial seed wall-loss period for each of their experiments, thus accounting for the experiment-to-experiment variation.

The OA/Sulfate correction method was proposed by Hildebrandt et al. (2009) using the organic and the sulfate mass concentration measured by the aerosol mass spectrometer (AMS). This approach assumes that the loss rate constant of organic species and sulfate are the same during an experiment as there are no processes affecting sulfate other than losses to the walls (e.g., no added SO$_2$ or other sulfate precursor). The corrected OA mass concentration is then calculated as,

$$C_{OA}(t) = \frac{C_{OA}^{sus}(t)}{C_{seed}(t)} C_{seed}(0)$$

(4)
where \( \frac{C_{\text{OA}}^{\text{sulf}}(t)}{C_{\text{seed}}(t)} \) is the AMS-derived organic to sulfate ratio and \( C_{\text{seed}}(0) \) is the seed concentration in the chamber when SOA formation starts. Several chamber studies have adopted this method (Henry and Donahue, 2012; Loza et al., 2012). Other variations of this method including use the ratio of OA to other inert tracers like black carbon (BC) which are present in experiments investigating the evolution of primary OA from combustion sources (Hennigan et al., 2011). This method involving the use of OA-to-tracer ratio is accurate when the OA and the tracer have the same size distribution during the experiment or when the loss rate constant is close to size independent. However, in experiments in which SOA condenses more onto smaller-sized particles, the size dependence of the loss rate can introduce significant uncertainty in the corrected results especially for time scales of several hours (Wang et al., 2018).

An alternative method for particle wall-loss correction is the use of models of aerosol dynamics. Pierce et al. (2008) developed the Aerosol Parameter Estimation (APE) model that simulates the processes of condensation/evaporation, coagulation and particle wall loss during a chamber experiment. By constraining the unknown parameters with the SMPS-measured particle size distribution, the model can predict SOA formation for each experiment accounting for wall losses. The predicted particle wall-loss rates are both size- and time- dependent. The APE model predicts the particle wall-loss rates by assuming specific functional forms of its dependence on particle size (Crump and Seinfeld, 1981). The model has performed well in experiments in which the reaction time scale was short, but produced more uncertain results in experiments with slower reacting systems. Nah et al. (2017) adopted a modified version of the APE model that calculates the size-dependent wall-loss rate necessary to reproduce the observed size distribution assuming Brownian coagulation was the only other particle process occurring in the chamber (i.e. no condensation/evaporation occurred during the analyzed portion of the experiment). The size-dependent, instantaneous particle loss rates were calculated directly from the SMPS-measured seed number size distribution at each time step. These instantaneous \( k(D_p) \) values were then averaged over the initial seed loss period of the experiment (or a separate experiment where \( k(D_p) \) was characterized). This determined \( k(D_p) \) can then be applied to the SOA formation period of experiments to correct for the size-dependent wall loss. This approach, focusing on specific wall-loss characterization experiments, has the advantage that the functional dependence of the wall-loss rate constant is directly calculated from the measurements by simply removing the effect of coagulation. Its disadvantage compared to APE is that it requires additional time/experiments for
seed measurements and can no longer address the potential time dependence of $k$ over the course of a complex experiment.

The aforementioned methods each has its own advantages and disadvantages, and may perform well for specific experiments and chambers. However, for long-lasting experiments such as SOA aging where particle size distribution may shift across a wide size range due to several generations of condensation, it is important to address both the time- and size-dependence of the particle loss rates for the purpose of SOA quantification. In this work, we adopt the modified APE model following Nah et al. (2017) and derived the size-dependent particle loss rate constants, $k(D_p)$, based on seed periods during the experiments. As an attempt to evaluate the time-dependence of the loss rates, we derive a second $k(D_p)$ at the end of each experiment with a second seed injection and loss characterization period. To probe the effect of electrostatic forces on particle wall loss, we regularly measured the $k(D_p)$ during the time period when the chamber was experiencing changes (e.g., changes in its surroundings, location or air motion inside). We explore the coagulation effect on the estimated particle wall-loss rates and particle number/volume concentration in both a 12 m$^3$ Teflon chamber and a smaller 1.5 m$^3$ Teflon reactor. We evaluate the performance of the aforementioned particle wall-loss correction methods for relatively complex aging experiments involving 2 or 3 generations of condensation of the $\alpha$-pinene ozonolysis products.

2. Experimental approach

2.1 Particle wall-loss rate constant measurements in the 12 m$^3$ CMU Teflon chamber

The CMU smog chamber is a 12 m$^3$ Teflon reactor (Welch Fluorocarbons) suspended in a temperature-controlled room. The same Teflon reactor was used in all experiments discussed in this work. Chamber is not renewed during the time span in the discussion of the paper. The walls of the room are covered with UV lights (GE10526 and 10244). Prior to each experiment, the chamber is flushed overnight with purified air under UV illumination to remove any residual particles and vapors. Purified air is generated by passing house air through a high-efficiency particulate air (HEPA) filter to remove particles, a Purafil filter to remove NO$_x$ and an activated carbon filter to remove any organics followed by a silica gel filter, keeping relative humidity (RH) below 20% throughout the experiments performed in this work.
A list of experiments performed in this work is presented in Table 1, together with the experimental conditions. To characterize the particle wall-loss rates, we performed experiments with ammonium sulfate (AS) particles during a full year, with one listed in Table 1. We used both 1 g L\(^{-1}\) and 5 g L\(^{-1}\) ammonium sulfate solutions to generate particles, with the latter producing more particles at larger sizes. The ammonium sulfate solution was fed to an atomizer (TSI, model 3076) at a constant rate of 90 mL h\(^{-1}\) to produce droplets. The droplets passed through a diffusion dryer and a neutralizer to produce dry ammonium sulfate particles. This process produced seeds with a number mode diameter of around 100 nm. The initial seed number concentration in the chamber ranged from 2-5×10\(^4\) cm\(^{-3}\), corresponding to mass concentration of 40-200 µg m\(^{-3}\) and surface area concentration of 1100-4600 µm\(^2\) cm\(^{-3}\). After injecting the particles, the particle wall loss was quantified for 3-4 hours. We measured the particle size distribution with a Scanning Mobility Particle Sizer (SMPS, TSI classifier model 3080; DMA model 3081; CPC model 3010 or 3772). The CPC was set to sample at a sampling rate was 0.3 L min\(^{-1}\) for experiments in both chambers. For a 5-hour experiment, the volume lost due to sampling is 0.09 m\(^3\). This is a small change for both the 12 m\(^3\) and the 1.5 m\(^3\) chambers. The SMPS scanning time was 5 min for each sample.

For a number of α-pinene ozonolysis experiments, we characterized the particle wall-loss rates twice, once before and once after the main experiment. The rationale behind the second injection is to ensure that the wall loss rate constant profile remains relatively consistent (no major change) throughout each experiment. Due to the length of these aging experiments, few particles were left after the main experiment for a robust characterization of the profile, and thus we inject additional seed particles for a second time. At the beginning of each aging experiment, we used the 1 g L\(^{-1}\) ammonium sulfate solution to generate seed particles to provide enough surface area for the vapors to condense on. From the 3-4 h wall-loss time period, we were able to characterize the initial \(k(D_p)\) profile for this experiment. At the end of each experiment, we injected additional ammonium sulfate seeds into the chamber using the same method with a more concentrated solution (5 g L\(^{-1}\)) (to generate bigger particles) in order to characterize the particle wall-loss rate constants a second time. Details about the experimental procedure for these aging experiments can be found in Wang et al. (2018).

### 2.2 Particle wall-loss rate constant measurements in a 1.5 m\(^3\) Teflon reactor
We performed additional particle wall-loss measurements in a dual smog chamber system consisting of a set of two identical Teflon chambers (1.5 m³ each). The two pillow-shaped chambers are mounted on metal frames with wheels on the bottom for portability. Details about the portable dual-chamber system can be found in Kaltsonoudis et al. (2018). The system was tested inside the laboratory in the present study. We used both ammonium sulfate and PSL particles (700 nm, Duke Scientific Corporation) for measurements in these chambers.

3. Data analysis
3.1 Particle loss rate constants

The particle loss rate constants derived from methods accounting for the coagulation of particles are denoted as \( k_c \) in the rest of the paper, while the apparent particle loss rate constants neglecting the role of coagulation as \( k_a \). Our analysis here focuses on the losses of the particles and not on the potential interactions approach implicitly assumes that of the particles deposited on the walls are removed from the system and stop interacting with the gas phase (Hildebrandt et al., 2009).

3.1.1 The coagulation-corrected particle loss rate constants, \( k_c(D_p) \)

The coagulation-corrected particle wall-loss rate constants were derived based on the model used by Nah et al. (2017). The model assumes that only two processes take place: particle wall loss and coagulation. With a given particle number size distribution at a specific time step, the model predicts how the distribution evolves at the next step assuming coagulation is the only process based on the Brownian coagulation kernel in Seinfeld and Pandis (2006). The model attributes the difference between the predicted size distribution and the measured one to particle wall loss. Then the model calculates the instantaneous wall-loss rate constants at each time step for each size. To obtain the coagulation-corrected particle wall-loss rate constant, \( k_c(D_p) \), the instantaneous rate constants are averaged over time. The reported uncertainty is calculated as \( \sigma_{k_c}/\sqrt{N-1} \) where \( \sigma_{k_c} \) is the standard deviation of \( k_c \) for a certain size bin and \( N \) is the total.
number of time steps used. To minimize the uncertainty of $k_c$, we used a time step of 15 min (3 SMPS samples) for the averaging of the measurements (3 SMPS samples). Only SMPS measurements from the seed wall-loss periods were used as the inputs for the model.

3.1.2 The apparent particle wall-loss rate constants, $k_a(D_p)$

The apparent particle loss rate constants, denoted as $k_a$ in this work, were derived from the size-dependent loss rate constant. Details can be found in Wang et al. (2018). This approach uses the SMPS-measured seed particle number size distribution as input, and calculates a first-order loss rate constant for particles of a certain size across all measured sizes. These $k_a$’s intrinsically represent the combined loss effect of both particle wall loss and coagulation.

For particles in size bin $i$, $k_{a,i}$ is found by:

$$\ln[N_{i_{sus}}(t)] = -k_{a,i}t + Q$$

(5)

where $N_{i_{sus}}(t)$ is the suspended aerosol number concentration at size bin $i$ measured by SMPS and $Q$ is an arbitrary constant.

3.2 Size-dependent particle loss correction

Both $k_c$ and $k_a$ are size-dependent. The corrected particle number concentration at size bin $i$, $N_{i_{tot}}(t)$, is calculated by,

$$N_{i_{tot}}(t) = N_{i_{sus}}(t) + k_{a/c,i} \int_0^t N_{i_{sus}}(t) dt.$$  

(6)

We can then derive the corrected particle volume concentration of size bin $i$, $V_{i_{tot}}(t)$.

The organic aerosol mass concentration corrected for wall losses can be then calculated during a seeded experiment using:

$$C_{SOA_{tot}}(t) = (V_{tot}(t) - V_{s}) \rho_{SOA},$$

(7)

where $V_{tot}(t)$ is the corrected total particle volume concentration summed across all sizes and $V_{s}$ is the corrected seed volume concentration right before SOA formation. For all practical purposes, $V_{s}$ is constant during the seed-only period after our particle wall-loss corrections.

4. Results and discussion

4.1 Role of coagulation in particle wall-loss processes
Fig. 1 shows the apparent \( (k_a) \) and coagulation-corrected \( (k_c) \) particle wall-loss rate constants as a function of particle size for the 1.5 m\(^3\) Teflon reactor (Exp. 1) and the 12 m\(^3\) chamber (Exp. 2) after both systems have remained undisturbed in the lab for weeks. The particle loss rate constants for a given size can be measured reliably only when there are enough particles of this size available in the system. In the experiments shown, the produced ammonium sulfate particle size distribution included few larger particles. As a result, the \( k \)'s at bigger particle sizes were quite had high uncertainty associated with them due to lack of particles, and are not shown were thus excluded from the figure. Since the difference in \( k_a \) and \( k_c \) are attributed to coagulation according to the aerosol dynamics model, coagulation was a significant loss process in Exp. 1 for particles with diameters smaller than 250 nm and for particles smaller than 150 nm in Exp. 2. In Exp. 1, the apparent loss rate constant for 100 nm particles was 0.5 h\(^{-1}\), while the actual rate constant after correcting for coagulation was only 0.2 h\(^{-1}\). For 200 nm particles, the corresponding values were 0.3 h\(^{-1}\) and 0.2 h\(^{-1}\) respectively. Note that the initial particle number concentration in Exp. 1 wasis one order of magnitude higher than that in Exp. 2 (Table 1), and thus coagulation played a more prominent role in Exp. 1. The coagulation effects were minor for particles larger than 250 nm in both cases. Once corrected for coagulation, the particle wall-loss rate constants indicated little size dependence for particles larger than 100 nm in both experiments. The corresponding values were 0.25 h\(^{-1}\) for the small reactor, and 0.1 h\(^{-1}\) for the 12 m\(^3\) chamber. The uncertainty of the aerosol dynamics model for the larger particles is significantly higher than that of the direct calculation based on Eq. 5 (for example, at \( D_p = 790 \) nm in the 1.5 m\(^3\) reactor for Exp. 1 the 1.5 m\(^3\) reactor the uncertainty of \( k_c \) is has 50 % while that of \( k_a \) only uncertainty vs. 7 %, for \( k_a \)). This is due to the reliance of the dynamics model on observed small changes of small number concentrations versus the linear regression that uses fit of the logarithm of the measured particle concentration in each size bin values. We suggest using twice the uncertainty of the linear regression as representative of the uncertainty of the rate loss constants of particles larger than 300 nm given the small impact of coagulation on particle concentrations and sizes in this range.

To evaluate the coagulation effect on particle number and volume concentrations, we corrected them for wall loss with both \( k_a(D_p) \) and \( k_c(D_p) \). The results for the 1.5 m\(^3\) reactor (Exp. 1) are shown in Fig. 2 as an example. The measured \( k_a \)'s were extremely uncertain due to lack of particles at small sizes and thus could not be used. To estimate the \( k_a(D_p) \)'s at \( D_p < 50 \) nm (and the \( k_c(D_p) \)'s at \( D_p < 70 \) nm), we used a linear fit of the \( k_a(D_p) \)'s from 50 to 70 nm (and of the \( k_c(D_p) \)'s
from 70 to 100 nm) to back extrapolate the $k_a$’s at smaller sizes. $k_c$, being the coagulation-corrected particle wall-loss rate constant, intrinsically excludes the impact of coagulation on particle number concentration. Since coagulation reduces particle number (but conserves mass), $k_c$-corrected particle number concentration is lower than the $k_a$-corrected one, with the difference attributed to the coagulation rate. Coagulation caused the particle number concentration to decrease by 27 % over a 5 h period in this case, but had, as expected, negligible effect on particle volume concentration. Please note that even if the overall effect of coagulation on total particle number is moderate, it is mostly concentrated in the lower end of the size distribution. As a result, the difference between $k_a$ and $k_c$ due to coagulation for particles smaller than 100 nm effect is almost an order of magnitude higher than the average difference for the rate constants of particles smaller than 100 nm. In this case there is little difference in the calculated total volume concentration, which is the most important quantity for SOA studies. However, this difference depends in general on the particle size distribution. If a significant part of the volume (or mass) is in particles with diameters less than 200 nm or so, the effect of coagulation will be significant for the corrected particle volume too. Nah et al. (2017) also studied the effect of coagulation on corrected SOA volume for the α-pinene ozonolysis system, and found that coagulation plays minor role in experiments with initial seed surface area of <3000 $\mu$m$^2$ cm$^{-3}$ while in experiments with high seed surface area (>8000 $\mu$m$^2$ cm$^{-3}$), the SOA can be substantially overestimated if one ignores coagulation. Our results are consistent with their low seed surface area experiments.

4.2 Particle wall-loss rate constants in the CMU chamber over three years

Fig. 3 shows the coagulation corrected $k_c(D_p)$ profiles together with their corresponding uncertainties in the 12 m$^3$ CMU smog chamber over a span of three years (Fig. S1 shows the uncertainty). All these measurements were performed during periods in which the chamber was undisturbed. These four curves are representative of the variability range of the size-dependent particle wall-loss rates in the CMU chamber under undisturbed conditions. The rate constants show a monotonic decreasing trend with sharp decrease initially until 100 nm due to diffusion dominating the wall-loss processes. Then the loss rate constants gradually decrease until 300 nm, after which they stay almost constant until the end of the measured size range. Using the $k_c(D_p)$ determined in 2017 as an example, $k_c$ decreased from 0.3 h$^{-1}$ at 50 nm, to 0.14 h$^{-1}$ at 100 nm, then gradually to 0.05 h$^{-1}$ at 300 nm and stayed constant until approximately 500 nm. The $k_c(D_p)$ profiles
over the past three years stayed fairly consistent with values equal to 0.32±0.03 h\(^{-1}\) at 50 nm, 0.16±0.03 h\(^{-1}\) at 100 nm, 0.10±0.02 h\(^{-1}\) at 200 nm and 0.07±0.01 h\(^{-1}\) at 300 nm. The behavior of the chamber after disturbances (e.g., repairs, upgrades) will be discussed in a subsequent section.

4.3 Applying different particle wall-loss correction methods to SOA aging experiments

The measured particle volume concentration time series of a typical aging experiment (Exp. 3) of α-pinene ozonolysis products in the 12 m\(^3\) CMU Teflon chamber is shown in Fig. 4. In this experiment there were three separate stages. We injected ammonium sulfate seeds both at the beginning (\(t = -4.5\) h) of the experiment and at \(t = 3.5\) h. At \(t = 0\), ozone was introduced into the chamber to react with α-pinene producing SOA in the dark. HONO was bubbled into the chamber twice at \(t = 0.5\) h and 1.2 h to produce OH radicals under UV illumination, leading to a second round of reactions in the system. The size-dependent \(k_a(D_p)\) and \(k_c(D_p)\) derived from the initial 4.5 h seed loss period differed by up to 0.2 h\(^{-1}\) for particles smaller than 100 nm and were practically the same in absolute values for particles larger than 100 nm (Fig. 4c). The size-independent loss rate constants \(k_1 - k_3\) were derived during the three periods when condensation/evaporation was minimal (based on both SMPS and AMS measurements). A value of \(k_1 = 0.05\) h\(^{-1}\) \((R^2 = 1)\) was derived from volume concentration measurements from \(t = -4.5\) to 0 h according to Eq. 2, \(k_2 = 0.04\) h\(^{-1}\) \((R^2 = 0.8)\) from \(t = 2\) to 3.4 h and \(k_3 = 0.03\) h\(^{-1}\) \((R^2 = 0.9)\) from \(t = 4.7\) to 8.4 h. One major contributor to the difference in these three \(k\)’s is the size dependence of the particle wall-loss rate constants. \(k_2\) was calculated from the period after three rounds of condensation (α-pinene ozonolysis and two doses of HONO). The particle size distribution shifted to larger sizes (Fig. 4b) and thus resulted in a smaller value compared to \(k_1\). \(k_3\) was derived from the final seed loss period when relatively large seed particles were present due to the higher concentration of the atomized ammonium sulfate solution. To prove this mathematically, we define a mass-weighted wall-loss rate constant, \(\bar{k}_{a,\text{period } 1-3}\), for each period. For each size bin \(i\) and a total of \(n\) size bins,

\[
\bar{k}_{a,\text{period } 1-3} = \frac{\sum_{i=1}^{n} v_i k_{a,i}}{\sum_{i=1}^{n} v_i} \tag{8}
\]

where \(v_i\) is the volume concentration for particles in size bin \(i\) and \(k_{a,i}\) is the particle loss rate constant for size bin \(i\). For Exp. 3, \(k_a\) for each size bin and a total of 110 size bins are shown in Fig. S2. And the volume concentration for particles in each size bin are taken from SMPS measurements averaged over the 3 periods, respectively. \(\bar{k}_{a,\text{period } 1-3}\) is found to be 0.6 h\(^{-1}\).
\[
\overline{k}_{\text{period}2} = 0.4 \, h^{-1} \quad \text{and} \quad \overline{k}_{\text{period}3} = 0.4 \, h^{-1} - \overline{k}_{\text{period}1} \quad \text{should be comparable to} \quad k_1 \quad \text{while} \\
\overline{k}_{\text{period}2} \quad \text{to} \quad k_2 \quad \text{and} \quad \overline{k}_{\text{period}3} \quad \text{to} \quad k_3. \quad \text{The} \ 0.1 \, h^{-1} \quad \text{difference between} \quad \overline{k}_{\text{period}1} \quad (\overline{k}_{\text{period}3}) \quad \text{and} \quad k_1 (k_3) \quad \text{can be attributed to the uncertainty of} \quad k_\alpha \quad \text{at each size bin (Fig. S2). However, the fact that} \\
\overline{k}_{\text{period}2} \quad \text{is} \quad 0.2 \, h^{-1} \quad \text{smaller than} \quad \overline{k}_{\text{period}1} \quad \text{at least show that the size dependence of} \quad k_\alpha \quad \text{is sufficient to explain the difference between} \quad k_1 - k_3. \\
\]

The particle wall-loss corrected SOA mass concentration (\( \rho = 1.4 \, \mu g \, m^{-3} \)) time series based on SMPS measurements using both the size-independent \( k_1 - k_3 \) and the size-dependent \( k_\alpha (D_p) \) and \( k_c (D_p) \) for Exp. 3 are shown in Fig. 5. Applying \( k_1 - k_3 \) to Eq. 3 resulted in corrected SOA mass concentration differing up to 20%. To estimate the \( k(D_p) \)'s at \( D_p < 50 \, nm \) and \( D_p > 300 \, nm \), we used a linear fit of the \( k(D_p) \)'s from 50 to 70 nm to back extrapolate the \( k \)'s at smaller sizes and assumed a constant \( k \) value equal to that at 300 nm for particles larger than 300 nm (Fig. S2). This assumption is justified because \( k_c \) remains relatively constant from format the 300-500 nm range based on our measurements (Fig. 3) when the chamber is undisturbed, and most of the SOA mass in this experiment was distributed in particles smaller than 300 nm (Fig. 4b). We corrected for total particle number concentration applying the size-dependent loss rates to Eq. 6, and then calculated the corrected SOA mass concentration using Eq. 7. Note that the \( k_\alpha (D_p) \)- and the \( k_c (D_p) \)-corrected SOA mass concentration time series were practically the same for this experiment, because the majority of the formed SOA mass condensed on particles with diameters exceeding 100 nm. If one is interested in the total produced SOA after 3.5 hours the differences among the results of the different corrections are 20% or less. If one is interested in the SOA produced during the aging phases the estimates vary by 25-30%.

Depending on which wall-loss rate constants are used, the corrected SOA mass concentration can vary by 20-30%. We recommend using the size-dependent wall-loss rate constants for the correction. However, when the chamber is undisturbed and the duration of the experiment lasts only a couple of hours, using the size-independent wall-loss rate constant derived from the initial 4-hour seed wall-loss period can give relatively accurate results (errors of 5% or less).

4.4 Effect of size-dependent wall loss on organic to sulfate ratio
Fig. 6 shows the AMS-measured organic to sulfate ratio (OA/Sulfate) for Exp. 34. In the beginning, the ratio increased to 0.84,2 at t=0.6 h due to the first generation of SOA formation. It then stayed practically constant until OH was introduced into the chamber at t=1.709 h. The second-generation of SOA formation led to an increase of the ratio to 1.04,6 at t=2.04,4 h. The ratio decreased gradually to 0.9 at t=3.5 h until the second introduction of HONO. This decrease could be explained as a loss of SOA due to photodegradation or other chemical processes such as SOA evaporation driven by organic-vapor uptake by the walls (Bian et al., 2015). Another explanation for the decreasing trend of OA/Sulfate during this period is the size dependence of the particle wall-loss rates (Loza et al., 2012). Particles of smaller sizes with higher organic to sulfate ratios can be lost to the walls at a faster rate, thus causing the OA/Sulfate to decrease during periods when wall loss is the dominant process in the chamber. The strong size dependence of the OA/Sulfate ratio in this experiment is indicated in Fig. 7. The organic mass distribution peaked at an aerodynamic vacuum diameter ($D_{va}=D_{vac,particle}$) equal to around 150300 nm, while the sulfate one at 210450 nm. This indicates that the majority of the organic vapors condensed onto smaller particles with a higher surface to volume ratio. Fig. 7b shows the OA/Sulfate derived from the AMS-measured mass distribution (averaged from t=2.14,4 to 3.5 h) as a function of the particle vacuum aerodynamic diameter. For particles with $D_{va}$ from 75200 to 150300 nm, the ratio dropped dramatically from 740 to 13. It then decreased gradually and stabilized at 0.4+ from $D_{va}$=150300 to 6004000 nm.

To further analyze the effect of size-dependent wall loss on OA/Sulfate, we adopted the method suggested in Wang et al. (2018). This approach allows the estimation of mass-weighted wall-loss rate constants for both species, $k_{SO_4}$ and $k_{OA}$, by discretizing the AMS-measured mass distribution in the diameter space and assigning the corresponding $k_c(D_p)$. (An SOA density of 1.4 μg cm$^{-3}$ was used to convert the AMS-measured $D_{va}$ to $D_p$). For periods during the experiment when particle wall-loss is the only process, the loss-corrected OA/Sulfate can be estimated as: $\text{(OA/Sulfate)}_m(t)\exp(\bar{k}_{SO_4} - \bar{k}_{OA})t$, where $(\text{OA/Sulfate})_m(t)$ is the measured OA/Sulfate.

For Exp. 43 in this work, we discretized the AMS-measured mass distribution (averaged from t=2.14,2 to 3.54,7 h) into 10 diameter bins and found $\bar{k}_{org} = 0.130,06$ h$^{-1}$ and $\bar{k}_{SO_4} = 0.1105$ h$^{-1}$. The particle wall-loss corrected OA/Sulfate for the chosen time period is shown in the inset of Fig. 6. The correction explains more than 70% of the decrease in the OA/Sulfate (over 1.4 h) in this experiment. Loss-corrected ratio remained relatively constant, indicating that the
decrease observed in the measured OA/Sulfate was caused by the size-dependent particle wall-loss process coupled with the different size distributions of the organics and sulfate played a major role in the observed decrease in OA/Sulfate.

### 4.5 Time dependence of particle wall-loss rates during an experiment

When the CMU chamber is undisturbed the wall loss rate constant is around 0.1 h\(^{-1}\) for particles larger than 100 nm. However, friction with the Teflon walls induced by small repairs (addition of a sampling line, replacement of lights, etc.) around the chamber can increase the loss rates dramatically and the effects can last for weeks. During these periods the size-dependent coagulation-corrected particle wall-loss rate constants, \(k_c(D_p)\), can change significantly during the course of an experiment. The results of such an experiment in a “disturbed” smog chamber are described below.

The comparison of the two \(k_c(D_p)\) profiles derived from the initial and the final seed periods for Exp. 54 are shown in Fig. 8, together with the raw and the corrected aerosol volume concentration time series. This is a similar aging experiment of \(\alpha\)-pinene ozonolysis products as Exp. 3, but with only one HONO injection. Before \(t=0\), ammonium sulfate seed particles were lost to chamber walls. At \(t=0\), ozone was added into the chamber to react with \(\alpha\)-pinene. The aerosol volume increased due to condensation of the first-generation products. At \(t=2.5\) h, HONO was introduced into the chamber and OH were produced at \(t=3\) h under UV illumination. The aerosol volume increased again due to additional SOA formation from the second-generation oxidation. At \(t=4\) h, we injected ammonium sulfate particles into the chamber to characterize the particle wall loss rates for a second time. The final \(k_c\)’s were statistically higher than their initial counterparts at every size, and both sets of \(k_c\)’s were higher than their usual values in the chamber (Fig. 3). Comparing the initial \(k_c\)’s with the averaged usual values under undisturbed chamber conditions, the initial \(k_c\) was 0.33 h\(^{-1}\) as compared to the usual 0.16 h\(^{-1}\) at 100 nm, 0.21 h\(^{-1}\) compared to 0.10 h\(^{-1}\) at 200 nm, and 0.15 h\(^{-1}\) compared to 0.07 h\(^{-1}\) at 300 nm. The particle wall-loss correction was applied when the ammonium sulfate seed was first injected into the chamber (as depicted in Figure 8a). The final \(k_c(D_p)\)-corrected volume concentration was higher than the one corrected using the initial \(k_c(D_p)\) by 37 % at \(t=4\) h. In this case, both \(k_c\) profiles were representative of the chamber condition, but during their corresponding time periods. The time dependence of \(k_c(D_p)\) during the course of this experiment introduced a 40% or so uncertainty in the corrected aerosol
mass or volume concentration. The change in the loss rates suggests a change in the electric field in the chamber during this experiment. This could be due to additional charge build-up or redistribution of the charges as the experiment progresses (lights are turned on and off, the chamber walls move due to the air motion from the temperature control system, etc.).

We define the chamber conditions under which these abnormally high loss rates and exacerbated time dependence of $k_c(D_p)$ were observed as “disturbed”. The $k_c(D_p)$ profiles shown in Fig. 3 were under “undisturbed” chamber conditions. Since electrostatic forces start to dominate the wall-loss process when particles are usually larger than 100 nm (McMurry and Rader, 1985), we postulate that excess electrostatic forces within the chamber are most likely the cause of the “disturbed” conditions. Friction created with the Teflon walls was found to be a major contributor to the exacerbated electrostatic forces and the “disturbed” chamber conditions.

Fig. 9 shows the $k_c(D_p)$ profiles measured over a span of five months after some major maintenance work (Jan. 2016) in the room where the chamber is suspended (Fig. S3 shows the uncertainty). During the one-week maintenance, friction with Teflon walls was created by partially deflating the chamber, moving and touching it repeatedly. The measured $k_c(D_p)$ profile changed drastically in shape for days after. The 16-day post-maintenance $k_c(D_p)$ profile presented an increasing trend from 75 nm to 300 nm, with particles bigger than 200 nm getting lost at a rate 3-4 times faster than before. Once we noticed the abnormally high particle loss rates in the chamber, we refrained from being in any form of contact with the chamber walls. The chamber was left suspended and full during those five months. Records of experiments performed in the chamber during that time support the previous statements. About a month later, the $k_c$’s recovered to the decreasing trend, but were in general high as compared to their pre-maintenance counterparts with values $> 0.2$ h$^{-1}$ at 300 nm. Three months after the maintenance, particles smaller than 100 nm recovered to its pre-maintenance values while particles bigger than 150 nm still had loss rate constants up to 0.1 h$^{-1}$ higher than before. Five months after, the $k_c(D_p)$’s made a full recovery, with values decreased further to 0.13 h$^{-1}$ at 100 nm, 0.09 h$^{-1}$ at 200 nm and 0.06 h$^{-1}$ at 300 nm. During the 5 months, the chamber was left fully inflated, stationary and suspended in the room. Only steps necessary for an experiment (overnight flushing, injection flow etc.) were taken. Exp. 54 was performed a month after the major maintenance when $k_c(D_p)$ was still in recovery, and thus the exacerbated electrostatic forces within the chamber likely played a major role in the extra time sensitivity of the $k_c(D_p)$’s in Exp. 54.
To test whether certain steps during an experiment cause changes in the particle loss rates, we explored potential impact on $k_c(D_p)$ of turning on the UV lights, injecting HONO and overnight flushing individually in separate seed experiments. Experiments 5-8 were designed to test each of these factors individually. These experiments were performed about 3 months after the maintenance when particle wall loss rates have almost recovered to its pre-maintenance values, indicating the chamber has mostly recovered to “undisturbed” conditions. Turning on the UV lights inside the room where the chamber is suspended can cause changes in the air circulation around the chamber walls, thus affecting the turbulence. Carrying HONO into the chamber with a clean air flow at a rate of ~ 5 L min$^{-1}$ for 20 mins may potentially impact the turbulence within the chamber. Though cleaning the chamber with overnight flushing may not have a direct impact of $k_c$ during the day of an experiment, flow rates higher than 100 L min$^{-1}$ into the chamber may well have an effect. The results of Experiments 5-8 are shown in Fig. 10 and none of the aforementioned processes had evident impact on $k_c(D_p)$. t-test results indicated that of the $k_c(D_p)$ profiles derived before and after each factor were statistically the same. We thus conclude that the usual steps taken during a typical SOA aging experiment do not have a significant impact on $k_c$ if the chamber is in its undisturbed state. However, when the chamber has been disturbed and the losses are already high they also become sensitive to routine changes in the experimental conditions.

4.6 Teflon chamber maintenance and operating procedure of chamber experiments

Routine seed experiments appear to be necessary for the quantification of the particle loss rates in Teflon chambers. Any deviation in the particle wall-loss rate constants from the usual values can be a sign of “disturbed” chamber conditions, which may result in higher particle loss rates and time sensitivity of $k_c(D_p)$ during an experiment. As discussed above, friction with the chamber walls can introduce excess electrostatic forces within the chamber and thus introduce significant uncertainty in the particle loss rates. In order to maintain minimum particle loss in Teflon chambers, one should refrain from creating any type of friction with the chamber walls such as touching the walls or having the walls rubbing against each other. When transporting the chambers such the dual-chamber system, it is ideal to leave them full or at least half-filled with air and fixed onto a rigid structure that can be packed during the transportation. This can minimize potential friction and shorten the recovery time for the particle loss rates. Other practices like using
metal gloves when it is absolutely necessary to touch the chamber can help reduce the build-up of static electricity on the chamber walls. Use of metal gloves is recommended when it is absolutely necessary to touch the chamber.

When the chamber is in a “disturbed” state, the $k_c(D_p)$’s can vary with time during the course of an experiment. It is thus vital in these cases to include two seed loss periods, one at the beginning and the other at the end, for each SOA experiment to characterize the $k_c(D_p)$, especially if the chambers are recently subjected to friction. When performing SOA experiments in a Teflon chamber, we recommend the following operating procedure:

1. injection of seeds and initial $k_c(D_p)$ characterization for 3-4 hours;
2. perform necessary steps for the SOA experiment and wait until the mass loadings in the chamber become low;
3. if the losses in step 1 are high, a second injection and another 3-4 hours of measurements for final $k_c(D_p)$ characterization are necessary.

5. Conclusions

Particle number losses in chamber experiments due to coagulation can be significant for small particles (< 150 nm under conditions in this work). It is thus important to correct for this coagulation effect when calculating the particle wall-loss rate constants especially for experiments in which the behavior of the nanoparticles is important (e.g., when they carry a significant fraction of the total particle mass).

The Teflon chamber used in this study appeared to operate in two different states: an undisturbed and a disturbed one. The chamber entered the second state after either major repairs or even after smaller changes (e.g., addition of a sampling line or replacement of a few lights) probably because it was touched by the researchers or because friction was created during the repairs causing charge build-up. The disturbed state could last for several weeks or even months. In this state the particle loss rates increased by more than a factor of 3-4 and their size dependence became more pronounced. There was significant variation of the losses from experiment to experiment and even within the same experiment. In the undisturbed state, the loss rate constant was less than 0.1 h$^{-1}$ for particles larger than approximately 200 nm and was constant from experiment to experiment. Under these conditions the cleaning of the chamber, turning on the
chamber lights, injection of reactants, etc., did not have a statistically significant impact on the loss rate constants.

The accuracy of the use of size-independent loss rate constants for the correction of the experimental results depends on the state of the chamber and the size distribution of the aerosol during the experiment. If the aerosol volume is dominated by particles larger than 200 nm and the chamber is undisturbed, the corresponding results can be quite accurate under conditions in this work. However, if the chamber has been disturbed or if the size distribution during some phase of the experiment includes a lot of ultrafine particles, significant errors can be introduced.

The correction based on the OA/Sulfate ratio can also introduce uncertainties under at least some conditions. The SOA mass distribution is usually shifted towards the smaller particles compared to that of the sulfate seeds. As a result, the losses of sulfate can be different than those of the organics. The sign and the magnitude of the error depends on both the differences between the two size distributions and also the size dependence of the losses in this specific experiment. A method to correct the OA/Sulfate ratio for these effects has been developed. In one of the experiment, this explains 70% of the apparent decreases of the ratio from 1.0 to 0.9 in less than one hour. The errors appear to be of the order of 20% or less, but may lead to problematic conclusions about potential processes taking place at longer timescales (e.g., photolysis and loss of SOA). Corrections similar to the one used in this work (taking into account the size-dependent losses and the size distributions of OA and sulfate) should be performed in the case of a disturbed chamber as a safeguard against higher errors.

Due to the above complexities, seed experiments for testing the particle loss rates in Teflon chambers should be performed regularly, probable before each experiment. If the rates are high a second measurement of the losses should be performed after the end of the experiment to constrain any potential changes. The use of size-dependent corrections accounting for coagulation effects is the preferred approach even if in a number of experiments when the chamber is undisturbed the errors introduced by neglecting the size dependence or the role of coagulation could be small. However, this depends a lot on the evolution of the aerosol volume distribution during the experiment and especially on the importance of the particles smaller than 200 nm or so for the objectives of the experiment.
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6. References


### Table 1: List of experiments and experimental conditions

<table>
<thead>
<tr>
<th>Exp.</th>
<th>Chamber volume (m³)</th>
<th>Type</th>
<th>Number of seed wall-loss periods</th>
<th>Initial seed number/surface/volume concentration</th>
<th>Notes</th>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(cm³)</td>
<td>(µm² cm⁻³)</td>
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<tr>
<td>1</td>
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<td>3.4×10⁴</td>
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Figure 1: The apparent (red symbols) and coagulation-corrected (black symbols) particle wall-loss rate constants as a function of particle size for a) the 1.5 m³ Teflon reactor and b) the 12 m³ CMU smog chamber after the two systems have been left undisturbed in the lab for weeks. The particle loss rate constants were derived based on SMPS measurements from Exp. 1 and Exp. 2. Only $k_a$’s with an $R^2 > 0.5$ are shown. The error bars correspond to one standard deviation. The grey area is the uncertainty associated with $k_c$. 

\[ k_a, k_c \]
**Figure 2:** The SMPS-measured (black symbols) and the particle loss corrected a) number and b) volume concentration using the $k_a(D_p)$ profile (red symbols) and the $k_c(D_p)$ profile (blue symbols) for Exp. 1.
Figure 3: The $k_c(D_p)$ profiles for the 12 m$^3$ CMU Teflon chamber over a span of three years. The particle wall-loss rate constants were derived based on SMPS measurements from experiments with only ammonium sulfate particles.
**Figure 4:** a) The SMPS-measured particle volume concentration time series for an aging experiment (Exp. 3) with 3 colored periods used to derive the corresponding size-independent particle wall-loss rate constants, $k_1$-$k_3$ (Eq. 2). The grey area indicates that the chamber was dark. The dashed lines mark the beginning and the end of bubbling HONO into the chamber twice; b) the averaged particle volume size distribution over the 3 periods used to develop $k_1$-$k_3$ based on the SMPS measurement for Exp. 3; c) the size-dependent particle wall-loss rate constants determined from SMPS-measured particle number concentration **from Period 1** for Exp. 3. Only $k_a$’s (red symbols) with an $R^2 > 0.5$ are shown. The error bars correspond to plus/minus one standard deviation. The grey area is the uncertainty associated with $k_c$ (black symbols).
**Figure 5:** Particle loss corrected SOA mass concentration ($\rho_{SOA} = 1.4 \, \mu g \, m^{-3}$) time series based on SMPS measurements using the size-independent $k$’s (open symbols) and the size-dependent $k(D_p)$’s (solid symbols) for Exp. 3. $k_1 - k_3$ were derived from the total mass concentration-based method (Eq. 2) when wall loss was the only process ($t_1 = -4.5-0h; \ t_2 = 2-3.4h; \ t_3 = 4.7-8.4h$). The $k_a(D_p)$ and the $k_c(D_p)$ profiles were derived from the two models based on the SMPS-measured number concentration of the seed wall-loss periods. The shaded area indicates that the chamber was dark. The dashed lines mark the beginning and the end of bubbling HONO into the chamber.
Figure 6: The organic to sulfate ratio time series derived from AMS measurements for Exp. 34 (data after the second HONO introduction is not shown). The inset is a blow-up of the OA/Sulfate from its maximum until the second HONO introduction. The black symbols are the size dependence corrected OA/Sulfate during that half hour. The shaded area indicates that the chamber was dark. The dashed lines mark the beginning and the end of the first HONO injection into the chamber.
a) $dM/d\log D_{va} \text{ (µg m}^{-3}\text{)}$

b) OA/Sulfate $D_{va}$ (nm)
Figure 7: a) The AMS-measured organic (green) and sulfate (red) mass distribution ($D_{va}$ from 75 to 600 nm) for Exp. 43; b) the dependence of the AMS-derived organic to sulfate ratio on particle vacuum aerodynamic diameter (75-600 nm). The results are based on particle time-of-flight (PToF) data averaged over ~2.51 ± 1.4 h (t=2.1-3.5 h).
Figure 8: a) The SMPS-measured (black symbols), the initial \( k_{c}(D_p) \)-corrected (red symbols) and the final \( k_{c}(D_p) \)-corrected (blue symbols) particle volume concentration time series for an aging experiment (Exp. 54), together with b) the initial (red symbols) and the final (blue symbols) \( k_{c}(D_p) \) profiles. The colored area are the uncertainties associated with the corresponding \( k_{c}(D_p) \). The grey area indicates that the chamber was dark. The two dashed lines mark the beginning and the end of HONO addition into the chamber. Ammonium sulfate seed particles were injected into the chamber at \( t=4 \) h.
Figure 9: The coagulation-corrected particle wall-loss rate constant, $k_c$, at each diameter derived from experiments with only ammonium sulfate particles in the 12 m$^3$ CMU Teflon chamber before and after some major maintenance in the room where the chamber is suspended. The chamber was partially deflated and its walls subjected to friction repeatedly during the maintenance.
Figure 10: The coagulation-corrected particle wall-loss rate constant, $k_c$, at each diameter for a) Exp. 56, b) Exp. 76, c) Exp. 87 and Exp. 98. The uncertainties associated with the corresponding $k_c(D_p)$ are either expressed as the grey area or the red error bars.