Timescale for hygroscopic conversion of calcite mineral particles through heterogeneous reaction with nitric acid

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Supplementary methods

Calibration of HNO$_3$ permeation tube and HNO$_3$ flow tube loss

The nitric acid permeation tube’s emission rate was empirically measured by bubbling 1.0 sLpm nitrogen gas from the HNO$_3$ source through the Teflon injector line into 500 mL of Na$_2$CO$_3$/NaHCO$_3$ buffer through a glass frit. 5.0 mL aliquots were removed every 30 minutes and the concentration of nitrate was determined by ion chromatography. The measured HNO$_3$ emission rate was highly linear with time ($R^2$=0.99) and equal to 370.4 ng/min (Fig. S1). The first two aliquots were excluded from the linear fit to eliminate the effect of HNO$_3$ ad/desorbing from the Teflon injection line and glass frit placed in the water, and to allow the system to reach steady-state. The measured HNO$_3$ emission rate is 73% of the manufacturer calibrated rate of 507 ng min$^{-1}$ at 30.0 C. This difference is attributed to irreversible loss of HNO$_3$ in the permeation tube holder and the lines and fittings between the source and end of the injection line.

The [HNO$_3$] exiting the kinetics flow tube was measured similar to the HNO$_3$ source calibration, by bubbling the HNO$_3$ containing gas flow exiting the flow tube into 500 mL of carbonate buffer, and determining the nitrate concentration via ion chromatography in 5.0 mL aliquots every 10 minutes. This produced a linear increase in [NO$_3^-$] with time ($R^2 > 0.95$), the slope of which corresponded to an observed HNO$_3$ emission rate of 325 ± 68 ng min$^{-1}$ (n=3). The difference between the HNO$_3$ source’s emission rate and that observed at the end of the flow tube is attributed to irreversible loss to the coated walls of the flow tube, and other surfaces.

CCN activation curve fitting

The lack of a clear minimum plateau from multiply-charged particles in the activation curves of reacted calcite aerosol (Fig. 5) makes it more difficult to accurately fit the curves with a sigmoid curve, defined as:

$$y = \frac{A_1 - A_2}{1 + e^{(x-x_o)/dx} - A_2}$$

where $y$ is the measured CCN/CN ratio, $A_1$ is the minimum value, $A_2$ is the maximum value, $dx$ is the rate of change, and $x_o$ is the midpoint between $A_1$ and $A_2$. The data is normally fit such that $A_1$ corresponds to the CCN/CN ratio of the multiply charged particles at small supersaturations, and $A_2$ is set to the scaled maximum CCN/CN ratio of ~1.0. Thus, $x_o$ corresponds to the CCN/CN=0.5 midpoint typically used to define the critical supersaturation ($s_c$) of the singly-charged particles of
interest. The lack of the multiply-charged plateau in the reacted aerosol makes it difficult to empirically constrain $A_1$ in this fit.

To resolve this issue the contribution of multiply-charged particles in the aerosol exiting the flow tube was directly measured using aerosol size distributions made with a scanning mobility particle sizer (SMPS) composed of a TSI Model 3081 DMA and TSI Model 3010 CPC. Aerosol and sheath flow rates of 0.40 and 4.0 sLpm were used for SMPS scans. The dried and denuded particles were sub-sampled by the SMPS after the flow tube. A typical size distribution is shown in Fig. S2a. The SMPS software’s multiple charge correction was not used here. As expected, the dominant mode is centred around 200 nm, which is produced by the original +1 charge 200 nm particles selected by the 1st DMA before the flow tube. The mode to the left are originally +1 200 nm particles that gain a +2 charge in the 2nd DMA, and therefore have a smaller mobility diameter, $D_m$. The mode on the right are originally +2 charge particles with $D_m = 200$ nm that only have +1 charge in the 2nd DMA; these represent the physically larger multiply-charged particles in the aerosol measured by the CCNc. The fraction of multiply charged particles was evaluated from the cumulative % size distribution shown in Fig. S2b. The fraction of particles larger than 250 nm was used to determine the fraction of multiply-charged particles in the aerosol and constrain the value of $A1$ in Eqn. S1. The resulting sigmoid fits are shown along with the data in Fig. 5. The uncertainty in the determination of $s_c$ is typically the product of the uncertainty in the CCNc’s SS calibration (3%), and in the particle size selection of the DMA ($\leq 5\%$). The lack of a multiply-charged plateau in the activation curve introduces additional uncertainty in the determination of $s_c$, though this is constrained by the SMPS measurement. The fraction of multiply-charged particles typically varied by 3% (absolute) between experiments. The value of $A1$ was varied by $\pm3\%$ (absolute) for fitting several typical activation curves with Eqn. (S1) and produced a change in $s_c$ no greater than $\pm0.01\%$ (absolute). A more conservative uncertainty of $s_c \pm0.02\%$ (absolute) is assigned for the reacted activation curves.

**Supplementary results and discussion**

**Partitioning of nitric acid to calcite under dry conditions**

The reactions performed at 0% RH produced a set of interesting results that suggest that under dry conditions the calcite particles became extensively coated by nitric acid that condensed on the partially reacted particles but that much of the HNO$_3$ was unable to actually react with the CaCO$_3$ core. Figure S3 displays two different ATPOMS nitrate measurements, in addition to the usual
NO$_2$/CaO$^+$ peak area ratio that has been used thus far (Fig. 4), for the 0% RH reactions. All three indicate that the nitrate signal at HNO$_3$ exposures $> 25$ ppb s was close to or larger than that measured from pure 200 nm Ca(NO$_3$)$_2$ particles. However, these particles never achieved critical supersaturations as small as pure calcium nitrate (Fig. S3d). The predicted critical supersaturation of 200 nm Ca(NO$_3$)$_2$ particles ($\kappa_{\text{app}} = 0.51$) is 0.058%, below the lower SS limit of the CCNc of $\sim 0.08\%$. Therefore, completely converted calcite particles that were 100% Ca(NO$_3$)$_2$ would be completely CCN active when measured in the CCNc, precluding the measurement of an activation curve. This was not observed, even at the largest HNO$_3$ exposure of 162 ppb s at 0% RH the particles were $\sim 40\%$ CCN-active at SS = 0.089%. However, the ATOFMS nitrate peak areas were much larger than from pure Ca(NO$_3$)$_2$ particles. The condensation of relatively large amounts of nitric acid to the surface of the partially reacted particle is the best explanation for this, particularly if HNO$_3$ produces larger $^{46}$NO$_2^-$ signals than Ca(NO$_3$)$_2$ does per molecule. The short reaction times in the flow tube may have limited the ability of the nitric acid reactant to react further with the calcium carbonate core. Additionally, the dry conditions may have limited the reaction to the particle surface if it became saturated with undissociated carbonic acid, as discussed in the main paper.
Supplementary figures

Figure S1. Calibration of HNO$_3$ permeation tube emission rate by bubbling gas stream into sodium carbonate buffer and measuring [NO$_3$] by ion chromatography in aliquots every 30 minutes. The first two aliquots (red symbol) are excluded from the linear fit.

Figure S2. (a) Typical SMPS size distribution of size-selected calcite aerosol after reaction in the flow tube. (b) The % cumulative plot is used to determine the fraction of multiply charged particles of $D_m > 250$ nm. The SMPS charge correction was not used here.
Figure S3. (a-c) Three ATOFMS nitrate metrics from calcite particles reacted with nitric acid at 0% relative humidity. Data from pure 200 nm calcium nitrate particles is included. (d) Critical supersaturation ($s_c$) of the same reacted particles versus the ATOFMS $\text{NO}_2^-/\text{CaO}^+$ absolute peak area ratio.